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REVISED AND ENTIRELY REWRITTEN (IN 1893)

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INTRODUCTION

TO THE ARTICLES RELATING TO ORGANIC CHEMISTRY

THE names used to denote ring formulæ are given below for convenience of reference

Since the publication of the last volume I have been assisted in the work of reading and making abstracts of original memoirs by Drs T Cooksey, T A Lawson, Samuel Rideal, Messrs J Wilkie, G N Huntly, J T Norman, and D A Louis I have also been assisted by Mr Arthur G Green and Mr Cecil W Cunningham in the work of revising the proof sheets I have great pleasure in thanking these gentlemen for the energetic and efficient manner in which they have carried out their share of the work

H FORSTER MORLEY

Nomenclature of Ring Formulæ

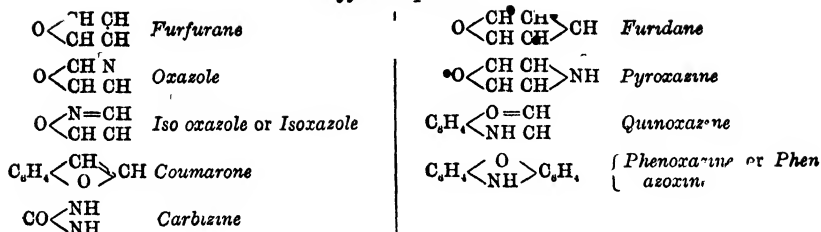
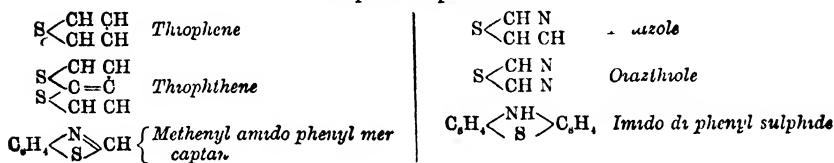
Hydrocarbons

$\text{CH} \begin{smallmatrix} \text{CH}_2 \\ \text{CH}_2 \end{smallmatrix}$ Trimethylene	$\text{C}_6\text{H}_4 \begin{smallmatrix} \text{CH} \\ \text{CH} \end{smallmatrix} \text{CH}$ Indonaphthene
$\text{CH} \begin{smallmatrix} \text{CH} \\ \text{CH} \end{smallmatrix} \text{CH}_2$ Tetramethylene	$\text{CH} \begin{smallmatrix} \text{CH} \\ \text{CH} \end{smallmatrix} \text{CH}$ Tetramethenyl
$\text{CH} \begin{smallmatrix} \text{CH} \text{CH}_2 \\ \text{CH} \text{CH}_2 \end{smallmatrix}$ Pentamethylene	$\text{CH} \begin{smallmatrix} \text{CH} \text{CH} \\ \text{CH} \text{CH} \end{smallmatrix}$ Pentamethenyl hydride

Nitrogen compounds

$\text{NH} \begin{smallmatrix} \text{CH} \text{CH} \\ \text{CH} \text{CH} \end{smallmatrix}$ Pyrrole	$\text{N} \begin{smallmatrix} \text{CH} \text{CH} \\ \text{CH} \text{CH} \end{smallmatrix} \text{CH}$ Puridine
$\text{NH} \begin{smallmatrix} \diagup \text{CH} \\ \diagdown \text{CH} \end{smallmatrix}$ Pyrrolole	$\text{N} \begin{smallmatrix} \diagup \text{CH} \\ \diagdown \text{CH} \end{smallmatrix} \text{CH}$ Puridazole
$\text{NH} \begin{smallmatrix} \text{CH} \text{N} \\ \text{CH} \text{CH} \end{smallmatrix}$ Glitoxaline	$\text{N} \begin{smallmatrix} \text{CH} \text{CH} \\ \text{CH} \text{CH} \end{smallmatrix} \text{N}$ Pyrazine
$\text{N} \begin{smallmatrix} \text{CH} \text{N} \\ \text{CH} \text{CH} \end{smallmatrix}$ Metapyrrole	$\text{N} \begin{smallmatrix} \text{CH} \text{N} \\ \text{CH} \text{CH} \end{smallmatrix} \text{CH}$ Pyrimidine
$\text{NH} \begin{smallmatrix} \text{CH} \text{N} \\ \text{CH} \text{N} \end{smallmatrix}$ or $\text{N} \begin{smallmatrix} \text{CH} \text{N} \\ \text{CH} \text{N} \end{smallmatrix}$ Triazole	$\text{N} \begin{smallmatrix} \text{CH} \text{N} \\ \text{CH} \text{N} \end{smallmatrix} \text{CH}$ Triazole
$\text{NH} \begin{smallmatrix} \diagup \text{CH} \\ \diagdown \text{CH} \end{smallmatrix}$ Osotriazole	$\text{N} \begin{smallmatrix} \diagup \text{N} \\ \diagdown \text{CH} \end{smallmatrix} \text{CH}$ Osotriazole
$\text{NH} \begin{smallmatrix} \text{N}=\text{N} \\ \text{CH} \text{N} \end{smallmatrix}$ Tetrazole	$\text{C}_6\text{H}_4 \begin{smallmatrix} \text{CH} \text{CH} \\ \text{N}=\text{CH} \end{smallmatrix}$ Quinoline
$\text{C}_6\text{H}_4 \begin{smallmatrix} \text{CH} \\ \text{NH} \end{smallmatrix} \text{CH}$ Indole	$\text{C}_6\text{H}_4 \begin{smallmatrix} \text{CH} \text{CH} \\ \text{CH} \text{N} \end{smallmatrix}$ Isquinoline
$\text{C}_6\text{H}_4 \begin{smallmatrix} \text{CH} \\ \text{N} \end{smallmatrix} \text{NH}$ Indazine	$\text{C}_6\text{H}_4 \begin{smallmatrix} \text{N} \text{CH} \\ \text{N} \text{CH} \end{smallmatrix}$ Quinoxaline
$\text{C}_6\text{H}_4 \begin{smallmatrix} \text{CH} \\ \text{NH} \end{smallmatrix} \text{N}$ Pseudo indazine	$\text{C}_6\text{H}_4 \begin{smallmatrix} \text{CH} \text{N} \\ \text{N}=\text{CH} \end{smallmatrix}$ Quinoxaline
$\text{C}_6\text{H}_4 \begin{smallmatrix} \text{CH} \\ \text{N} \end{smallmatrix} \text{C}_6\text{H}_4$ Acridine	$\text{C}_6\text{H}_4 \begin{smallmatrix} \text{CH} \text{CH} \\ \text{N}=\text{N} \end{smallmatrix}$ Cumuline
$\text{C}_6\text{H}_4 \begin{smallmatrix} \text{N} \\ \text{N} \end{smallmatrix} \text{C}_6\text{H}_4$ Phenazine	

INTRODUCTION.

Oxygen compounds*Sulphur compounds*

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Articles by Mr MUIR are initialed M M P M

UNSIGNED ARTICLES are by Dr MORLEY

ABBREVIATIONS.

I JOURNALS AND BOOKS

When an author has been mentioned in an article, he is usually referred to thereafter in that article by his initial only

A	Liebig's Annalen der Chemie
A A	Annales de la Sociedad Cientifica Argentina
A Ch	Annales de Chimie et de Physique
P Am A	Proceedings of the American Academy of Arts and Sciences.
Am	American Chemical Journal
Ann M	Annales des Mines
Am S	American Journal of Science
A C J	Journal of the American Chemical Society
Am Ch	American Chemist
Am J	American Journal of Pharmacy
Pharm	
An	The Analyst
A Ph S	Proceedings of the American Philosophical Society.
Ar N	Archives néerlandaises—The Hague
Acad	Mémoires de l'Académie des Sciences
Ar Ph	Archiv der Pharmacie
Ar Sc	Archives des Sciences phys et nat
B	Berichte der deutschen chemischen Gesellschaft.
B A	Reports of the British Association
Bt	Bulletin de la Société chimique de Paris
B B	Berliner Akademie Berichte
B C	Biedermann's Centralblatt für Agricultur Chemie
B J	Berzelius' Jahresberichte
B M	Berliner Monatsberichte
C S Mem	Memoirs of the Chemical Society of London
C J	Journal of the Chemical Society of London
C J Proc	Proceedings of the Chemical Society of London
C N	Chemical News
C R	Comptes rendus hebdomadaires des Séances de l'Académie des Sciences Paris
C C	Chemisches Central Blatt
D P J	Dingler's polytechnisches Journal
Fr	Fresenius Zeitschrift für analytische Chemie
G	Gazzetta chimica italiana
G A	Gilbert's Annalen der Physik und Chemie
H	Hoppe Seyler's Zeitschrift für physiologische Chemie
I	Proceedings of the Royal Irish Academy
J	Jahresbericht über die Fortschritte der Chemie und verwandter Theile anderer Wissenschaften
J C T	Jahresbericht für Chemische Technologie
J M	Jahrbuch für Mineralogie
J de Ph	Journal de Physique et des Sciences accessoires
J Ph	Journal de Pharmacie et de Chimie
J pr	Journal für praktische Chemie
J Th	Jahresbericht über Thierchemie
J R	Journal of the Russian Chemical Society
J Z	Jenaische Zeitschrift für Medizin und Naturwissenschaft
L V	Landwirthschaftliche Versuchs Stationen
M	Monatshefte für Chemie und verwandte Theile anderer Wissenschaften
M S	Le Moniteur Scientifique
Mém S	Mémoires de la Société d'Artuel.
d'A.	
Mém B	Mémoires couronnés par l'Académie de Bruxelles

ABBREVIATIONS.

N	Nature.
N Ed P. J.	New Edinburgh Philosophical Journal.
N J P	Neuer Jahresbericht der Pharmacie
N R P.	Neues Repertorium für die Pharmacie.
N J T	Neues Journal von Trommsdorff
P M	Philosophical Magazine
P	Poggendorff's Annalen der Physik und Chemie
P B	Beilatter zu den Annalen der Physik und Chemie.
Pf	Pfûger's Archiv für Physiologie
Pr E	Proceedings of the Royal Society of Edinburgh.
Ph	Pharmaceutical Journal and Transactions
Ph C	Pharmaceutisches Central Blatt
Pr	Proceedings of the Royal Society
P R I	Proceedings of the Royal Institution of Great Britain.
P Z	Pharmaceutische Zeitschrift für Russland
R T C	Recueil des travaux chimiques des Pays Bas
R P	Repertorium für die Pharmacie
Q J S	Quarterly Journal of Science
S	Schweigger's Journal der Chemie
Scher J	Scherer's Journal der Chemie
S C I	Journal of the Society of Chemical Industry
Sitz W	Sitzungsberichte der K Akademie zu Wien
T or Tr	Transactions of the Royal Society
T E	Transactions of the Royal Society of Edinburgh.
W	Wiedemann's Annalen der Physik und Chemie
W J	Wagner's Jahresbericht
Z	Zeitschrift für Chemie
Z B	Zeitschrift für Biologie
Z f d g	Zeitschrift für die gesammten Naturwissenschaften.
Natur- wiss	
Z K	Zeitschrift für Krystallographie und Mineralogie
Z P C	Zeitschrift für physikalische Chemie
Bn	Handbuch der organischen Chemie von F Beilstein, 2te Auflage
E P	English Patent.
G P	German Patent
Gm	Gmelin's Handbook of Chemistry—English Edition
Gm-K	Gmelin-Kraut Handbuch der anorganischen Chemie
Gerh	Traité de Chimie organique par Charles Gerhardt
K	Lehrbuch der organischen Chemie von Aug Kekulé
G O	Graham Otto Lehrbuch der anorganischen Chemie [5th Ed]
Stas,	Stas' Recherches, &c
Rech	} Aronstein's German translation is referred to as <i>Chem Proport</i>
Stas,	
Nouv R.	
Th	Thomsen's Thermochemische Untersuchungen

II TERMS AND QUANTITIES, &C, FREQUENTLY USED

Aq	Water, e.g. NaOHAq means an aqueous solution of caustic soda
aq	18 parts by weight of water
A'	} Residues of mono, di, and tri basic acids. Thus, in describing the salts of a monobasic acid NaA', CaA', AlA', may be written, HA' standing for the acid. For a dibasic acid we should write Na ₂ A'', CaA'', Al ₂ A'', &c
A''	
B' B'' &c	
	Stand for bases of the ammonia type, in describing their salts. Thus the hydrochloride would be B'HCl or B''2HCl, according as the base is monacid or diacid, &c
conc	Concentrated.
dil.	Dilute
g	gram
mgm.	milligram.
mm	millimetre.
mol	molecule
oil	liquid nearly, or quite, insoluble in water
pp	precipitate
to ppt	to precipitate.
ppg	precipitating
ppd.	precipitated.

ABBREVIATIONS.

xi

sol	soluble in.
insol	insoluble in
v e sol	very easily
v sol	very
m sol	moderately
sl. sol	slightly
v sl & l	very slightly
v	see
cf	compare
c	about
[°]	a melting point
(°)	a boiling point
H	Hardness (of minerals).
At w	Atomic weight
Mol w or	Molecular weight.
D	Density
cor	corrected
uncor	uncorrected
v	in vapour
V D	vapour density, i.e. density of a gas compared with hydrogen or air
S G	Specific gravity compared with water
S G $\frac{10}{15}$	" " at 10 compared with water at 0°
S G $\frac{15}{4}$	" " " 15° " " " 4°
S G $\frac{12}{1}$	" " " 12°, compared with water of which the temperature is not given
S H	Specific heat
S H v	" " of a gas at constant volume
S H p	" " " " " pressure
H C	Quantity of heat, in gram units, produced during the complete combustion of the mass of a solid or liquid body represented by its formula, taken in grams
H C v	Heat of combustion in gram units of a gram molecule of an element or compound, when gaseous, under constant volume
H C p	The same, under constant pressure
H F	Quantity of heat, in gram units, produced during the formation of the mass of a solid or liquid body represented by its formula, taken in grams, from the masses of its constituent elements expressed by their formulae, taken in grams
H F v	Heat of formation of a gram molecule of a gaseous compound from the gram molecules of its elements under constant volume
H F p	The same, under constant pressure
H V	Heat of vaporisation of a liquid, i.e. gram units of heat required to change a gram molecule of the liquid compound at B P into gas at same temperature and pressure
T C	Thermal conductivity (unit to be stated)
S V	Specific volume, or the molecular weight of a gaseous compound divided by the S G of the liquid compound at its boiling point compared with water at 4°
S.V.S	Specific volume of a solid, or the mass of the solid expressed by its formula, taken in grams, divided by its S G
E C	Electrical conductivity (the unit is stated in each case)
C E (10° to 20°)	Coefficient of expansion (between 10° and 20°)
S	Solubility in water
S (alcohol)	" " alcohol
μ , &c	Index of refraction for hydrogen line B
$R_{\frac{1}{2}}$	" " " sodium " D, &c
R_{∞}	Molecular refraction for sodium light, i.e. index of refraction for line D minus one, multiplied by molecular weight, and divided by S G at 15° compared with water at 0°
R_{∞}	The same, S.G. being determined at 15°-20° and referred to water at 4°
[a] _D	Cauchy's formula (Bruhl's R_{λ})
[a]	Specific rotation for sodium light
[a]	" " " neutral light $[a] = \frac{100}{p} \times \frac{a}{d}$ a = observed rotation for 100 mm of liquid. d = S.G. of liquid. p = no of grammes of active substance in 100 grammes of liquid

M M	Molecular magnetic rotatory power = $\frac{m \times \alpha}{d' \times m'}$, where m = molecular weight of the body of S G = d , α = angle of rotation under magnetic influence, α' = angle of rotation of water under same influence, and m' = molecular weight of water (18)
Ac . .	Acetyl C_2H_5O
Bz . .	Benzoyl C_6H_5O
Cy . .	Cyanogen CN
Et . .	Ethyl C_2H_5
Me . .	Methyl CH_3
Ph . .	Phenyl C_6H_5
Pr . .	Normal Propyl $CH_2CH_2CH_3$
Pr . .	Isopropyl $CH(CH_3)_2$
R, R' etc	Alcohol radicles or alkyls
prim	primary
sec	secondary
tert	tertiary
n	normal
m, o, p .	meta—ortho—para
c	consecutive
i	irregular
s	symmetrical
u	unsymmetrical,
ψ	pseudo
v	attached to nitrogen
α	Employed to denote that the substituent is attached to a carbon atom which is next, next but one, or next but two, respectively, to the terminal carbon atom. The end to be reckoned from is determined by the nature of the compound. Thus $CH_3CHBrCO_2H$ is α bromo propionic acid
β	
γ	
ω	denotes that the element or radicle which follows it is attached to a terminal carbon atom
α, β, γ , etc	indicate position in an open chain, only
1, 2, 3, etc	indicate position in a ring only
(α), (β), etc	Used when α, β , &c are employed in a sense different from the above, e.g. (α) dibromo camphor
(B)	Baeyer's Nomenclature
(Py)	benzene ring
	pyridine ring
	Thus (B 1 3) dichloroquinoline, means a meta dichloroquinoline in which the chlorine atoms are both in the benzene ring
	While (Py 1 3) dichloroquinoline, means a similar body, only the chlorine atoms are in the pyridine ring. The numbers are counted from two carbon atoms which are in different rings, but both united to the same carbon atom
(A)	denotes the central ring in the molecule of anthracene, acridines, and azines
exo	means that the element or radicle it precedes is in a closed ring
exo-	" " " " not in a benzene ring
allo	denotes isomerism that is not indicated by ordinary formulæ, thus maleic acid may be called <i>allo</i> -fumaric acid
thio	denotes displacement of oxygen by sulphur
sulpho	" the group SO_2H , except in the word sulphocyanide
sulphydro-	" the group SH
	Tribromonitrobenzene sulphonic acid {1 2 3 4 5} means that the three bromines occupy positions 1, 2, and 3, the nitro group the position 4, and the sulpho group the position 5

* Denotes that the formula to which it is affixed has not been determined by analysis. But it by no means follows that formulæ without this mark are those of analysed compounds.

All temperatures are given in degrees Centigrade unless when specially stated otherwise.

Wave lengths are given in 10^{-7} mm.

Formulæ, when used instead of names of substances, have a qualitative meaning only.

Thomsen's notation is used in thermochemical data.

DICTIONARY OF CHEMISTRY.

INDIGO CARBOXYLIC ACID *v* INDIGO

INDIGO-ARMINE *v* INDIGO

INDIGO-SULPHONIC ACID *v* INDIGO

INDIGOTINE *v* INDIGO

INDIGO-WHITE *v* INDIGO

INDILEUCINE *v* INDIGO

INDIN $C_{16}H_{16}N_4O_2$ Formed by the action of potash upon isatyde, thio isatyde, or di thio isatyde, formed also by heating isatin (Laurent, *A Ch* [3] 3, 471) Obtained also by boiling a solution of dioxindole in glycerin for a long time (Knop, *Z* 1865, 273) Deep rose coloured powder or minute needles Insol water, *v* sl sol alcohol and ether Dissolves in H_2SO_4 , forming a red solution, whence it is ppd unchanged by water A solution in concentrated alcoholic KOH deposits black crystals, apparently either $C_{16}H_{16}KN_4O_2$ or, more probably, potassium indate $C_{16}H_{16}KN_4O_2$.

Di bromo-indin $C_{16}H_{16}Br_2N_4O$ Formed by the action of bromine on indin or on di thio isatyde (Laurent) Formed also by heating di bromo isatyde at 220° (Erdmann *J pr* 22, 265) Violet black powder, sl sol alcohol

Di chloro-indin $C_{16}H_{16}Cl_2N_4O_2$ Obtained by heating di chloro isatyde either alone or with alcoholic potash (E) Dirty violet powder, insol water, alcohol, and $HClAq$ Forms a yellowish solution in $KOHAq$ from which HCl ppts yellow flakes

Tetra chloro indin $C_{16}H_4Cl_4N_4O$ Formed by heating di chloro isatyde either with alcoholic potash or alone below 200° (E) Dirty violet powder

Di nitro-indin $C_{16}H_4(NO_2)_2N_4O_2$ Formed by boiling indin or hydrindin with HNO_3 (L) Bright violet powder, insol water, *v* sl sol alcohol and ether Forms a dark brown solution in $KOHAq$

Indin di-sulphonic acid $C_{16}H_4(SO_3H)_2N_4O_2$? Formed by oxidising hydrindin di-sulphonic acid with INO_3 , with K_2FeCy_6 , with $NaOCl$, or even by exposing it in alkaline solution to the air (G a A Schlieper, *A* 120, 24) Red deliquescent crystals, *v* sol water, sl sol alcohol, insol ether Its solution dyes silk and wool scarlet Ammonium sulphide reduces it to hydrindin di sulphonic acid It forms a purple solution in $KOHAq$, and on warming the solution it becomes pale red, probably through assimilation of water On adding HCl to the pale red solution a yellow pp is formed ($C_{16}H_{16}(SO_3H)_2N_4O_2$?), which, when heated, quickly changes to indin disulphonic acid Salts — K_2A 5aq lustrous red needles — Ag_2A bulky brown needles — BaA 2aq slender dark reddish-brown needles or crystals

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powder M sol water, insol $BaCl_2Aq$, alcohol and cold $HClAq$

Hydrindin $C_{12}H_{12}N_4O_2$? Formed by the action of alcoholic potash upon indin, isatyde, thio isatyde, or di thio isatyde (Laurent *A Ch* [3] 3, 475) White, or pale yellow powder or needles (from alcohol), insol water, sl sol boiling alcohol Boiling HNO_3 converts it into a violet powder Hydrindin dissolves in warm aqueous KOH, and on cooling there separate pale yellow needles of a salt $C_{12}H_{12}KN_4O_2$ 3aq, which is decomposed by washing with water, leaving hydrindin

Hydrindin disulphonic acid, so called, $C_{12}H_8(SO_3H)_2N_4O_2$? Formed by the action of ammonium sulphide on indin sulphonic acid or on isatin sulphonic acid (G a A Schlieper, *A* 120, 20) Colourless radio crystalline mass, which becomes reddish when exposed to air *v* sol water, m sol alcohol, insol ether In alkaline solution it is oxidised by air to indin sulphonic acid — BaA 4aq white scales, *v* sol water, *v* sl sol $BaCl_2Aq$

Leucindin disulphonic acid $C_{16}H_{16}(SO_3H)_2N_4O_2$ Formed by boiling the preceding with baryta water (G a A Schlieper, *A* 120, 33) White crystalline mass, *v* sol water, sl sol alcohol Not reddened by the action of air on its alkaline solutions On evaporating with $HClAq$ there is formed indin disulphonic acid — BaA 5aq colourless crystals Its solution is not ppd by $AgNO_3$, till NH_4Aq is added

INDIPURPURIN is identical with *Indirubin*

v INDIGO

INDIRETIN *v* INDIGO

INDIRUBIN *v* INDIGO

INDIUM In At w 1134 Mol w unknown, as V D of element has not yet been determined [176°] (Winkler, *J pr* 102, 273)

S G $^{20^\circ}$ 7.11 to 7.147 (Reich a Richter, *J pr* 93, 480), $^{150^\circ}$ 7.362, $^{160^\circ}$ 7.421 (Winkler, *J pr* 95, 414, 102, 273) S H (0° to 100°) 0.5695 (Bunsen, *P* 141, 1) C E (0° to 100°) 0.000459 (Fizau, *C R* 68, 1125) Characteristic lines in emission spectrum 4510.2, 4101.3, 4071.6, 4032.7, 3852.8, 3834.7, 3257.8, 3255.5, 3038.7, 3008, 2982.3, 2940.8, 2889.8, 2559.5, 2527.1, 2351.3, 2306.9 (Hartley, *T* 1884, 102)

The observation of two indigo blue lines in the spark spectrum of a specimen of the *zinc-blende* of Freiberg in 1863, led Reich a Richter to the discovery of indium (*J pr* 89, 441, 90, 175, 92, 480)

Occurrence —In some *zinc blendes* from Freiberg (c 1 p c), also from Durham (Flight, *B* 10,

B

2054), and from Bohemia (Kachler, *J pr* 96, 447), in some Italian *galenas* (Denegri, *B* 11, 1249), in various zinc ores (Tanner, *J* 1874 1227), in the fumes from zinc-ovens (Böttger, *J pr* 98, 26), in some tungsten ores (Hoppe-Seyler, *A* 140, 247)

Preparation—Indium is more readily prepared from zinc which has been made from indium containing blends than from zinc-blende itself. The zinc is treated, for some days at the ordinary temperature, or for a shorter time at boiling temperature, with so much dilute H_2SO_4 or HClAq that a small quantity remains undissolved, the residue contains In, with some Zn, and Pb, Cu, Cd, As, and Fe. There are many methods for obtaining In from this residue, that of Bayer (*A* 168, 372) is simple, and yields very pure In. The residue is washed, treated with a few drops of dilute H_2SO_4 (to remove any basic Zn salts), again washed thoroughly with hot water, dissolved in HNO_3 (any SnO_2 which may be present is allowed to remain), evaporated with excess of H_2SO_4 until all HNO_3 is removed, and treated with water, the solution now contains sulphates of In and the other metals which may be present, but the greater part of the PbSO_4 remains insoluble. Large excess of NH_4Aq is added, whereby hydroxides of In and Fe are pptd with small quantities of hydroxides of Zn, Cd, Pb, and Cu, the ppt is well washed and dissolved in the smallest possible quantity of HClAq , the solution is boiled after addition of NaHSO_4 until the smell of SO_2 is almost gone, a basic sulphite of In, $2\text{In}_2\text{O}_3 \cdot 3\text{SO}_2$, is thus pptd as a fine crystalline powder. The ppt is free from salts of Cu, Zn, and Cd. If much Fe should have been present in the original Zn, small quantities of Fe salts may be pptd by the action of the air during filtration, in this case pptn should be conducted in CO_2 , or the ppt should be dissolved in NaHSO_4 and repptd by boiling. The ppt may contain Pb salts and traces of alkali, it is dissolved in SO_2 in which Pb sulphite is insoluble, after filtration In sulphite is pptd, free from Na salts, by boiling (cf Winkler, *J pr* 102, 278, Böttger, *J pr* 98, 26, R E Meyer, *A* 150, 137. For methods of preparing In from zinc-blendes v Reich a Richter, *J pr* 89, 441, 90, 175, 93, 480, Weselsky, *J pr* 94, 443, Richter, *J pr* 94, 414, Stolba, *D P J* 198, 223). The pptd basic In sulphite may be dissolved in H_2SO_4 after boiling off SO_2 , addition of NH_4Aq ppts $\text{In}_2\text{O}_3 \cdot \text{H}_2\text{O}$, which when strongly heated yields In_2O_3 . In is obtained from the oxide (1) by heating in a stream of pure H, (2) by mixing with pure C, and heating to a very high temperature, (3) by heating with an equal weight of Na cut in small slices, under a layer of fused NaCl in a porcelain crucible placed in a larger Hessian crucible, decomposing the alloy of Na and In so formed by water, and melting with Na_2CO_3 (Winkler, *J pr* 102, 275). Böttger (*J pr* 107, 89) recommends to ppt In from solutions in H_2SO_4 by placing a stick of pure Zn in the liquid, he washes the pptd metal with water, presses it with the finger, then between paper, and when quite dry fuses it under dry KCN.

Properties—A silver white, lustrous, ductile metal; softer than lead, leaves a mark when

rubbed on paper. Non-crystalline. Electro-negative \sim Zn and Cd; much less volatile than these metals. Unchanged in air at ordinary temperature, but burns to In_2O_3 with blue-violet flame and brownish fumes, when strongly heated in air. Combines directly with Cl, Br, I, and S when heated. Boiling water is not decomposed by In. Soluble in dilute acids with evolution of H and formation of salts In_2X_2 , where $\text{X} = \text{SO}_4, 2\text{NO}_3$, &c.

The at w of In has been determined (1) by synthesis of In_2O_3 from an (Ruch a Richter, *J pr* 92, 484, Winkler, *J pr* 94, 8, 102, 282, Bunsen, *P* 141, 28), (2) by analyses of In_2S_3 (R a R, *lc*) (3) by decomposing NaAuCl_4 by In, and determining the Au (Winkler, *J pr* 102, 212), (4) by determining the VD of InCl_3 , InCl , and InCl (Nilson a Pettersson, *C J* 53, 814), (5) by determining the SH of In (Bunsen, *P* 141, 1).

The atom of In appears to be monovalent (in InCl), divalent (in InCl_2), and trivalent (in InCl_3), in gaseous molecules, as the lower chlorides are decomposed by water with formation of In and InCl , it is probable that in solutions of its haloid compounds the atom of In is directly combined with at least three monovalent atoms.

In is distinctly metallic, with acids it evolves H and forms salts. A few basic, and some double, salts are known. In forms an ammonia alum $\text{InO}_3 \cdot \text{H}_2\text{O}$, reacts towards acids as a salt forming hydroxide. In is closely related to Al and Ga, less closely to Tl, it is also related to the other earth metals Sc, Yt, La, and Yb (v EARTH, METALS OF THE, vol 11 p 424). The investigation of In compounds is as yet far from complete.

Detection and Estimation—In salts colour the flame blue violet. Hoppe-Seyler (*A* 140, 247) boils c 1 gram of an In ore with *aqua regia*, neutralises by soda, filters, adds Na acetate, and ppts by H_2S , he dissolves the ppt in acid, and repts, the In_2S_3 is then tested in the flame after moistening with HClAq . The spectral lines 4510.2 and 4101.3 are very characteristic. Boiling with NaHSO_4 causes pptn of r, fine crystalline ppt $2\text{In}_2\text{O}_3 \cdot 3\text{SO}_2 \cdot 8\text{H}_2\text{O}$, this salt is used for the estimation of In.

Indium bromide InBr . White crystalline tablets, v sol water, formed by heating In in a stream of CO , charged with Br, and subliming (R Meyer, *A* 150, 429). VD not determined.

Indium chlorides. In combines with Cl in three proportions, forming InCl , InCl_2 , and InCl_3 .

INDIUM MONOCHLORIDE InCl . Mol w 148.77. VD at c $1100^\circ\text{--}1400^\circ = 78.16$ (Nilson a Pettersson, *C J* 53, 821). Obtained as a reddish black, vitreous, radiated, crystalline mass, by distilling InCl_3 (q v) on to In (rather more than the calculated quantity), heating for a little in a sealed tube, and distilling in a stream of CO , (N a P, *lc*). When melted InCl forms a dark-red liquid, in thick layers appearing almost black. Deliquescent, gradually decomposes in moist air, and quickly in water, to In and InCl_2 .

INDIUM DICHLORIDE InCl_2 . Mol w 184.14. VD at $1000^\circ\text{--}1400^\circ = 99.62$ (Nilson a Pettersson, *C J* 53, 820). White radiated crystals, obtained by heating In to its melting point in a current of dry HCl free from air until an amber-

coloured liquid is obtained, and removing any adhering HCl by heating in dry at free CO_2 . Unchanged in dry air, but deliquesces in ordinary air. Decomposed by water to form $\text{InCl}_3 \cdot \text{Aq}$ (N & P, *lc*)

INDIUM TRICHLORIDE InCl_3 . Mol wt 219.51 V D at c $8.7^\circ = 106.9$ (Nilson & Pettersson, *C J* 53, 818), at bright red heat V D = 113.88 (V & C Meyer, *B* 2, 611). InCl_3 does not appreciably volatilise at 140° , at c 600° volatilisation is slow and V D is rather higher than that calculated for InCl_3 , the normal V D is attained between 600° and 850° , at temperatures towards 1000° dissociation begins, probably into Cl and InCl , or InCl (N & P, *lc*). InCl_3 is prepared by heating In or a mixture of In_2O_3 with C, in a stream of dry Cl. Nilson & Pettersson heated molten In in dry air free HCl, and then gently heated the InCl_3 thus formed in air free, dry, Cl, and finally distilled in a current of dry, air free, CO_2 . White, lustrous tablets, deliquescent, sol in water with production of heat, the solution may be evaporated on a steam bath almost unchanged, but at higher temperatures decomposition occurs, probably with production of oxychlorides.

InCl_3 combines with KCl, NaCl, LaCl, and PtCl_4 to form crystalline double salts (R Meyer, *A* 150, 144, Nilson, *B* 9, 1059). The compound $2\text{InCl}_3 \cdot 6\text{KCl} \cdot 3\text{H}_2\text{O}$ separates in quadratic crystals from a solution of the mixed salts in proper proportions on evaporation.

Indium cyanide vol in p 332

Indium hydrosulphide The white pp produced by adding NH_4 sulphide, or H_2S , to solution of an In salt, after addition of tartaric acid and NH_4Aq , is probably a hydrosulphide, when dried, H_2S is evolved and In_2S_3 remains (R Meyer, *A* 150, 429).

Indium hydroxide v *Indium oxides and hydroxide*

Indium iodide InI_3 V D not determined. Yellow, crystalline, very hygroscopic, may be distilled in dry CO_2 , easily melted to a dark reddish brown liquid. Prepared by heating In in I vapour (R Meyer, *A* 150, 144, 429).

Indium oxides and hydroxide In forms two oxides, InO and In_2O_3 , the V D of neither has been determined, the existence of intermediate oxides is probable. In_2O_3 forms at least one hydrate $\text{In}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$. The oxides are basic, only salts corresponding to In_2O_3 have been isolated, although the lower oxide is said to dissolve in dilute acids without decomposition into In and In_2O_3 .

Indium monoxide InO Mol w unknown, as the oxide has not been volatilised. Obtained by heating In_2O_3 in H at c 300° until the oxide becomes almost black and water is no longer evolved. Forms a light, loose, powder, which quickly oxidises to yellow In_2O_3 , if brought into air before it is quite cold. Very pyrophoric. InO is changed by conc HNO_3 to In_2O_3 , with evolution of NO , dilute acids dissolve it slowly without apparent formation of In_2O_3 or In (Winkler, *J pr* 94, 1, 95, 414, 98, 344, 102, 273).

Indium sesquioxide In_2O_3 . Mol. w unknown as the oxide has not been volatilised. SG 7.179, SH $(0^\circ-100^\circ)$ 0.807 (Nilson & Pettersson, *B* 18, 1459). Obtained by ppg solution of an In salt by NH_4Aq , washing and heat-

ing the pp, or by heating In to full redness in air, also by strongly heating In carbonate or nitrate. A yellow powder, becomes brown on heating but yellow again when cold. (It is doubtful whether pure In_2O_3 is yellowish or white.) Very infusible, reduced to metal by heating with C, or in H, or with Na, reduction in H begins at c $190^\circ-200^\circ$, and at c 300° InO is formed (v *supra*). Soluble, in acids, quickly on heating, forming salts In_2X ($\text{X}=\text{SO}_4, 2\text{NO}_3$, &c).

Oxides intermediate between InO and In_2O_3 , possibly exist. By heating In_2O_3 in H to c 200° , a greyish blue body is obtained probably In_2O_3 , at c 230° a green substance, probably In_2O_3 , remains (Winkler, *lc*).

Indium hydroxide $\text{In}_2\text{O}_3 \cdot \text{H}_2\text{O}$ or $\text{In}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$ Formed by ppg a solution of an In salt by NH_4Aq , washing the pp and drying at 100° . The pp by NH_4Aq is gelatinous and resembles AlO_2H_3 , in air it dries to horny semitransparent lumps. Insol in NH_4Aq , easily sol in KOH or NaOH or NaOHAq , dissolves in acids to form salts In_2X ($\text{X}=\text{SO}_4, 2\text{NO}_3$, &c). A series of very unstable hydrates of In_2O_3 probably exists (cf Carnelley & Walker, *C J* 53, 88).

Indium oxybromide The white amorphous solid formed by heating In_2O_3 in Br vapour is probably an oxybromide, this substance is said not to be decomposed by heating with acids or alkalis (R Meyer, *A* 150, 137).

Indium oxychloride When $\text{InCl}_3 \cdot \text{Aq}$ is boiled down to dryness, the white residue is probably an oxychloride.

Indium salts Not many salts of In have been isolated and examined. They are obtained by dissolving In or In_2O_3 in acids, and evaporating, and in some cases by ppn from other In salts in solution. The chief salts are the carbonate, nitrate, sulphate, and a basic sulphite (v *Carbonates*, *Nitrates*, &c). The sulphate forms an ammonia alum, $\text{In}_2(\text{SO}_4)_3 \cdot (\text{NH}_4)_2\text{SO}_4 \cdot 24\text{H}_2\text{O}$, but with K_2SO_4 and Na_2SO_4 it forms double sulphates $\text{In}_2(\text{SO}_4)_3 \cdot \text{K}_2(\text{Na})\text{SO}_4 \cdot 8\text{H}_2\text{O}$.

Indium sulphide In_2S_3 Mol w unknown as compound has not been gasified. Obtained by ppg a neutral or feebly acid solution of an In salt by H_2S , and drying the pp, also by heating together In and S, or In_2O_3 and S. A yellowish-grey solid. If In_2O_3 is heated with S and Na_2CO_3 , and the fused mass is treated with water, In_2S_3 remains as lustrous tablets resembling mosaic gold (Winkler, *lc*). In_2S_3 is infusible, with acids it gives In salts and H_2S (Reich & Richter, *lc*). Heated in air it is burnt to In_2O_3 . According to Winsinger (*Bl* (2) 49, 452) In_2S_3 is obtained in aqueous solution in a colloidal form, by passing H_2S into In_2O_3 suspended in water, the dark yellow liquid thus obtained can be freed from H_2S by boiling, it is coagulated by acetic acid and also by salts. For reactions of alkali sulphides with In salts v R Meyer, *lc*.

Potassium, and sodium, indium sulphides, $\text{K}_2\text{In}_2\text{S}_3$, $\text{Na}_2\text{In}_2\text{S}_3$. Formed by fusing together 1 pt In_2O_3 , 6 pts S , and 6 pts K_2CO_3 or Na_2CO_3 . The K salt remains as hyacinth red quadratic tablets on lixiviating the fused mass with water, the Na salt goes into solution and separates on standing as $\text{Na}_2\text{In}_2\text{S}_3 \cdot \text{H}_2\text{O}$ which on drying gives $\text{Na}_2\text{In}_2\text{S}_3$ (R. Schneider, *J pr* [2] 9, 209).

M M P M

INDOGEN & INDIOXYL**INDOGENIC ACID & INDIOXYLIC ACID****INDOGENIDE OF BENZOYL ALDEHYDE**

$C_{11}H_{11}NO$ & $C_8H_7<\frac{CO}{NH}>CCH_2C_6H_5$ *Benzylidene-ψ-indoxyl* [176°] Obtained by heating indoxyllic acid with benzaldehyde (Baeyer, *B* 16, 2197) Long flat orange needles Sol alcohol and chloroform to yellowish red solutions, which have a yellowish green fluorescence In H_2SO_4 and strong HCl it dissolves with a deep red colour, in alcoholic KOH with a greenish blue, which gives the indigo spectrum

(α)-INDOGENIDE OF ψ ISATIN is INDIRUBIN

(β)-Indogenide of ψ isatin is INDIGO

Indogenide of ethyl ψ isatin is *Ethyl indirubin* v INDIGO

INDOGENIDE OF p-NITRO-BENZALDEHYDE $C_{13}H_{10}N_2O_3$ &

$C_8H_7<\frac{CO}{NH}>CCH_2C_6H_4(NO_2)$ *p Nitro-benzylidene-ψ-indoxyl* [273°] Formed by adding an acetic acid solution of *p* nitro benzaldehyde to an aqueous solution of indoxyl acidified with HCl Red needles (Baeyer, *B* 16, 2199)

INDOGENIDE OF PYRUVIC ACID $C_{11}H_8NO_4$

& $C_8H_7<\frac{CO}{NH}>CMeCO_2H$ [197°] Prepared by adding HCl to an aqueous solution of indoxyl and pyruvic acid (Baeyer, *B* 16, 2199) Red needles Easily soluble in alcohol and acetone Dissolves with a red colour in alkalis, with a blue colour in conc H_2SO_4

INDOGENE & INDIGO**INDOLE** C_8H_7N & $C_8H_7<\frac{CH}{NH}>CH$ *Ketole*

Mol w 117 [52°] (251°) (Ciamician & Zatti, *B* 22, 1980) VD 4.45 (calc 4.05) Occurs in small quantity in human excrement (Briegleb, *J pr* [2] 17, 133)

Formation — 1 By passing the vapour of oxindole over heated zinc dust (Baeyer, *A* 140, 295, *Suppl* 7, 56, Engler & Jancke, *B* 9, 1411) — 2 By distilling with zinc dust the yellow product of the action of tin and HCl on indigo (B) — 3 By fusing *o* nitro cinnamic acid with KOH and iron filings (Baeyer & Lummerling, *B* 2, 679, *Z* [2] 6, 213, Beilstein & Kuhlberg, *A* 163, 141) — 4 Formed to the extent of 5 p c in the passage through a red hot tube of diethyl-*o*-toluidine, in less quantity from di methyl-*o*-toluidine, in small quantity from methyl ethyl aniline, ethyl acetanilide, and di ethyl aniline, and in trace only from ethyl aniline when subjected to like treatment (Baeyer & Caro, *B* 10, 692, 1262) — 5 By digesting albumen with pancreas and water at 40° to 45° for several days (Nencki, *B* 8, 336, Kühne, *B* 8, 206) According to Harris and Tooth (*J Physiol* 9, 220) its formation is due to a special micro-organism — 6 By distilling albumen (1 pt) with KOH (8 pts), the yield being about 25 p c (Engler & Jancke, *B* 9, 1411, Nencki, *J pr* [2] 17, 98) — 7 By distilling (3,4,1) nitro propenylbenzoic acid with lime (Widmann, *B* 15, 2552) — 8 By boiling aniline with di chloro acetic aldehyde or with di chloro di ethyl oxide (Berlinerblau, *M* 8, 180) — 9 By heating *o* chloro-*o*-amido styrene with NaOEt at 165° (Lipp, *B* 17, 8067) — 10 Together with ethane, ethylene, and propylene, by the distillation of *o* cumidine

$C_8H_7(NH_2)(C_6H_5)$ over red hot PbO (Fileti, *G*, 13, 381) — 1 By the dry distillation of *o* tolyloxamic acid (Mauthner & Suida, *M* 7, 238) — 12 From phenyl-ethylene diamines by successive oxidation with CrO_3 and distillation over zinc dust (Prud'homme, *Bl* [2] 28, 55c) — 13 By fusing carbosyryl with potash (Morgan, *C N*, 36, 239) — 14 In small quantity, by heating the phenyl hydrazide of pyruvic acid with $ZnCl_2$ at 200° (E Fischer, *B* 19, 156°) — 15 By distilling calcium phenyl amido acetate with calcium formate (Mauthner & Suida, *M* 10, 253)

Preparation — 1 By heating aniline (50g) with an equal volume of water with inverted condenser and gradually adding di chloro ether (25g) After boiling for an hour the excess of aniline is distilled off, the residue heated for 5 hours at 220° (Berlinerblau, *M* 8, 18°) — 2 By distilling a mixture of calcium phenyl amido acetate with calcium formate, extract the distillate with ether, shaking the ether with dilute acid to remove aniline, evaporating and distilling with steam The indole is purified by conversion into the picrate The yield is 5 p c of the theoretical quantity (M & S)

Properties — Colourless laminae, m sol not water, v sol alcohol, ether, and ligroin Readily volatile with steam An alcoholic solution, acidified by HCl, dyes pine wood yellow Has a peculiar, but not very powerful, odour It is a very weak base, with conc $HClAq$ it forms a sparingly soluble salt, decomposed by boiling water

Reactions — 1 Suspended in water and subjected to the action of ozonised oxygen indole is converted into indigo and resinous products (Nencki, *B* 8, 727, 1517) Indigo appears in the urine after subcutaneous injection of indole — 2 Aqueous CrO_3 yields a bulky violet brown pp, insol ether, chloroform, and benzene, sl sol alcohol, forming a red solution sol conc $HClAq$ (L & J) — 3 Heated with MeI it gives di methyl quinoline dihydride (Ciamician & Zatti, *B* 22, 1980) — 4 When to an aqueous solution of indole there is added a few cc of $HClAq$, and then a large quantity of fuming HNO_3 (previously partially freed from nitrous fumes by evaporation), a red pp is formed which may be purified by solution in alcohol and precipitation by adding ether This substance, so called nitroso indole nitrate $C_{10}H_{13}(NO)_2HNO_3$, forms minute red needles, v sl sol water and ether, v sol alcohol nearly insol dilute nitric acid It detonates sharply when heated Alcoholic ammonium sulphide reduces nitroso indole nitrate to so called hydrazo indole $C_{10}H_{13}N_2$ (?), which forms yellow needles, sol alcohol and ether, melting at 140° to a deep-blue mass Acids and alkalis convert it into a dark brown colouring matter termed azo indole by Nencki

Picrate $C_{12}H_9N_3O_7$ Long red glittering needles

Acetyl derivative $C_{10}H_9Nac$ [183°] From indole and Ac_2O at 190° (Baeyer, *B* 12, 1314) Long needles (from water) or four sided pyramids (by sublimation)

(β)-Acetyl-indole $C_{10}H_7<\frac{C CO CH_3}{NH}>CH$ *Methyl indol ketone* [190° or] Obtained by heating

indole (a)-carboxylic acid, prepared from methylketole, with ten times its bulk of Ac_2O for 7 hours at 220° (Carlo Zatti, *Rend. Accad. Linc.* [4] 4, 184, B 22, 662) White needles Sublimes in colourless plates, sl sol cold, v sol warm water and benzene, may be extracted from its aqueous solution by ether Gives indole when heated with conc HClAq It readily yields an oxim and an oily phenyl hydrazide Potash fusion converts it into indole (β) carboxylic acid [214°] identical with that obtained by oxidising skatole

Picrate [183] Needles, sl sol cold benzene

Oxim $\text{C}_8\text{H}_7\text{C}(\text{NOH})\text{Me}$ [144° - 147°] White needles

Di-acetyl indole $\text{C}_8\text{H}_5\text{C}(\text{COCH}_3)_2$ [147° - 150°]

Prepared by heating indole (a)-carboxylic acid with Ac_2O and separated from the preceding by its greater solubility in water containing Na_2CO_3 and in boiling benzene (Zatti) Needles (by sublimation), sl sol boiling water When boiled with aqueous KOH or K_2CO_3 , it yields (β) acetyl indole

References - Di chloro indole, Benzyl indole, Methyl indole, Methyl ethyl indole, Ethyl indole

Isoindole is Di phenyl hydrazine (q v)

Di indole v Indoline under Indigo

Retin indole $\text{C}_8\text{H}_7\text{NO}$? Obtained by treating di chloro indole (chloro oxindole chloride) with HI dissolved in HOAc passing in SO and ppk with NaOH (Bayer, B 12, 1413) Amorphous, insol NaOHAq , v sol alcohol and ether Not volatile with steam Its solution in HOAc mixed with HCl colours pine wood red On dry distillation it yields indole

INDOLE (a) CARBOXYLIC ACID $\text{C}_8\text{H}_7\text{NO}_2$

$\text{C}_8\text{H}_5\text{C}(\text{COOH})_2$ [201°]

Formation -1 By heating the phenyl hydrazide of pyruvic ether with ZnCl_2 at 195° (E Fischer, B 19, 1567, A 236, 140) -2 By fusing (a) methyl indole (methyl ketole) (1 pt) with KOH (15 pts), the acid being prepd by H_2SO_4 from the dissolved product, the yield is over 50 p.c (Ciamician & Zatti, *Rend. Accad. Linc.* [4] 4, 746) -3 From acetyl (a) methyl indole $\text{C}_8\text{H}_5\text{C}(\text{NMe})\text{COCH}_3$ by potash fusion (Ciamician & Magnanini, B 21, 673)

Properties - Needles (from water) M sol hot water and benzene, v sol alcohol and ether May be sublimed in plates, but suffers partial decomposition thereby At 230° it splits up into indole and CO_2 Pine wood acidified with HCl is not coloured yellow by it. With isatin and H_2SO_4 it gives a reddish violet colouration It forms a pyruic acid compound crystallising in slender golden needles Its Ba salt is soluble Ac_2O at 220° forms (β) acetyl indole and di acetyl-indole

Anhydride $\text{C}_{10}\text{H}_7\text{N}_2\text{O}_3$

$\text{C}_8\text{H}_5\text{C}(\text{CH}_3\text{COON})_2$ [312° - 315°] Mel

w Confirmed by Raoult's method (Magnanini, B 23, 2508) Formed by boiling the acid with

Ac_2O using an inverted condenser. Yellow needles

Methyl ether MeA' [152°] Needles

Indole (B)-carboxylic acid

$\text{C}_8\text{H}_5\text{C}(\text{COOH})_2$ [214°] Formed by fusing (β) methyl indole (skatole) with KOH (Ciamician, Magnanini & Zatti, B 21, 673, 1929) Obtained also by fusing (β) acetyl indole with caustic potash (Zatti, B 22, 664) Plates (from water), decomposed on fusion Sl sol benzene and boiling water, m sol alcohol and ether, almost insol ligroin Its aqueous solution evolves CO_2 on boiling With isatin and H_2SO_4 it gives a brownish violet colour Its ammonium salt gives pps with CuSO_4 and with FeCl_3 . It does not yield a picric acid compound.

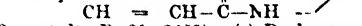
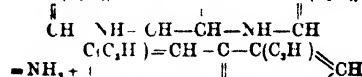
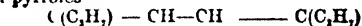
Indole di carboxylic acid

$\text{C}_{10}\text{H}_7\text{C}(\text{COOH})_2$ [above 250°] Obtained by saponifying its acid ether, which is formed when $\text{CO Et C}_8\text{H}_5\text{N}_2\text{H CMe CO Et}$, the product of the condensation of hydrazido benzene ether with pyruvic ether, is heated with zinc dust (Roder, A 236, 169) Slender needles, sol hot alcohol and acetic acid, sl sol ether and water Does not colour pine wood Decomposed on fusion giving off CO_2 and yielding a product that behaves like indole towards pine wood.

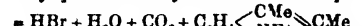
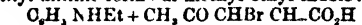
Mono ethyl ether EtHA' [250°] Yellow needles (from HOAc)

INDOLES Alkylated derivatives of indole may be formed synthetically in the following ways

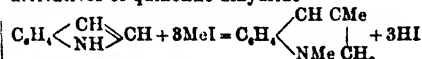
(a) By removal of NH_2 from the phenyl hydrazides of ketones, aldehydes, or ketonic acids Thus with the phenyl hydrazide of acetone $\text{C}_8\text{H}_5\text{N}_2\text{H CMe} = \text{NH}_2 + \text{C}_8\text{H}_5\text{C}(\text{CH}_3)_2$, and in like manner from the phenyl methyl hydrazide of pyruvic acid $\text{C}_8\text{H}_5\text{N}_2\text{H Me CMe CO}_2\text{H} = \text{NH}_2 + \text{C}_8\text{H}_5\text{C}(\text{CH}_3)_2\text{CO}_2\text{H}$ (E Fischer, A 236, 116) (b) By elimination of NH_2 from alkylated di pyrroles -



(Dennstedt, B 21, 3429) (c) By heating (β) bromo β acetyl propionic acid with aromatic amines (Wolff, B 21, 3360) Thus o- or p-toluidine gives tri methyl indole, (β) naphthylamine gives di methyl (β) naphthindole, while ethyl aniline forms di methyl ethyl-indole.



Indoles may be converted by alkyl iodides into derivatives of quinoline dihydride



INDOLINE v Indigo

INDONAPHTHENE

The hypothetical hydrocarbon C_{10}H_7 or $\text{C}_8\text{H}_5\text{C}(\text{CH}_3)_2$ which stands in the same relation to indole that naphthalene stands to quinoline (Baeyer & Perkin, jun, B 17, 122) The name *indene* has also been proposed for

thesame hydrocarbon (Roser, *A* 247, 132), v also METHYL INDONAPHTHENE Besides the methods mentioned below, indonaphthene derivatives may be formed by condensation of naphthalene de

riative Thus $C_6H_4 \begin{smallmatrix} \diagup CO \\ \diagdown CHCl \end{smallmatrix} \begin{smallmatrix} CO \\ \diagdown CCl_2 \end{smallmatrix}$, a product of

the chlorination of (β)-naphthoquinone is converted by cold aqueous NaOH into

$C_6H_4 \begin{smallmatrix} \diagup C(OH)CO_2H \\ \diagdown CHCl \end{smallmatrix} \begin{smallmatrix} CO_2H \\ \diagdown CCl_2 \end{smallmatrix}$ (Zincke, *B* 20, 2894) which

may be oxidised by chromic acid to

$C_6H_4 \begin{smallmatrix} \diagup CO \\ \diagdown CHCl \end{smallmatrix} \begin{smallmatrix} CO \\ \diagdown CCl_2 \end{smallmatrix}$ [59°]

INDONAPHTHENE DIHYDRIDE CARBOXYLIC ACID $C_{16}H_{16}O_4$, ϵ

$C_6H_4 \begin{smallmatrix} \diagup CH_2 \\ \diagdown CH_2 \end{smallmatrix} CH CO_2H$ *Hydrindonaphthene carboxylic acid* [130°] *S* 833 at 100°

Formation—1 By heating the corresponding di carboxylic acid until the evolution of CO_2 has ceased, and distilling the product (Baeyer a Perkin, *jun*, *B* 17, 122)—2 By the action of (1 mol of) ω -di bromo xylene (o xylenebromide) upon aceto acetic ether (1 mol) and NaOEt (2 mols) and saponification of the product with alcoholic KOH (Scherks, *B* 18, 378)

Properties—Colourless needles (from water), may be distilled when quickly heated. Sl sol cold water. On oxidation with $KMnO_4$, it yields phenyl-glyoxylic acid. It forms a bulky white silver salt, sl sol hot water.

Indonaphthene dihydride dicarboxylic acid

$C_6H_4 \begin{smallmatrix} \diagup CH_2 \\ \diagdown CH_2 \end{smallmatrix} C(CO_2H)_2$, [199°] Formed by the action of di ω bromo o xylene on malonic ether (1 mol) and NaOEt (2 mols) in ether (Baeyer a Perkin, *jun*, *B* 17, 122, Perkin, *jun*, *C* 1 53, 7), the resulting ether being saponified. Trimetric plates, m sol hot water, alcohol, and ether, sl sol cold water. When heated above 200° it gives off CO_2 and leaves the mono carboxylic acid— Ag_2A'' white amorphous pp becoming crystalline, sl sol water.

INDONAPHTHOQUINONE $C_{16}H_8O_2$, ϵ

$C_6H_4 \begin{smallmatrix} \diagup CO \\ \diagdown CO \end{smallmatrix} CH_2$, [131°] *Di keto indonaphthene* *Di keto hydrindene* Formed by warming its carboxylic ether $C_6H_4 \begin{smallmatrix} \diagup CO \\ \diagdown CO \end{smallmatrix} CH CO_2Et$ with alkalis (Wislicenus, *A* 246, 351). Best obtained by acidifying an aqueous solution of the sodium derivative $C_6H_4 \begin{smallmatrix} \diagup CO \\ \diagdown CO \end{smallmatrix} CNa CO_2Et$, extracting with ether, and leaving the ethereal solution to stand, when CO_2 is given off.

Properties—Small needles (from benzene ligroin), v sol hot alcohol and benzene, m sol ether and hot ligroin, v sl sol cold ligroin and water. Dissolves in alkalis with intense yellow colour.

Reactions—1 Gives a phenyl hydrazide

$C_6H_4 \begin{smallmatrix} \diagup C(N.HPh) \\ \diagdown CO \end{smallmatrix} CH_2$, [163°] crystallising in yellow needles, insol water and ligroin, v e sol ether and benzene. With phenyl hydrazide at 100° it gives rise to a di phenyl hydrazide

$C_6H_4 \begin{smallmatrix} \diagup C(N.HPh) \\ \diagdown C(N.HPh) \end{smallmatrix} CH_2$, which forms flesh-coloured plates [171°], and when dissolved in H_2SO_4 , is coloured deep bluish green by $FeCl_3$ (W

Wislicenus a Kötze, 252, 73)—2 Forms a nitroso erivative $C_6H_4 \begin{smallmatrix} \diagup CO \\ \diagdown CO \end{smallmatrix} CNOH$

[198°] crystallising from HOAc in triangular plates—3 *Benzoic aldehyde* at 120° forms $C_6H_4 \begin{smallmatrix} \diagup CO \\ \diagdown CO \end{smallmatrix} CCHPh$ [160°]—4 Gives with

bromine a compound $C_6H_4 \begin{smallmatrix} \diagup CO \\ \diagdown CC \end{smallmatrix} CBr_2$, [177°] crystallising in plates, m sol hot alcohol and HOAc, v sol ether and CS_2 (cf Zincke, *B* 20, 3216). This di bromo derivative, which is also formed by the action of bromine on $C_6H_4 \begin{smallmatrix} \diagup CC \\ \diagdown C(NH) \end{smallmatrix} CBr$ and on

$C_6H_4 \begin{smallmatrix} \diagup CO - C(NH_2) \\ \diagdown C(NH) \end{smallmatrix} CH$, is converted by the action of alkali into bromoform, phthalic acid, and the compound $C_6H_4 \begin{smallmatrix} \diagup C(OH) \\ \diagdown CO \end{smallmatrix} CBr$ [118°], whence

Br forms again the original $C_6H_4 \begin{smallmatrix} \diagup CO \\ \diagdown CO \end{smallmatrix} CBr_2$,

while Cl forms $C_6H_4 \begin{smallmatrix} \diagup CO \\ \diagdown CO \end{smallmatrix} CClBr$ [147°] (Zincke, *B* 21, 2394). The corresponding $C_6H_4 \begin{smallmatrix} \diagup CO \\ \diagdown CO \end{smallmatrix} CCl_2$ [125°] is formed by the action

of chlorine on $C_6H_4 \begin{smallmatrix} \diagup CO \\ \diagdown CCl_2 \end{smallmatrix} \begin{smallmatrix} CO \\ \diagdown CCl_2 \end{smallmatrix}$ dissolved in aqueous Na_2CO_3 .

Indonaphthoquinone carboxylic ether

$C_6H_4 \begin{smallmatrix} \diagup CO \\ \diagdown CO \end{smallmatrix} CH CO_2Et$ [78°] Formed by the action of NaOEt (2 mols) on phthalic ether (1 mol) and subsequent treatment with acetic ether, the product $C_6H_4 \begin{smallmatrix} \diagup CO \\ \diagdown CO \end{smallmatrix} CNa CO_2Et$ being treated with dilute H_2SO_4 (Wislicenus, *A* 246, 349). Slender yellow needles, insol water, v e sol alcohol, ether, benzene, and ligroin. $FeCl_3$ colours its alcoholic solution deep red. Dissolves in aqueous Na_2CO_3 , expelling CO . If its alkaline solution be boiled and then acidified CO_2 is evolved and indonaphthoquinone liberated. It forms a phenyl hydrazide. If its acid solution is boiled with water there is formed $C_{16}H_{16}O_4$, which dyes wool violet red and forms crystalline $C_{16}H_{16}BrO_4$ [196°] and $C_{16}H_{16}Br_2O_4$ [242°].

Salts— $C_{16}H_8O_2Na$ aq yellow needles (from water). Gives with MeI the compound $C_6H_4 \begin{smallmatrix} \diagup CO \\ \diagdown CO \end{smallmatrix} CMe CO_2Et$, crystallising in prisms [74°], v e sol alcohol and ether, insol water, sol NaOHAq with rose colour— $(C_{16}H_8O_2)_2Cu$ green crystals (from alcohol).

INDONE The ketone $C_6H_4 \begin{smallmatrix} \diagup CO \\ \diagdown CH \end{smallmatrix} CH$, a number of the haloid derivatives C^* which have been described by Roser (*A* 247, 132). Indone may also be viewed as the anhydride of di oxy INDONAPHTHENE.

INDOPHANE $C_{22}H_{14}N_2O_4$. A blue substance, resembling indigo, produced, together with naphthyl purpuric acid, by adding a hot concentrated solution of KCy (45 g) to 1 nitro naphthol (? g) dissolved in boiling water (2 litres) to which just enough NH_3 has been added to effect

solution (Hlasiwetz, Sommaruga, *C C* 1871, 617) Violet mass with green lustre, insol. water, alcohol, ether, benzene, and C_{12} , m sol H_2SO_4 , and hot $HOAc$, forming purple solutions. May be sublimed. Nitric acid converts it into a brownish red body, soluble in alkalis. Ferrous sulphate mixed with lime reduces it. Aqueous KOH forms $C_{12}H_7KN_2O_4$, a body greatly resembling indigo, NaOH acts in like manner.

INDOPHENAN ($C_{11}H_7NOS$)₂. Formed by shaking isatin (1 pt.) with H_2SO_4 and benzene that contains thiophene $C_4H_5NO_2 + C_6H_5S = H_2O + C_{12}H_7NOS$ (Baeyer, *B* 12, 1309, 16, 2188, 18, 2637, V Meyer, *B* 15, 2893, 16, 1465, P Meyer, *B* 16, 2269, Gumpert, *J pr* [2] 32, 278) Blue powder, which exhibits a coppery lustre when rubbed, or small needles (from alcohol phenol). Insol water, benzene, and ligroin, v sl sol alcohol, ether, and CS_2 , sl sol $HOAc$. In H_2SO_4 it forms a blue solution, whence it is pptd by water. Cannot be sublimed. May be reduced by zinc dust and $HOAc$ to a colourless body which is re oxidised to indophenin on exposure to air. Bromo-, di bromo-, and methyl indophenins may be formed in like manner by using bromo-, di bromo-, or methyl isatin in place of isatin in the above preparation.

INDOPHENOLS Quinonimides of the formula $C_6H_4 \begin{smallmatrix} \diagup NH \\ \diagdown NR' \end{smallmatrix}$, where R' is an aromatic radicle containing hydroxyl (*c/* INDIMINES). Thus when a mixture of di methyl *p* phenylene diamine and (a) naphthol is oxidised with NaOCl or $K_2Cr_2O_7$ there is formed the dark blue indophenol

$C_6H_4 \begin{smallmatrix} \diagup NMe_2Cl \\ \diagdown C_{10}H_7OH \end{smallmatrix}$ (Pabst, *Bl* [2] 38, 161),

while *p* amido phenyl piperidine mixed with phenol and oxidised by K_2FeC_4 forms a similar

body $C_6H_4 \begin{smallmatrix} \diagup N C_6H_5Cl \\ \diagdown N C_6H_5OH \end{smallmatrix}$ (Leilmann & Geller, *B* 21, 2288).

INDOANTHIC ETHER $C_{11}H_{11}NO$, *ie.* $C_6H_4 \begin{smallmatrix} \diagup CO \\ \diagdown NH \end{smallmatrix} > C(OH)CO_2Et$ [107] Prepared

by the careful oxidation of indoxylic ether with $1c.Cl_2$ (Baeyer, *B* 15, 775) Yellow needles or monoclinic prisms. Sol water and ether. On further oxidation it gives $CO_2H C_6H_4 NH CO CO_2Et$. On reduction it is reconverted into indoxylic ether. Decomposed by alkalis with formation of anthranilic acid.

Nitrosamine $C_6H_4(NO)NO$, [113°] Light yellow needles or tables. Sol alcohol and ether, sl sol water.

INDOXYL C_6H_7NO *ie.* $C_6H_4 \begin{smallmatrix} \diagup C(OH) \\ \diagdown NH \end{smallmatrix} > CH$ (stable form) or $C_6H_4 \begin{smallmatrix} \diagup CO \\ \diagdown NH \end{smallmatrix} > CH_2$ (transition form).

Formation—1 By warming potassium indoxyl sulphate with HCl or H_2SO_4 .—2 By heating indoxylic acid alone or with H_2SO_4 (Baeyer, *B* 14, 1744).—3 By boiling *o* nitro phenyl acetylene with acid ammonium sulphite, and treating the product with zinc dust and ammonia (Baeyer, *B* 15, 56).—4 By reducing isatogen-sulphurous acid with zinc dust and ammonia.

Properties—Oil. Not volatile with steam.

In the pseudo form $C_6H_4 \begin{smallmatrix} \diagup CO \\ \diagdown NH \end{smallmatrix} > CH_2$ it is not known in the free state, but di substitution products of ψ indoxyl are stable. Nevertheless in doxyl forms condensation products, called indogenides, with bodies containing a CO group, in which reactions it must be supposed first to change into the pseudo form.

Reactions—1 In alkaline solution it is rapidly oxidised in the air to indigo. $FeCl_3$ forms a white amorphous body, which is immediately converted by HCl into indigo (E. Baumann & Tiemann, *B* 13, 415).—2 Indoxyl is not affected by H_2SO_4 , or conc $HClAq$, but dilute $HClAq$ converts it into an amorphous red substance.—3 A solution in H_2SO_4 , treated with *o* nitro-phenyl propionic acid gives indoin.—4 On warming with a solution of Na_2CO_3 and *o* nitro-phenyl propionic acid, indigo is produced.—5 By the action of Na_2CO_3 upon an alcoholic solution of isatin and indoxyl there is formed indirubin, while with bromo isatin the product is bromo indirubin (Baeyer).—6 Bromine gives tri bromo aniline (E. Baumann & Tiemann, *B* 12, 1192).—7 In alkaline solution $K_2S_2O_8$ forms indoxyl sulphuric acid.—8 Diazo benzene chloride forms $C_6H_4 \begin{smallmatrix} \diagup C(OH) \\ \diagdown N_2Ph \end{smallmatrix} > CH$ [236°] (Baeyer, *B* 16, 2190).

Nitrosamine $C_6H_4 \begin{smallmatrix} \diagup C(OH) \\ \diagdown N(NO) \end{smallmatrix} > CH$ Formed

by the action of nitrous acid on indoxyl (Baeyer, *B* 16, 2188). Slender yellowish needles. When boiled with HCl it gives indigo.

Isonitroso derivative of ψ indoxyl

$C_6H_4 \begin{smallmatrix} \diagup CO \\ \diagdown NH \end{smallmatrix} > CNOH$ See oxim of ψ ISATIN.

Ethyl derivative $C_6H_4 \begin{smallmatrix} \diagup C(OEt) \\ \diagdown NH \end{smallmatrix} > CH$.

Formed by heating the ethyl derivative of indoxylic acid. Oil, volatile with steam. Smells like indole, and colours pine wood, acidified with HCl, brownish red. Its picric acid compound $C_{10}H_{11}NOC_6H_2(NO_2)_3OH$ forms brown needles. Its nitroso derivative

$C_6H_4 \begin{smallmatrix} \diagup C(OH) \\ \diagdown N(O) \end{smallmatrix} > CH$ forms yellow prisms [85°], sol alcohol, insol water, and converted into indigo by heating with HCl (Baeyer, *B* 15, 781).

Indoxyl sulphuric acid $C_6H_7NSO_4$. Occurs in the urine of herbivora and, in smaller quantity, in that of man (Schunck, *P M* [4] 14, 288, Hoppe Seyler, *C C* 1864, 511, H 8, 79, E. Baumann & Tiemann, *B* 13, 408). When indole is introduced into a dog by injection or in food large quantities of indoxyl sulphuric acid appear in the urine (Jaffé, *Fr* 11, 358, E. Baumann & Brieger, *H* 3, 254). Indoxyl sulphuric acid is also formed by adding $K_2S_2O_8$ to a solution of indoxyl in aqueous KOH (Baeyer, *B* 14, 1745).

Potassium salt—KA'. Plates (from hot alcohol). V sol water, v sl. sol cold alcohol. Not affected by KOH even at 170°. Decomposed by hot dilute HCl or H_2SO_4 , or even by water at 120°, into indoxyl and H^+SO_4 . When warmed with $FeCl_3$ and a little HCl it is entirely converted into indigo. Indigo is also formed when the salt is heated alone.

INDOXYLIC ACID C_8H_7NO , $\epsilon\epsilon$

$$C_8H_7\langle\begin{smallmatrix} C(OH) \\ NH \end{smallmatrix}\rangle \rightarrow CO_2H \text{ (stable form) or}$$

$$C_8H_7\langle\begin{smallmatrix} CO \\ NH \end{smallmatrix}\rangle \rightarrow CH \cdot CO_2H \text{ (transition form) } [123^\circ].$$

Formed by saponifying its ether with fused NaOH at 180° . White crystalline pp., sl sol water. In dilute alkaline solutions it is converted into indigo by the action of air or oxidising agents. On heating it splits off CO_2 , forming indoxyl.

Indoxyllic ether $C_8H_7N(OH)CO_2Et$. Colourless prisms, $[121^\circ]$, dissolves in alkalis and is reprecipitated by CO_2 . Prepared by reduction of isatogenic ether or *o*-nitro phenyl propionic ether. Heated with H_2SO_4 , it gives indigo sulphonic acid. On oxidation it gives successively indoxanthidic ether $C_{12}H_{11}N_2O_5$, indoxanthic ether $C_{11}H_{11}NO_4$, and hydrogen ethyl oxaloxyl *o*-amido benzoate $CO_2Et \cdot CO \cdot NH \cdot C_6H_4 \cdot CO_2H$.

Acetyl indoxyllic ether
 $C_8H_7N(OAc)CO_2Et$ white needles, $[138^\circ]$
Ethyl indoxyllic ether
 $C_8H_7N(OEt)CO_2Et$ large colourless crystals, $[98^\circ]$
Ethyl-indoxyllic acid
 $C_8H_7N(OEt)CO_2H$ plates, $[160^\circ]$. Oxidised to indigo by $FeCl_3$, but not in alkaline solution.
Nitroso-ethyl-indoxyllic acid
 $C_8H_7N(NO)(OEt)CO_2H$ flat golden needles. Decomposes at about 200° . Formed by the action of nitrous acid on ethyl indoxyllic acid (Baeyer, B 14, 1742, 15, 775, 16, 2189, Forrer, B 17, 376).

INDUCTION When H and Cl are mixed in equal volumes and exposed to sunlight a measurable time elapses before chemical change begins. Bunsen & Roscoe, who examined this phenomenon (*T* 1857 355), regarded the mixture as resisting chemical change, and they used the term *induction* to express the gradual overcoming of this resistance. The term has also been used by Wright, who noticed a similar phenomenon in the reduction of metallic oxides by CO and H (*C J Trans* 1879 475, 1880 757), v **CHEMICAL CHANGE**, vol 1 p 749.

M M P M

INDULINES (O N Witt & Thomas, *C J* 43, 112). Coloured compounds formed by the action of such bodies as azo, amido azo, nitro, or nitroso-compounds on the hydrochlorides of certain aromatic amines with elimination of NH_3 . Aniline hydrochloride gives a dark-blue colouration with amido azo benzene. Hofmann & Geyger (*B* 5, 474) supposed that only one dye stuff was formed $C_{12}H_{11}N_3 + C_6H_7N = C_{18}H_{13}N_3 + NH_3$. In reality a great number of dye-stuffs are got. If the mixture of diazo benzene anilide (1 pt), aniline (4 pts), and aniline hydrochloride (1 pt), which has been left until the diazo benzene anilide has changed into amido azo-benzene, be heated at 100° for 24 hours, crystals of azophenine $C_{18}H_{13}N_3$ separate. This forms garnet coloured leaflets (from aniline) $[237^\circ]$ (v **AZOPHENINE**). If the mixture of aniline, aniline hydrochloride, and amido azo-benzene, after heating at 100° till azophenine is formed, be further heated at 125° - 130° for 12 hours, the azophenine disappears, and the mass turns blue. If it be slowly cooled crystals are formed. These may be separated from the

mother-liquor (which contains 'induline B'), washed with spirit and with boiling water, and recrystallised from aniline mixed with aniline hydrochloride. So obtained, induline B forms glistening brown leaflets, $C_{18}H_{13}N_3 \cdot HCl$. It is insol cold, sol hot, spirit, and v sol aniline. Alcoholic NaOH liberates the free base, which forms purple solutions with alcohol and with aniline. Induline B is sky blue. Its sulphonic acid resembles that of induline B.

Induline B $C_{18}H_{13}N_3$ is formed by heating a mixture of amido azo benzene (100 pts), aniline hydrochloride (130 pts), and aniline (300 pts), for 5 hours at 110° , then adding more aniline hydrochloride (65 pts), and heating to 165° - 170° . The hydrochloride, $B \cdot HCl$, separates as green crystals. The free base separates from aniline as brilliant green needles. The solution of this base is of a dark bluish purple colour. The hydrochloride is insoluble in the usual solvents, and is hardly soluble in aniline. It dissolves in phenol and in H_2SO_4 , forming a greenish blue solution.

Induline B hydrochloride is v sol spirit, sl sol water, v sol a conc aqueous solution of aniline hydrochloride. The base forms a brown red spirit solution. The sulphonic acid is insol water, but its neutral alkaline salts form reddish blue aqueous solutions.

Formation of indulines is due to the dehydrogenating action of amido azo benzene, which itself becomes aniline and *p*-phenylene diamine, the latter then entering into the reaction. Azo benzene, phenyl amido azo benzene, chrysoidine, nitroso dimethyl aniline, and even amido azo naphthalene, may be substituted for amido azo benzene, but if instead of aniline *o*-toluidine or naphthylamine be used, red dyes are got (azo toline, &c). This would indicate that azophenine was an oxidation product of aniline, and not derived from the amido azo benzene. In the preparation of azophenine by heating aniline (4 pts) with aniline hydrochloride (1 pt) and amido azo benzene (2 pts) at 85° , ammonium chloride and *p*-phenylene diamine are constant by products, indicating that phenylation and reduction of the amido azo benzene has taken place (Witt, *B* 20, 1534). When azo phenine (400 g) is boiled with alcohol (40 litres) and strong H_2SO_4 (2 kilos) for 10 hours quinone dianilide (19½ g) is formed. The compound

$$C_6H_5\langle\begin{smallmatrix} NPh \\ NPh \end{smallmatrix}\rangle$$
 is converted by heating with aniline at 100° into azophenine (Haudrows's 1, *M* 9, 414). These reactions confirm the formula

$$C_6H_5\langle\begin{smallmatrix} NPh \\ | \\ NPh \end{smallmatrix}\rangle$$
 proposed by Kimich and by

Fischer & Hepp for azophenine, which is also consistent with its formation from nitroso di-

$$\text{phenylamine } C_6H_5\langle\begin{smallmatrix} N \cdot HPh \\ | \\ N \cdot O \end{smallmatrix}\rangle \quad (\text{O Fischer \& Hepp,}$$

B 20, 2479, 21, 676, 2617). By the action of aniline on azophenine, induline (6 B) of the formula $NPh \cdot C_6H_5\langle\begin{smallmatrix} NPh \\ | \\ N \end{smallmatrix}\rangle \cdot C_6H_5$ might be formed. A mixture of 3 B and 6 B induline is formed by heating *p*-nitroso di-phenyl amine with aniline and aniline hydrochloride in alcoholic solution at 130° . By heating *p*-nitroso-di-

phenyl amine (1 pt) with aniline hydrochloride (1 to 2 pts) and aniline (4 pts) at 75° – 140° , a very rich yield of induline is obtained, which chiefly consists of 6B. If pure azophenine is heated at 140° with equal parts of aniline and aniline hydrochloride 6B induline is almost exclusively formed (Fischer a Hepp, *B* 20, 2484). By the action of azo-, nitro-, or nitroso compounds on *m*- or *p*-diamines, indulines which are soluble in water are formed, e.g. *paraphenylene blue*.

Bosinduline $C_{22}H_{19}N_3$, etc

$C_{10}H_7(NPh)_2 \xrightarrow{NPh} C_6H_5$, [225°] Formed by the action of nitroso phenyl (a)-naphthylamine hydrochloride on aniline and aniline hydrochloride at 120° (O Fischer, *B* 21, 2621). Formed also by oxidising triphenyl tri amido-naphthalene $C_{10}H_5(NHPh)_3$, [124]. Red plates, almost insol water, m sol alcohol, v sol benzene. Reduced by HI and P at 150° to a yellowish red base and aniline. Zinc and HCl give naphthalene and aniline.

Salts $\rightarrow B \cdot HCl$ 14 aq reddish brown prisms $\rightarrow b \cdot H \cdot PtCl_5$, $\rightarrow b \cdot H_2SO_4$ aq $B \cdot HNO_3$, [231°] — Picrate plates, insol water.

INDYL PHENYL KETONE CARBOXYLIC

ACID $C_{10}H_7CO_2C_6H_5 \cdot \begin{smallmatrix} CH \\ NH \end{smallmatrix} > CCO_2H$ *Benzoyl indole carboxylic acid* [285°]. From the *iso* benzoyl phenyl hydrazide of pyruvic acid by heating with $ZnCl_2$ at 220° (Ruhemann a Blackman *C J* 55, 617). White needles, sl sol boiling water. Decomposes on fusion.

INOGEN v Muscle

INOSIC ACID $C_{10}H_{11}N_3O_{11}$? Occurs in the mother liquor in the preparation of creatine from flesh (Liebig, *A* 62, 317, *A Ch* [3, 33, 129]. Occurs to the extent of $\frac{1}{2}$ pc in the flesh of hens. Pigeon's flesh and human flesh contains none (Gregory, *A* 64 106, Schlossberger *A* 66, 82). It is found in some fish (Limpricht, *A* 113, 301).

Preparation — The filtrate from which creatin has separated is concentrated, then mixed with alcohol till it acquires a milky consistence, and left to itself for a few days. It then deposits a crystalline mixture containing the inosates of Na and Ba. The crystals are dissolved in hot water and $BaCl_2$ added. The Ba salt crystallises on cooling.

Properties — Uncrystallisable, v sol water, forming a solution with a flavour of broth, whence alcohol ppt it as an amorphous powder. Insol ether.

Salts — K_2A'' 7 aq long, four sided prisms, v sol water insol alcohol — Ba_2A'' 6 aq prisms *S* 25 at 16° .

INOSITE $C_6H_7O_7$, *Phaseomannite* [217°] (Maquenne), [225° cor] (Fick) (319° in vacuo) (M) SG (of $C_6H_7O_7 \cdot 2aq$) Δ 1524 SG (of $C_6H_7O_7$) Δ 1752 S (of $C_6H_7O_7 \cdot 2aq$) 10 at 12° (T a V), 13 at 19° (Fick) H C 666500 H F 311500 (Berthelot a Recoura, *C R* 105, 14). Discovered by Scherer (*A* 73 322, 81, 375) in muscular substance (*is, iods, muscle*). It occurs especially in the heart, but also in the lungs, kidneys, liver, and spleen of oxen (Cloetta, *A* 99, 289) in the brain of oxen (W Müller, *A* 103, 140), in human kidneys, and in the urine in a case of Bright's disease to the amount of 1 p c,

and sometimes in healthy urine (Kulz, *Fr* 16, 135). Found also in birds, in cephalopoda (Krukenberg, *J Th* 1881, 343), and in the flesh of a young porpoise (O Jacobsen, *A* 157, 227). Inosite occurs in green kidney beans (*Phaseolus vulgaris*) (Vohl, *A* 99, 125, 105, 330), in the green rods and unripe seeds of peas (*Pisum sativum*), in unripe lentils (*Ervum Lens*), in the unripe fruits of the common acacia (*Robinia Pseud-Acacia*), in the heads of the common cabbage (*Brassica oleracea*, var *capitata*), in foxglove leaves, in extract of digitalis, in the leaves of the dandelion (*Taraxacum dens leonis*), in potato shoots, in green asparagus and in its berries, in *Lactaria piperatus*, in *Clavaria crocea* (Marmé, *A* 129, 222), in the leaves of the common ash (*Fraxinus excelsior*) (Gintl, *J pr* 104, 491), in grape juice (Hilger, *A* 160, 333), in young vine leaves (Neubauer, *Fr* 12, 45), in walnut leaves (Tanret a Villiers, *A Ch* [5] 23, 389, *C R* 86 486, Maquenne, *C R* 104, 225).

Preparation — 1 Walnut leaves are extracted with water. The brown liquid is treated, at 100° , with excess of milk of lime, filtered, ppd with lead acetate, again filtered and treated with ammonia and lead acetate. The crude lead compound of inosite is treated with dilute sulphuric acid, and the inosite is ppd with alcohol and ether. It can be further purified by recrystallisation from hot dilute acetic acid (Hilger, Maquenne). — 2 The fresh-chopped tissue is covered with water, and allowed to stand for 24 hours in a cool place with frequent stirring, the liquid is then separated and the residue pressed, the solution is heated with a little acetic acid to separate albumin and hæmatin, then strained, evaporated on the water bath to one tenth, ppd with neutral acetate of lead and filtered, and the filtrate is mixed with basic acetate of lead, whereupon inosite is thrown down in combination with oxide of lead, accompanied by uric acid, cystine, and other substances. The pp, after washing, is decomposed under water by H S, and the liquid filtered from the sulphide of lead, it then sometimes deposits crystals of uric acid, and when evaporated to a small bulk on the water bath, and mixed with alcohol till a turbidity is produced, it yields crystals of inosite (Cloetta). — 3 Crystals of inosite may often be obtained by evaporating an aqueous liquid containing it, and adding three volumes of alcohol to one of the boiling liquid (Cooper Lane, *A* 117 118).

Properties — Monoclinic crystals (containing $2aq$) $a \cdot b \cdot c$ 1 090 1 1535, β = $111^{\circ} 39'$. Effloresces in dry air. At 100° it gives off its water of crystallisation. It has a sweet taste and is optically inactive. Sl sol cold dilute alcohol, insol absolute alcohol and ether. Does not combine with NaCl or KCl. With lead subacetate it forms a gelatinous pp which may be $C_6H_7O_7 \cdot 2PbO$. Inosite does not react with phenyl hydrazine (E Fischer, *B* 17, 582). Inosite is not turned brown by boiling alkalis, or by boiling dilute H_2SO_4 . It does not reduce Fehling's solution but forms a green pp. It will not undergo alcoholic fermentation. In presence of decaying cheese or of putrid meat and chalk it forms butyric and lactic acids. Sodium amalgam does not affect inosite. Inosite does not form an acid solution with borax.

Reactions—1 Evaporated with *nitric acid* it leaves a residue which gives with CaCl_2 in ammoniacal solution a rose red colouration (colour test)—2 When a solution of inosite is evaporated nearly to dryness and a drop of mercuric nitrate solution is added there is formed a yellow pp which turns red on warming (Gallior, *Fr* 4, 264)—3 Heated with oxalic acid it gives CO_2 and formic acid According to Lorin (*Bl* [2] 48, 235), this indicates that it is a polyhydric alcohol—4 HIAq at 170° gives a trace of benzene, phenol, and tri iodo phenol [153°]²—5 It forms no addition products with chlorine and bromine—6 It does not reduce boiling ammoniacal AgNO_3 , but on adding KOH a mirror is formed—7 *Chromic acid* oxidises it to CO_2 and formic acid in the cold—8 KMnO_4 gives CO_2 —9 Conc HNO_3 at 100° forms oxalic acid, but in an open dish on the water bath it gives on evaporation a white substance which dissolves in water, and the solution, mixed with alcohol and evaporated, then deposits black crystals of tetra oxy quinone HNO_3 also forms rhodozone acid (Maquenne, *Bl* [2] 47, 290) According to Maquenne this reaction, and *reaction 4* indicate that inosite is hexa oxy benzene hexahydrate

Hexa-acetyl derivative $\text{C}_6\text{H}_6(\text{OAc})_6$, [212°] (23° *in vacuo*) From inosite and AcCl or Ac_2O Sublimes at 200° Insol water, sol hot alcohol and Ac_2O Saponified by alkalis and strong acids (Maquenne, *C R* 104, 1719, *Bl* [2] 48, 58) According to Fick (*C C* 1887, 452), the acetyl derivative is $\text{C}_6\text{H}_6\text{Ac}_6\text{O}_6$ and melts at 216° cor

Hexa-benzoyl derivative $\text{C}_6\text{H}_6(\text{OBz})_6$, [258°] Minute needles Insol all usual solvents

Hexa nitroxyl derivative $\text{C}_6\text{H}_6(\text{ONO})_6$, 'Nitro inosite' [120°] Formed by adding a solution of inosite in fuming HNO_3 to conc H_2SO_4 Maybe crystallised from alcohol, a crystalline tri-nitroxyl derivative being left in the mother liquor Trimetric tables, insol water, v sol alcohol Detonates when struck Cold alcoholic KOH decomposes it forming KNO_3 , the inosite being completely broken up When boiled with alcohol and a little H_2SO_4 , the products are nitrous ether and inosite

INULIN ($\text{C}_6\text{H}_{10}\text{O}_5$)_x, or $\text{C}_{72}\text{H}_{124}\text{O}_{62}$, *Helennin* *Alantinn* *Menyanthin* *Dahlin* *Synanthrin* *Sinistrin* Mol w 1980 (by Raoult's method Brown & Morris, *C J* 55 464) [160°] SG 1.35 (Kiliani), 1.36 (Payen), 1.46 (Dubrunfaut), 1.47 (Dragendorff) [a_D^{20}] = -35° (Dubrunfaut, *C R* 42, 803), -36.5° (Lescœur & Morelle, *C R* 87, 216), -40° (Lefranc, *J Ph* [5] 2, 216) HC 684900 (Berthelot & Vieille, *Bl* [2] 47, 868, *A Ch* [6] 10, 459) HF 230600 (B & V), 197000 (Von Rechenberg) A starch like substance occurring in the roots of elecampane (*Inula Helennium*) (Valentin Rose, *A D* 1804), dandelion (*Taraxacum dens leonis*), chicory (*Cichorium Intybus*), feverfew (*Pyrethrum Parthenium*), and meadow saffron (*Colchicum autumnale*), in the tubers of the potato, the dahlias, and the Jerusalem artichoke (*Helianthus tuberosus*), in Lerp manna (from *Eucalyptus dumosa*), and in certain lichens (*Lichen fraxineus* and *Lichen fastigiatus*) Sinistrin, from the sea onion (*Urginea Scilla*), is perhaps identical with inulin

Preparation—The finely divided roots are boiled v h water in presence of a small quantity of CaCl_2 . The filtrate is cooled by a freezing mixture, and, after thawing the ice, filtered again The *residue* is redissolved in hot water, and the filtered solution again subjected to the action of a freezing mixture, the process being repeated until the inulin is perfectly white The inulin is then freed from traces of levulose by treatment with 97 p.c alcohol, and is finally dried over sulphuric acid It still, however, contains about 6 p.c proteid matter corresponding to 1 p.c nitrogen, and also minute traces of inorganic substances (Kiliani, *A* 205, 147)

Properties—White powder, resembling starch Tasteless Occurs also in a gum like or horny modification, probably as a hydrate Sol cold water, v sol hot water, insol alcohol The powder is composed of minute spheres (Sachsse) It melts at 160° , becoming changed to amorphous 'pyro inulin' (Prantl, *N R* 19, 513, 577, 641) Inulin is very hygroscopic, and even when dried at 100° its composition appears to be $(\text{C}_6\text{H}_{10}\text{O}_5)_x\text{H}_2\text{O}$ (Kiliani) The aqueous solution of inulin does not form a jelly like that of starch It is not coloured blue by iodine It dissolves in cold aqueous KOH and the solution, when acidified, deposits after a while unaltered inulin It dissolves in ammoniacal cupric oxide solution (Cramer, *J pr* 73, 16) the solution yielding after a few hours a copious blue pp (Schlossberger, *J pr* 73, 373) It dissolves in an ammoniacal solution of nickel oxide It does not ppt lead acetate or sub acetate, but with a solution of lead acetate containing ammonia it gives a white pp Inulin is not pptd by salts of Fe, Cu, Hg Ag, or Au Inulin forms with alkalis unstable compounds which are soluble in water and pptd there from by alcohol When carbonised it emits an odour of burnt sugar It reduces ammoniacal gold and silver solutions, but not Fehling's solution It is not fermentable, nor is its optio activity affected by diastase It is not converted into levulose by invertin A solution of inulin dissolves PbO Baryta water gives a pp soluble in excess of a solution of inulin, and not decomposed by CO_2

Reactions—1 When heated with water at 100° it is slowly but completely converted into levulose—2 Boiling dilute acids quickly convert it into levulose—3 Dilute *nitric acid* oxidises it to racemic, oxalic, glycollic, and formic acids, but forms neither saccharic nor mucic acids (Tollens, *A* 249, 220)—4 By long contact with bromine it yields bromoform, oxalic acid, and CO_2 —5 With bromine and silver oxide it yields glycollic acid (from the intermediate formation of levulose)—6 When heated with baryta water small quantities of lactic acid are formed—7 Sodium-amalgam has no action on inulin—8 With chloro sulphuric acid (ClSO_3H) at 0° it forms $\text{C}_6\text{H}_6(\text{SO}_3\text{H})_6(\text{OCl})(a_D^{20}) = +11.5^\circ$ whence warm water produces levulose (Clausson, *J pr* 128, 27)

Acetyl derivatives These are best formulated as derivatives of $\text{C}_{12}\text{H}_{22}\text{O}_{11}$.

Tri-acetyl derivative $\text{C}_{12}\text{H}_{17}\text{Ac}_3\text{O}_{11}$ Formed by boiling inulin (1 pt) with Ac_2O (1 pt) and HOAc (2 pts) for 15 minutes (Ferrouillat & Savigny, *Z* [2] 5, 509, Lescœur & Morelle,

OR 87, 216) Amorphous yellowish mass, v sol water and alcohol, in either Lavo-ratory
Tetra acetyl derivative $C_{12}H_{11}Ac_4O_{10}$
 From inulin by heating with Ac_2O (F & S, Schützenberger, *A Ch* [4] 21, 234) Amorphous Lavo-ratory

Hexa acetyl derivative $C_{12}H_{11}Ac_6O_{18}$
 Amorphous Lavo-ratory

Hepta acetyl derivative $C_{12}H_{11}Ac_7O_{21}$
 Dextro-rotatory

Metinulin When inulin is heated with glycerin alone, or when it is heated with dilute acids, the product contains bodies resembling dextrin. They are not very soluble in water and on heating yield soluble metinulin and inuloid which are not so powerfully lavo-ratory as the unaltered inulin. Optically inactive derivatives are also formed, but they are not identical with levulosan (Höning & Schubert, *M* 8, 529). Metinulin was prepared by Dragendoff by heating inulin (1 pt) with water (5 pts) in a closed vessel, and ppg with alcohol. Inuloid is a similar or identical substance, said by Popp (*A* 156, 199) to occur in the Jerusalem artichoke. It is twice as soluble in water as inulin.

INVERTIN : FERMENTATION, vol 11 p 541

IOD or **iodo** Use of this prefix applied to inorganic compounds, for Iodo compounds and Iodo salts v the element the iodo compound of which is sought for or the salts to the name of which iodo is prefixed. Thus *stichloride of lead* will be found under *Iodine* and *iodo chromate of potassium* under *CHROMATES*.

IODATES and PERIODATES Salts of iodic and periodic acids v **IODINE**, Or acids of, pp 21, 23

IODATION v Iodo compounds

IODHYDRIC ACID *HI* (*Hydroiodic acid*, *Hydrogen iodide*) Mol w 127.53 Solidifies at -55° (Faraday, 7 1845 170) V D 63.22 S H p (21° -100°, equal mass of water = 1) 0.55

(Strecker, *W* 17, 447) S H p (20°) 1397, S H v

(100°) 1396 (Strecker, *l.c.*) I C of HI_{Aq} v Kohlrausch, *P* 159, 233 S G of HI_{Aq} saturated at 0° - 199 (De Luynes *A Ch* [4] 2, 34), 20 (Vigier, *Bl* [2] 11, 125) Vapour pressure of liquid HI at -17.8° - 2.9 atmos, at 0 - 4 atmos, at 15.5° - 5.8 atmos (Faraday, *T* 1845 170, values approx only) H F $[HI, I]$ - 60.36 gaseous H and solid I to form gaseous HI, $[H, I]$ - -436 gaseous elements at 1.0° , $[H, I, Aq]$ - 13,170 (*Th* 2, 35) HI does not occur in nature

Formation - 1 By direct union of H and I, by passing a stream of H over I at a full red heat (Courtois, *A Ch* 88, 305, Merz & Holzmann, *B* 22, 869) Cornu-vander passes H and I vapour over Pt black heated to 300° - 400° (*A Ch* [3] 80, 242, 34, 77, v also Blundell, *P* 2, 216, Lemoine, *C R* 85, 34) - 2 HI is formed by decomposing a metallic iodide by another acid, but a portion of the HI is generally decomposed with separation of I, and in some cases the HI reacts with the other acid, e.g. when H_2SO_4 is added to KI, SO_2 and H_2S are produced as well as HI and I. To prepare a solution of HI in water, Glover adds an exactly equivalent quantity of H_2SO_4 to BaI_{Aq} , and separates from ppt $BaSO_4$, (*P M* [8] 19, 92) -

3 By the reaction between Na_2SO_4 (Mène, *C R* 28, 478), or $Na_2S_2O_3$ (Gladstone, *P M* [3] 35, 345), I, and H_2O (cf formation of HBr, vol 1 p 532) - 4 By decomposing PI_3 by water (Kolbe, *J pr* 15, 172, Vigier, *Bl* [2] 11, 125, Pettenkofer, *A* 138, 57) Bieckher (*C C* 1863 207) recommends to make PI_3 by adding P to I in CS_2 and distilling off the CS_2 - 5 Etard & Moissan (*B* 18, 1862) heat I with colophony

Preparation - 1 Washed H_2S is passed into water in which a little finely powdered I is suspended ($2HI_{Aq} + 2L_{Aq} = 4HIAq + 2S$), more I is added little by little, and the passage of H_2S is continued. When all the I has been added, and no brown colour is produced on shaking the liquid, the separated S is agglomerated by briskly agitating the vessel, the liquid is filtered, H_2S is removed by gently warming, and the liquid is distilled, $HIAq$ of SG c 1.7 distils at 120° - 128° . An inverted untubulated retort with wide neck is a suitable vessel, the H_2S is passed down the neck of the retort by a tube dipping into the water in which the I is suspended. The $HIAq$ thus prepared may be used for making gaseous HI or a more conc solution. About 2 pts I are dissolved in the $HIAq$ made as described, and the solution is dropped from a tap funnel on to amorphous P (in a flask) moistened with $HIAq$ of the same concentration, the HI produced is passed through a wide U tube containing glass beads and some amorphous P moistened with $HIAq$ (to convert any I vapour into HI) $[PI_3 + 3H_2O + Aq = H_3PO_4 + 3HI]$. If dry HI is wished for, the gas is passed through a $CaCl_2$ (or better CaI_2) tube and then over P_2O_5 , and collected by downward displacement. If $HIAq$ is to be prepared, the gas coming from the first U tube is passed into the tubulus of a retort, the neck of which dips a very little way beneath the surface of water kept cold by ice; this arrangement prevents the water from rushing back into the U tube. The flask containing P in $HIAq$, into which the solution of I is dropped, should be only very gently warmed for some time, as H_3PO_4 is among the first products of the reaction, and when this is heated H_3PO_4 and PH_3 are formed, the latter of which combines with HI to form PH_4I (v Bannow, *B* 6, 1498) - 2 L Meyer recommends the following method of preparing $HIAq$ by the formation and decomposition by water of PI_3 (*B* 20, 3381) 100 g I in a retort, with the neck inclined upwards, are moistened with c 10 g water, a tap funnel (or a funnel into the neck of which fits a glass rod) is fitted into the tubulus of the retort, 5 g amorphous P and 10 g water are placed in the funnel, and a single drop of this is allowed to flow into the retort, after a little a drop or two more is allowed to follow, and a little later the liquid is added in larger quantities. HI passes off and may be collected in water as described under 1. No heating is required. If more than one drop of the P in water is added at first an explosion usually follows. The I carried over is nearly all deposited on the neck of the retort. 3 14 pts KI are warmed with 20 pts I and 1 to 1½ pts P with a little water (Müllon, *J Ph* 28, 99, Roscoe, *C J* 13, 146) - 4 Na_2SO_3 (6 pts) is rubbed up with water (1 pt), and after warming, I (3 pts) is slowly added (Mène, *C R*.

28, 478, cf Gladstone, *P M* [3] 85, 845) — 5 60 g of copaiba oil are slightly warmed in a 500 c.c. retort connected with a reversed condenser, 20 g of I are added little by little, and the temperature is allowed to rise, after a few minutes a regular and steady stream of HI comes off, when this slackens the retort is allowed to cool somewhat, more I added, and heating is recommenced, about 145 g HI may be obtained from 150 g I (Bruylants, *B* 12, 2059)

S Kemp (*P M* [3] 7, 444) says that liquid HI may be prepared by placing H persulphide in one limb of a sealed tube and a little I in the other, and gently warming the persulphide, H_2S is evolved and liquefied, and the I dissolves in the liquid H_2S , on then adding a drop of water (this is done by a special arrangement of the tube) HI is produced and liquefied, no details are given as to how the HI is separated from the H_2S

Properties — HI is a colourless, strongly acid gas, it is incombustible and extinguishes flame. Dry HI is unchanged when kept in closed tubes in the dark (Lemoine, *J* 1877 138) HI is readily liquefied (*v ante*), liquid HI is colourless (Kemp, *P M* [3] 7, 444, says it is yellowish), at -55° it solidifies to a colourless, ice like mass (Faraday, *T* 1845 170) Liquid HI is a very bad conductor of electricity (Bleekrode, *W* 3, 161, Hittorf, *W* 4, 374) HI is very soluble in water, the solution contains a strong acid, affinity of HIAq is about the same as that of HClAq (*v AFFINITY*, vol 1 p 82) HIAq is a colourless, strongly acid, fuming liquid, the solution saturated at 0° has S.G. 1.99 to 2.0, this solution gives off much HI when warmed to 40° – 50° . When HIAq containing more than c 57 p.c HI is distilled, HI is evolved until the S.G. becomes c 1.67–1.70 when the B.P. becomes approximately constant at 127° (at 774 mm.), and the liquid contains 57.0 p.c HI, if the original liquid contains less than 57 p.c HI water distils over until the acid of 57 p.c is produced, which then distils at 127° almost unchanged (Roscoe, *C J* 13, 160) By passing dry H through HIAq at 15° – 19° , an acid of constant concentration, 60.3 to 60.7 p.c HI, is obtained, at 100° the constant acid contains 58.2 to 58.5 p.c HI (Roscoe, *lc*) Topsøe gives the following S.G. and composition of HIAq (*B* 3, 403, cf Wright, *C N* 23, 242) —

Temp	S.G. referred to H_2O at same temp.	P.c HI	Temp	S.G. referred to H_2O at same temp.	P.c HI
0			0		
13.5	1.017	2.29	13	1.413	40.43
"	1.052	7.02	"	1.451	43.39
"	1.077	10.15	"	1.487	45.71
13	1.095	12.21	"	1.528	48.22
13.5	1.102	13.09	13.5	1.542	49.13
"	1.126	15.73	13	1.573	50.75
"	1.164	19.97	12.5	1.603	52.43
13.8	1.191	22.63	14	1.630	53.93
"	1.225	25.86	13.7	1.674	56.15
13.5	1.254	28.41	13	1.696	57.28
"	1.274	30.20	12.5	1.703	57.42
13	1.309	33.07	13.7	1.706	57.64
"	1.347	36.07	12	1.708	57.74
"	1.382	38.68			

HI is absorbed by charcoal, according to Favre 22,000 gram thermal units are produced for every 128 grams HI absorbed (*A Ch* [5] 1, 209)

Reactions — 1 HIAq is decomposed by electrolysis to H_2 and H (Riche, *J* 1858 101) 2 HI is slowly decomposed in sunlight, the decomposition proceeds without limit (Lemoine, *J* 1877 139) When mixed with oxygen and exposed to sunlight, even when the gases are dry, decomposition occurs, and proceeds the more rapidly the greater the mass of O (dry HCl and HBr are not decomposed under the same conditions) (Richardson, *C J* 51, 801) HIAq is readily decomposed with separation of I by exposure to air — 3 HI is decomposed by heat to H and I, slowly at 180° quickly a 440° (Hautefeuille, *Bl* [2] 7, 198), the dissociation of HI has been exhaustively studied by Lemoine (*v DISSOCIATION*, vol 1 p 400) — 4 Heated with oxygen H_2O and I are formed — 5 Chlorine forms HCl and I, with excess of Cl, ICl_3 is produced, bromine reacts similarly — 6 Sulphur and selenium form HI_2 or H_2Se , and iodide of S or Se (Hautefeuille, *Bl* [2] 7, 198), in presence of water I reacts with H_2S to form HIAq and S (*v PREPARATION* No 1) — 7 HI without action on amorphous phosphorus at 100° , but with ordinary phosphorus, even at the ordinary temperature, it forms P_2I_4 , HIAq slowly reacts with excess of P to produce H_3PO_4 and PH_3I ($2P + HIAq + 2.1 O = PH_3I + H_3PO_4$) (Damoiseau, *J* 1880 272) — 8 Conc. nitric acid decomposes gaseous HI instantaneously — 9 Conc. sulphuric acid forms I, and also SO and H_2S — 10 Sulphur dioxide forms S, I, and H_2O — 11 Iodic acid reacts with HIAq to form I and H_2O — 12 Very many oxidizers separate I with simultaneous formation of H_2O , e.g. H_2O_2 , $HClO_4$, $IClO_3$, chromates — 13 Many metals form iodides and evolve H , with HIAq — 14 Metallic oxides and carbonates generally form iodides and H_2O , metallic peroxides form the same products and also separate I — 15 HI produces pps of iodides when added to solutions of salts of metals which form insoluble iodides, e.g. salts of Hg, Ag, Cu, Pd — 16 Potassium permanganate produces KIO_3 — 17 HI reacts with carbon compounds generally, especially with such as contain the group OH, as an energetic reducing agent (cf Berthelot, *Bl* [2] 9, 8, 91, 173, 265)

Combinations — 1 HIAq dissolves several metallic iodides, e.g. BiI_3 , HgI_2 , PtI_2 , and PtI_4 , some of the solutions thus formed react with alkalis and alkaline hydroxides to form salts, which are probably derived from acids containing I and the metal whose iodide was dissolved in HI, e.g. by dissolving AuI_3 in HIAq and adding KOH the salt $KAuI_4$ is obtained, by similar reactions the salts Na_2PtI_6 , $MgPtI_6$, &c., are formed. Such reactions render it probable that HI combines with many metallic iodides, and that the products frequently react as acids, one or two such acidic compounds have been isolated, e.g. H_2PtI_6 , $9H_2O$ (cf BROMHYDRIC ACID, Reaction 7, vol 1 p 583, and CHLORHYDRIC ACID, Reaction 13, vol 1 p 8) — 2 HI combines with ammonia to form NH_4I , $[NH_2^+HI^-] = 43,462$ (*Th* ~, 75) — 3 With phosphine HI combines to form PH_4I , $[PH_3^+HI^-] = 24,100$ (Ogier, *C R* 89, 705)

The solution of HI in water is attended with production of much heat $[HI, Aq] = 19,207$ (*Th* 2

84) Thomsen's measurements of the heat of solution of HIAq point to the existence of a hydrate $\text{HI} \cdot \text{H}_2\text{O}$, the results are similar to those obtained for HClAq , but cannot be represented by so simple a formula (cf *CHLORHYDRIC ACID*, vol ii p 8)

The heat produced by diluting HI in $n\text{H}_2\text{O}$ with quantities of water varying from 200 to 300 H_2O is given by Thomsen as follows (Th 3, 76) —

n	$[\text{HI} \cdot n\text{H}_2\text{O}, (300 - n)\text{H}_2\text{O}]$
2	6670
3	4400
5	1830
10	630
20	220
50	70
100	30

The acids HCl , HBr , and HI are very similar in their properties and reactions. All dissolve very freely in water, forming solutions of monobasic acids, the affinities of which are great and approximately equal. All combine with certain metallic haloid compounds, especially with those of Hg , Au , Pt , and Pd , to form compounds which are best regarded as distinct acids. The thermal reactions attending the syntheses of the three acids show a gradation. Thomsen (Th 2, 39) gives the following data —

$\text{X} (\text{H}, \text{X})$ gaseous, at 180° , from gaseous elements.
 X 22,153
 Br 12,413
 I — 436
 $\text{X} (\text{H}, \text{X}, \text{Aq})$
 Cl 39,315 gaseous Cl at 19°
 Br 32,197 (calculated on assumption that Br l 18,619) and I are gaseous at 19°

Of the three acids HI is the most easily decomposed by heat and by oxidisers (cf *HALOGEN ELEMENTS*, vol ii p 665, where HF is compared and contrasted with the other halogen acids H^+X^-)

M M P M

IODHYDRIN (GLYCERIN)

Di-iodhydrin: DI IODO PROPYL ALCOHOL

IODIC ACID: IODINE Oxyacids of p 19

IODIDES Binary compounds of I with more positive elements, i.e. with any element except O , F , Cl , or Br . Iodides of almost all metals and binary compounds of I with all non metals except B , have been isolated. Most metallic iodides may be obtained by direct combination of the elements, many are obtained by heating I with metallic oxides, or by dissolving metals or their oxides in HIAq . I reacts with alkalis and alkaline hydroxides to form iodides and iodates. Some non metallic iodides are formed by direct union of the elements, e.g. iodides of H , Br , Cl , Se , S , and P . NI_3 is obtained by the reaction between I and NH_3Aq , Cl , by the reaction of AlI_3 on a mixture of CCl_4 and CS_2 . O and I combine indirectly, oxides of I are obtained by the action of oxidisers on I . The non metallic iodides as a class are more easily decomposed by heat than the chlorides or bromides, oxide of I is much more stable than oxide of Cl , and no oxide of either Br or F has yet been isolated. If two chlorides or bromides of a specified metal are known, the more stable iodide of that metal as a general rule corresponds to the lower chloride or bromide, e.g. SbCl_3 and SbCl , exist, but only SbI_3 , FeCl_3 and FeCl are stable, but I ,

FeI , exists it very easily goes to FeI_2 and I , similarly with CuI , which exists only in solution, and very readily parts with I becoming CuI , whereas CuCl_2 is more stable than CuCl . In their reactions with water metallic iodides are usually less easily decomposed than the corresponding bromides and chlorides. Metallic iodides as a class dissolve in water without change, some, however, form oxyiodides, e.g. SbI_3 and BiI_3 , and some form oxides and HIAq , e.g. SnI_2 . Metallic iodides, generally speaking, are not so readily volatilised as chlorides or bromides, most of them are unchanged by heat, but some are decomposed to metal and I , e.g. iodides of Au , Pt , and Pd . As a whole, then, the metallic iodides are not so numerous or so varied in composition as the chlorides or bromides, and they are more stable towards heat and the action of water than the chlorides and bromides, they are also less easily reduced, e.g. by H or CO , than the chlorides or bromides.

The heat of formation of a metallic iodide is usually considerably less than that of the bromide or chloride of the same metal. The following data are taken from Thomsen, —

X	$[\text{Na}, \text{X}, \text{Aq}]$
Cl	193,020
Br	171,160
I	140,600
X	$[\text{Ca}, \text{X}^2, \text{Aq}]$
Cl	187,230
Br	165,360
I	134,940
X	$[\text{Al}^3, \text{X}^3, \text{Aq}]$
Cl	475,650
Br	410,040
I	318,780

The difference between the heat of formation in aqueous solution of a chloride and the analogous bromide of the same metal is approximately 21,850, and the difference in the case of a chloride and analogous iodide is approximately 22,250. Some metallic iodides exhibit differences in crystalline form and SG, for instance SbI_3 forms hexagonal crystals, and also exists in trimetric and in monoclinic forms, CdI_2 probably exists as a white salt SG 5.644 unchanged at 250° , and also as a brownish compound SG 4.626, which begins to change at 40° (vol i p 656), there are also differences between BiI_3 according as it is prepared by sublimation or by pptn.

Iodides are usually decomposed when heated in Cl or Br with production of chlorides or bromides and I , heated in HCl they generally form chlorides and HI . Heated with conc H_2SO_4 , or conc HNO_3 , I is separated, and SO_2 (also sometimes H_2S), or NO , is evolved. I is separated from iodides by the action of many oxidising agents, such as CrO_3 , ferric salts, MnO_2 , &c.

Many metallic iodides dissolve freely in solutions of the alkali iodides, frequently with formation of double iodides. An aqueous solution of KI dissolves much I , with formation of KI_2 , but the greater part of the I thus dissolved is pptd on largely diluting the liquid. Some other periodides analogous to KI_2 are known, part of the I in such compounds is more loosely combined than the rest, and can generally be removed very easily, CuI_2 in solution, for instance,

is reduced to CuI by shaking with CS_2 . Some iodides, e.g. PtI_2 , HgI_2 , combine with HI to form compounds which are best regarded as metallic acids (H_2PtI_4 , H_2HgI_4 , &c.) Some of the resemblances and differences between the three allied classes, chlorides, bromides, and iodides, are considered in the article HALOGENS, BINARY COMPOUNDS OF THE (vol 11 p 666)

M M P M

IODINE I At w 126.53 Mol w 253.06 (*v. infra*) [113°–115°] (Stas), [114.15°] (Ramsay & Young, *C J* 49, 460) solidifies at 113.6° (Regnault, *J* 1856 41), (200°) (Stas), (184.35°) at 760 mm (B & Y, *I c*) Sublimes *in vacuo* without melting (L Meyer, *B* 8, 1627) S G 4.917 at 40.3°, 4.886 at 60°, 4.857 at 79.6°, 4.841 at 89.8°, 4.825 at 107°, 4.004 liquid at 107°, 3.866 liquid at 151°, 3.796 liquid at 170°, vol increases for 1° by 0.00235 (Billet, *J* 1855 46) ∇D c 250°–1000° 125–127, c 1500° 68 (*v Properties*, p 10) S H (solid 9°–98°) 0.5412 (Regnault, *A Ch* [2] 73, 1) S H p for I vapour at 206°–377° (equal wt of water = 1) 0.3369

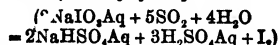
(Strecker, *W* 17, 85), $\frac{S H p}{S H v}$ 1.294 at 220°–375°

(Strecker, *W* 13, 20) Heat of fusion = 11,710, heat of vaporisation = 23,950 (Favre & Silbermann, *A Ch* [3] 37, 461) For vapour pressures of solid I from 58° to 114°, and of liquid I from 114° to 185°, *v* Ramsay & Young, *C J* 49, 458 S V S 25.9 S 0.18 at 10°–12° (Wittstein, *J* 1857 123), 0.15 at 6.3° (Dossios & Weith, *Z* 1869 379) S in glycerin = 1 (Regarding solubility *v* further under *Properties*) The absorption spectrum of I vapour shows numerous fine lines extending from the red to the violet. As the thickness of the layer increases absorption becomes very marked in the red, but even when the whole of the red part is obscured, the violet portion remains without bands (Plucker, *J* 1863 109, Thalen, *P* 139, 503, *c* Conroy, *J* 1876 146, and Salet, *A Ch* [4] 28, 29) The emission spectrum shows many bright lines in the yellow-green and yellow. By using an induction current of low tension lines are seen coincident with the dark absorption lines (Salet, *I c*, Wullner, *P* 120, 158), at red heat a continuous spectrum is observed. I crystallises very easily from alcohol or HIAq , or by sublimation, the crystals are trimetric, $a b c = 4866 : 1 : 7324$ (Mitscherlich, *B* 1855 416)

Iodine was discovered by Courtois in 1811 in the mother liquor of kelp after preparation of Na_2CO_3 , Davy (*T* 1814 74) and Gay Lussac (*A Ch* 88, 311, 319, 91, 5) showed I to be an element, and established its relationship to chlorine

Occurrence—Certain mineral waters contain small quantities of free I. According to Wanklyn (*C N* 54, 300), the water of Woodhall Spa, near Lincoln, is coloured slightly brown by free I. Iodides of Na and Mg, and iodate of Mg according to Sonstadt (*C N* 25, 196, 231, 241), occur in small quantities in sea-water, alkaline iodides are found in very many mineral waters, in the ashes obtained by burning sea-plants, and some sea animals, and in certain specimens of Chili saltpetre, dolomite, phosphates, and some other minerals. Iodides of Hg, Ag, and Pb occur in small quantities in Mexico

Formation—1 By decomposing alkaline iodide by MnO_2 and H_2SO_4 , or by Cl, or by $\text{K}_2\text{Cr}_2\text{O}_7$ and H_2SO_4 , or LiNO_3 .—2 By heating an acidified solution of an iodide with $\text{FeCl}_3 \cdot \text{Aq}$. 3 By reduction; NaIO_3 by SO_2



Preparation—The greater part of the I of commerce is prepared from kelp, which is the product of burning sea plants, the kelp is lixiviated with hot water, and the liquid is evaporated in successive stages until most of the salts except NaI , Na_2S , and $\text{Na}_2\text{S}_2\text{O}_3$ have been removed. Enough H_2SO_4 is added to decompose the S compounds. After exposure to the air for some time S separates and SH_2 and SO_2 escape. The liquid is then run off and distilled with addition of more H_2SO_4 and MnO_2 , and the I is collected in a series of tubulated receivers fitted one into the other. For details of manufacture *v* DICTIONARY OF APPLIED CHEMISTRY. The I obtained as described is dried and sublimed. By slow resublimation fairly pure I is obtained, but ICy cannot thus be removed.

The chief impurities in commercial I are chlorides, water, traces of various salts, and sometimes ICy .

Stas (*Chem. Proport.* 137) gives two methods for preparing pure I.—1 KI is dissolved in its own weight of water, the solution is saturated with I (about 4 parts I are required to 1 part KI), water is added until formation of a permanent pp begins. After settling, the liquid is poured off and shaken with $\frac{2}{3}$ of the quantity of water required to bring down all the I which can be pptd by this method (the amount of water is determined by a preliminary experiment on a small quantity of the liquid). The separated I is washed by decantation until the washings are free from K, distilled with water from a large retort, allowed to drain, placed over dry Ca_2NO_3 , which is frequently changed (all other drying materials bring impurities into the I), and finally twice distilled after mixing with 5 p.c. finely-powdered pure BaO. The last traces of H_2O and traces of HI are thus removed.—2 Iodide is prepared by adding powdered I to cold conc. NH_4Aq until the dark brown liquid is nearly colourless, the compound is washed by decantation with cold conc. NH_4Aq until NH_4I is removed, placed on a funnel, the neck of which is drawn to a fine point, and covered with cold water. When the black colour of the compound changes to brownish, and the wash water is coloured yellowish brown, the moist iodide is placed in ten times its weight of water contained in a large glass balloon, and slowly heated on a water bath to 60°–65°, when decomposition occurs with formation of crystals of I, solution of I in NH_4IAq , and a white salt, which is probably NH_4IO_3 . When the change seems completed the liquid is warmed to 100° for a few minutes, if the temperature is at once raised above 65° decomposition occurs very rapidly, at 100° it proceeds with explosive violence, at the ordinary temperature the change is very slow. After cooling, the solid which separates is thrown on to a funnel with drawn out neck and washed with water, it is then distilled with water from a retort (NH_4IO_3 is not

volatilised), and the I is dried over Ca_2NO_3 , and finally by distilling with pure BaO as in '.

Recovery of iodine from laboratory residues Beilstein (*Z* 1870 528) recommends to evaporate with excess of Na_2CO_3 , to heat the residue until it is white, to add excess of H_2SO_4 , and to pass in NO oxides, obtained by heating starch with HNO_3 , until all I has separated, to wash the I in cold water, dry over P_2O_5 , and sublime slowly. The stream of NO oxides is best obtained by adding 15 grams starch to about 90 grams fairly conc HNO_3 in a large flask (the acid must not be added to the starch, else the latter may cake on the bottom of the flask and cause it to break), heating till red fumes appear, and then removing the flame and cooling from time to time, if necessary, by cold water. If insoluble iodides, e.g. HgI_2 , are present, Henry (*B* 2, 599) heats with water and granulated Zn or Fe filings, whereby soluble ZnI_2 or FeI_3 is produced. The final sublimation of the recovered I may conveniently be conducted as described by Mohr. The rim of a flat porcelain basin is ground with sand, so that a glass plate placed on the basin touches the rim in every part, the I is placed in the basin, a little finely powdered KI is strewn on the surface, the glass plate is placed in position and bound to the rim by a strip of paper, and the basin is placed on a sand tray and very slowly heated. The process of sublimation should occupy several days. If any ICl is present it reacts with the KI to give ICl and I.

Properties—A greyish black, soft, solid with metal like lustre. Perfectly pure I is described by Stas as quite black, whether solid or liquid. Only the thinnest plates of I are transparent (cf. Schultz Sellack, *P* 140, 334). I is very easily vapourised, the vapour corrodes the skin and mucous membranes, unsaturated vapour has a violet colour, saturated vapour appears deep blue in thin layers (Stas), a layer 10 centims thick is quite impervious to light (cf. Andrews, *C* N 24, 75). I vapour shows orange yellow fluorescence (Lommel, *W* 19, 356), solid I does not fluoresce (Stokes). Crystals of I polarise light (Conroy, *J* 1876 147). I is a non conductor of electricity (Jolly, *P* 37, 420). I dissolves freely in solutions of alkaline oxides and in HIAq , it is more soluble in solutions of NH_3 salts than in water. Dossios a Weith (*Z* 1869 379) give the following numbers for the S of I in KIAq

S.G. at 7°	P.c.			S.G. at 7°	P.c.		
	KI	I			KI	I	
1.0234	1.802	1.173		1.1382	8.663	7.368	
1.0433	8.159	2.303		1.1637	10.036	8.877	
1.0668	4.628	8.643		1.1893	11.034	9.949	
1.0881	5.935	4.778		1.2110	11.893	11.182	
1.1112	7.201	6.037		1.2293	12.613	12.060	

I is more soluble in solution of tannic acid than in water, addition of 0.15 parts of the acid increases S to 42 at 30° , and when 3.3 parts acid are present S is 24 at 12° (Koller, *Z* 1866 890). I is very sol in CS_2 , CHCl_3 , ether and alcohol, C_2H_6 and various hydrocarbons, also sol in SO_2 (Sestini, *Z* 1868 718), and in SO , (Weber, *J* pr 25, 224), solutions in ether or alcohol contain HI (cf. Carles, *Ph* [3] 5, 88). Addition of CS_2 to an aqueous solution of I causes withdrawal of most of the I from the water, according to Berthelot a. Jungfleisch

o 400 parts of the I go into solution in the CS_2 for each part remaining in the water (*C* R 69, 338). Solutions of I in CS_2 , CHCl_3 , C_2H_6 , and other liquid hydrocarbons are violet, solutions in ether, alcohol, and some other solvents are reddish brown. The reddish solutions absorb light in the violet end of the spectrum up to midway between D and E, conc violet solutions absorb the rays of mean refrangibility to the limits of the yellow and blue, very conc solutions absorb all rays except the ultra violet (cf. Vogel, *B* 11, 919).

The at w of I has been determined —1 By changing AgI into AgCl (Berzelius, *P* 14, 558, Dumas, *J* 1859 3) —2 By synthesis of ZnI_2 (Gay Lussac, *A* Ch 91, 5) —3 By synthesis of AgI (Marignac, *Bibl Univ Genève*, 46, 367, Stas, *Chem. Propriété*) —4 By determining the quantity of KI required to ppt a known quantity of Ag dissolved in HNO_3 (Marignac, *l.c.*) —5 By analysis of AgIO_3 (Stas, *l.c.*) —6 By determining V D of many binary compounds —7 By measuring S H of I.

Attempts have been made to determine the mol w of I in solution. Paterno a Nasini (*B* 21, 2153) measured the lowering of the freezing point of benzene and acetic acid produced by dissolving I in these liquids, the results point to the existence of molecules of I_2 in very dilute benzene solutions, and more complex molecules in more conc solutions, the numbers got with acetic acid lead to a mol w between I and I_2 . Loeb (*C* J 53, 805) determined the vapour pressure of I in solution in CS_2 and ether, ethereal solutions of I are reddish brown, solutions in CS_2 are violet, Loeb's results point to the mol w I_2 in red solutions, and a mol w between I_2 and I in violet solutions.

Determinations of the S G of I vapour, by Gay Lussac, Dumas, Bineau, Deville a Troost, up to c 1000° gave values from 8.73 to 8.62 (calc for $\text{I}_2 = 8.75$) V Meyer (*B* 13, 401, 1723, 14 1453) obtained the following results —8.86 at 293° , 8.72 at c 586° , 6.76 at c 842° , 5.75 at c 1027° , 5.7 at c 1570° . Crafts a Meier (*B* 13, 870, *C* R 92, 39) obtained similar results, the S G remained constant (8.8) to c 700° , even when pressure was diminished, but decreased above that temperature, and decreased the more rapidly the more the pressure was lowered, the S G decreased with rising temperature rapidly to a certain point and then more slowly, at a pressure of 230 mm S G became constant at 1400° – 1500° and was = 4.6, at 152 mm S G became constant at c 1400° , at 76 mm a constant value for S G was obtained at c 1350° . At c 1700° V D corresponds with molecular weight = I (Biltz a Meyer, *B* 22, 725). There can be little doubt that these results prove a gradual dissociation of I_2 into I (S G calc for I = 4.375) (cf. DISSOCIATION, vol II p 394, v also Naumann, *B* 13, 1050). The molecule I_2 is more easily dissociated than either of the molecules Br_2 or Cl_2 .

The atom of I is monovalent in gaseous molecules. I acts as a non metallic element, it is negative to all elements except O, F, Cl and Br. I combines directly with most of the metals, and with some of the non metals, e.g. H, Br, Cl, Se, S, and P, binary compounds of I with all non metals except B have been isolated. I is closely related to F, Cl, and Br, the relationship is con

sidered in the article HALOGEN ELEMENTS, vol II p 664, cf also IODIDES in this vol p 13

Reactions—1 I dissolves very slightly in water, the solution probably contains traces of HI. According to Cross a Higgins (C J 35, 225) I dissolves slightly in water at 100° in a sealed glass tube with production of a little alkaline iodide and iodate. Electrolysis of Iaq yields HIO_2Aq (Rich², J 1858 101)—2 I does not react directly with oxygen, even when I and O are heated to c 300° in presence of spongy Pt (Wehsarg, B 17, 2896) Ogier (C R 86, 722) says that if I vapour and O are subjected to the silent discharge, all the oxides of I are produced—3 *Hydrogen peroxide* produces HIAq and O, but conc HIAq is decomposed by H_2O_2 with separation of I—4 *Hydrogen sulphide* is decomposed by I in presence of water forming HI and S, dry H_2S does not react with I—5 With solutions of *caustic alkalis*, I forms alkaline iodide and iodate, according to Van Deventer a Van't Hoff, KIO is also formed (the proof of this is indirect, C C 1888 362)—6 *Ammonia gas* forms NH_4I and N (*in Combinations*, No 2), ammonia solution forms NH_4IAq and N iodide—7 When I is shaken with lime suspended in water, a bleaching liquid is obtained which reacts as if it contained Ca hypoiodite Ca(IO) and CaI_2 (Lunge a Schock, B 15, 1883)—8 I vapour heated with oxides or carbonates of the alkaline earth metals, in presence of oxygen yields periodates without formation of oxyiodides, with lead oxides various oxyiodides are produced (Cross a Sigura, C J 33, 405)—9 I is oxidised to I_2O_5 by nitric acid, chromic anhydride, chlorates, and some other oxidisers—10 I in presence of water acts as an oxidiser towards some salts and other easily oxidised compounds, arsenous oxide and arsenites are oxidised to As_2O_5 and arsenates in alkaline solutions. Sulphurous acid is oxidised to H_2SO_4 , sodium thiosulphate is oxidised to $\text{Na}_2\text{S}_4\text{O}_6$ and a little NaHSO_4 (Pickering, C J 37, 128)—11 With some carbon compounds I reacts to substitute I for H, but the HI formed tends to reproduce the original compound, so that the reactions of I with C compounds are much less marked than those of Cl and Br—12 I dissolved in CS_2 reacts with ppt *arsenous sulphide* (not with impure) to give AsI_3 and S, at a higher temp As_2S_3 and I are reformed, by distilling As_2S_3 and I in the ratio As_2S_3 6I and heating the distillate in a sealed tube to 72° (AsI_3)₂SI₂ was obtained (Schneider, J pr [2] 36, 498)

Combinations—No combination of I with H_2O has been isolated—1 With most metals, and with many non metals, especially with H, Cl, Br, S, Se, P, As. According to Holzmänn (B 22, 869) Na is scarcely attacked by I when the pure elements are heated to 300°. Fe and I combine by shaking Fe filings with I in presence of water, FeI_2 is produced, and also Fe_2O_3 and HI probably by the formation and decomposition of FeI_3 (Fleury, J Ph [5] 16, 529)—2 *Ammonia* forms several compounds with I, the dry gas is absorbed forming a dark blue liquid, the volume of NH_3 absorbed varies with temperature, at 20° it corresponds to 3NH_3 , 2I, at 80° to NH_4I , at 0° to $(\text{NH}_4)_2\text{I}_2$, at -10° to $(\text{NH}_4)_3\text{I}_3$ (Raschig, A 241, 253, cf Bineau, A Ch [3] 15, 80, Millon, A 62, 54)

Detection and Estimation—Free I is detected in solution by the colour which it gives to CS_2 or CHCl_3 ; and by the formation of a deep blue colour when a drop of starch paste is added (cf Béchamp, Fr 14 66, Mylius, B 20, 688, Personne, C R 74, 617, Duclaux, C R 74, 533, Goppelsröder, P 119, 57, Hlawetz, W A B 1867 181). In soluble iodides is detected by adding a very little Cl water, or a drop of a solution of NO_2 in conc H_2SO_4 , and then a little starch-paste. Iodates are reduced to iodides by SO_2Aq , alkaline iodates mixed with all alkali iodides and a drop of a weak acid, e.g. tartaric, give free I. Many insoluble iodides may be converted into soluble ZnI_2 by treatment with Zn and dilute $\text{H}_2\text{SO}_4\text{Aq}$. Most non metallic iodides are decomposed by water or caustic alkali giving HI or alkaline iodide. I is estimated gravimetrically by pptn as AgI , iodates are reduced by SO_2Aq and then pptd by AgNO_3Aq . Volumetrically it is estimated by titration with standard $\text{Na}_2\text{S}_2\text{O}_3\text{Aq}$ or standard As_2O_3 dissolved in NaHCO_3Aq . The water in samples of ordinary I may be determined by mixing with a weighed quantity of Hg , rubbing up with a little alcohol, and carefully drying at 100°, loss of weight = water (Bolley, D P J 126, 40). The quantity of I may be determined by dissolving in a conc solution of an alkali sulphate, filtering, pptg by AgNO_3Aq , washing ppt with NH_4Aq to remove AgCl and AgBr , boiling with H_2O containing a little AgNO_3 to remove AgSO_4 , washing the residual AgI , and weighing it, after drying, in the usual manner (Hesse, A 122, 225).

Iodine, bromides of Two bromides of I are known, IBr and IBr_3 , but neither has been gasified, attempts to prepare IBr_3 , analogous to ICl_3 , have failed (Bornemann, A 189, 183). The combination of I with Br has been studied by Balard (A Ch 32, 337), Lagermark (B 7, 907), and Bornemann (A 189, 201).

Iodine monobromide IBr Formula not necessarily molecular, but probably so, from analogy to ICl which has been gasified. Prepared by heating I with rather more than the calculated quantity of Br to 25°–50° in a retort, removing excess of Br by a stream of CO_2 , and allowing to cool. Forms dark greyish, iodine like crystals, melting at 36°, probably sublimes with only partial decomposition, smells of Br, the vapour attacks the eyes and mucous membranes. The vapour appears red in thick strata (Gernez, C R 74, 1190). Soluble in alcohol, CHCl_3 , CS_2 , and ether, slowly decomposed by water with separation of I. Berthelot (A Ch [5] 21, 370) gives the H of solid IBr from solid I and liquid Br as $[\text{I}, \text{Br}] = 2470$. A hydrate $\text{IBr} \cdot 5\text{H}_2\text{O}$ described by Löwig is merely a mixture of IBr and ice.

Iodine pentabromide IBr_5 A dark brown liquid obtained by heating excess of Br with I, soluble in water, with alkali solutions forms bromide and iodate, solution in sunlight decomposes to HBr and HIO_3 .

Iodine, carbide of, better called Carbon iodide, v vol I p 689

Iodine, chlorides of I combines rapidly with Cl even at -90° (Donny a Mareska, C R 20, 817). Two chlorides of I have been isolated, ICl and ICl_3 , ICl can be gasified without decomposition, ICl_3 decomposes when heated. Kammerer (J pr 83, 83, cf B 8, 489, note) asserted

the probable existence of a tetrachloride ICl_4 , but the non existence of this compound has been proved (cf Hannay, *C J* 35, 169), Brenken (*B* 8, 187) failed to procure any higher chloride by passing Cl over strongly cooled ICl . By adding I to liquid Cl , Hannay (*C J* 35, 169) obtained a reddish liquid when the elements were present in the ratio $\text{I} : 5\text{Cl}$, but on removing pressure or raising temperature Cl was at once evolved.

IODINE MONOCHLORIDE ICl Mol w 161.9 [212°], (101°), S.G. d_4^{20} 3.18223 (Thorpe, *C J* 37, 174, H.F. [I , Cl] = 2,100 (solid ICl from gaseous Cl and solid I), 6,700 (solid ICl from gaseous constituents) (Berthelot, *A Ch* [5] 21, 170). V.D. 89.29 at 120°; 81.2 at 512° (Hannay, *C J* [2] 11, 919). Absorption spectrum of vapour v. Ger. *z.*, *C R* 74, 660.

Preparation—1 By passing dry Cl over carefully dried I till liquefaction occurs and refining by distillation with a gram or two of I . 2 By heating an intimate mixture of I with 4 parts KClO_3 and distilling the product from powdered KClO_3 ($\text{I}_2 + 3\text{KClO}_3 \rightarrow \text{KClO}_4 + \text{KIO}_3 + \text{KCl} + \text{O}_2 + \text{ICl}$)—3 By boiling I with excess of *aqua regia* diluting with water, shaking with ether and evaporating the ethereal liquid (Bunsen, *A* 54, 1).

Properties— ICl seems to exist in two forms, a reddish brown oily liquid (Ray Lussac, *A Ch* 91, 5), and a highly red solid (Schutzenberger, *Z* 5, 1). Thorpe (*C J* 37, 175) says that ICl remains liquid in a closed vessel for weeks even in a freezing mixture but solidifies on addition of a minute fragment of ICl_3 . If the liquid ICl is exposed to the air it solidifies after a time and is partially changed to ICl_3 and I (cf Hannay, *C J* [2] 11, 915; Bornemann, *A* 189, 183). According to Stortenbeker (*R T C* 7, 152), two forms of solid ICl exist one forming dark red needles melting at 27° obtained by cooling the liquid ICl to -25°, and another crystallising in dark red plates melting at +14° obtained by crystallising the liquid between -10° and +5°. ICl smells of I and Cl , it attacks all mucous membranes rapidly and forms very bad sores if let fall on the skin, decolourises indigo, does not give blue colour with starch (cf Christomanos, *B* 9, 134). ICl dissolves in alcohol, also in HClAq . Can be distilled with only very slight decomposition (Schutzenberger, *Z* 5, 1, cf Bornemann, *A* 189, 183).

Reactions—1 *Water* decomposes ICl forming HCl , HIO_3 , and I (Hannay, *C J* [2] 11, 920). According to Schutzenberger (*l.c.*) a compound IClHCl is formed (besides HIO_3 and I) by the reaction of water with ICl (cf also Trapp, *J* 1854 310, and Bornemann, *A* 189, 183)—2 *Potash solution* produces KIO_3 and KI with separation of I (H , *l.c.*)—3 *Ammonia solution* forms NH_4Cl and N iodide which combines with a portion of the NH_3 (Bunsen, *A* 84, 1)—4 Distilled repeatedly with *potassium iodide*, KCl and I are formed—5 Dissolves in *carbon disulphide*, when a saturated solution is distilled CCl_4 and CSCl_2 are formed (H , *l.c.*)—6 With many *metals* ICl reacts to form chlorides and iodides (H , *l.c.*), e.g. with Hg , Sn , Sb , Bi , Al , Mg , Na —7 With *mercuric oxide*, *cupric oxide*, and *lead peroxide*, forms chlorides and iodides

with evolution of O and separation of I —8 *Hypochlorites* and *chlorates* produce iodates with evolution of Cl (Henry, *B* 3, 892)—9 *Mercuric chloride* forms a pp of HgI_2 (Kane, *J pr* 11, 250)—10 *Stannous chloride* produces SnCl_4 and SnI_4 .

Combination—With *hydrogen chloride* to form HCl ICl , a yellow, volatile, unstable compound, produced by dissolving ICl in water, extracting with ether, and evaporating (Schutzenberger, *C R* 84, 389).

IODINE TRICHLORIDE ICl_3 Mol w unknown, decomposed by heat to ICl and Cl S.G. d_4^{20} 3.1107 (Christomanos, *B* 10, 789) [25°] (Trapp, *J* 1854 310), [33°] (Christomanos, *l.c.*) H.F. (Berthelot, *A Ch* [5] 21, 370) [I , Cl_3] = 21,700 solid ICl_3 from gaseous constituents, 16,300 solid product from solid I and gaseous Cl , [ICl , Cl_2] = 9,500 solid product from solid ICl and gaseous Cl . According to Brenken (*B* 8, 487) ICl_3 is partially decomposed when it is melted Stortenbeker (*R T C* 7, 152) gives M.P. as varying from 20° to 60°, but says that at pressure of 16 atmos the M.P. is definite and = 101°.

Preparation—1 By leading excess of dry Cl over I or ICl until yellowish red crystals are formed, and then subliming at as low a temperature as possible in a stream of Cl (Brenken *B* 8 187)—2 By the action of HCl on warm powdered HIO_3 , Cl_2 is evolved (Ditte, *A* 156, 335)—3 By the action of PCl_5 on I_2 —4 Christomanos (*B* 10, 134, 789) recommends to lead dry Cl and dry HI into a well cooled glass receiver, keeping the Cl in excess ($\text{HI} + 4\text{Cl} = \text{HCl} + \text{ICl}_3$), if HI is in excess the reaction $\text{HI} + \text{ICl}_3 = \text{HCl} + 2\text{ICl}$ occurs). To prepare small quantities of ICl_3 , say in a bulb tube Christomanos leads dry Cl through the tube, cools the place where the ICl_3 is to be deposited, then allows dry HI to stream through the tube, and finally leads dry Cl again, all parts of the apparatus being gently warmed except that where the ICl_3 is to be formed.

Properties—A citron yellow, crystalline, deliquescent solid. Very disagreeable smell, attacks the eyes and nose. Can be kept unchanged only in dry Cl (Hannay, Christomanos). In air sublimation with partial decomposition begins even at -12° (C). Complete decomposition into ICl and Cl occurs at c 70°-80° even in an atmosphere of Cl at pressure of 760 mm (Melikoff, *B* 9, 490). According to Brenken (*B* 8, 187) ICl_3 does not melt when heated, but decomposes even in an atmosphere of Cl into ICl and Cl at c 25°, the temperature varying according to the pressure. ICl_3 is sol. water, with partial change to ICl , HCl , and HIO_3 , at 100° this change occurs suddenly and completely (Bornemann, *B* 10, 121, Christomanos, *l.c.*) Sol. benzene, sol. conc. H_2SO_4 .

Reactions—1 With *water* to form ICl , HCl , and HIO_3 , in cold water a part of the ICl_3 remains unchanged, in hot water the change is complete, at 100° it occurs very rapidly—2 With *caustic alkalis*, chloride and iodide, chlorate and iodate, of the alkali are produced (Christomanos, *l.c.*)—3 With excess of *ammonia*, NH_4Cl , NH_4I , and NI_3 are formed (C)—4 *Acetic acid* produces Cl and I (C)—5 *Hydrogen iodide* in excess forms ICl and HCl (C)—6 *Carbon disulphide* produces some S chloride, the solution contains

Si_2I_2 (C) (cf Weber, P 128, 459) —7 Carbon dioxide passed over ICl_2 forms a little COCl_2 (C) 8 ICl_2Aq shaken with silver oxide forms AgCl and HIO_3 , heated with excess of Al_2O_3 , Ag peroxide (Philipp, B 3, 4) —9 Hydrogen has no action at the ordinary temperature, when the ICl_2 is slightly warmed $\text{HCl} + \text{ICl}$ are produced, at a higher temperature HCl , HI , and I are obtained (C) —10 Potassium and phosphorus burn in ICl_2 , forming chlorides and iodides (C) —11 ICl_2 oxidises hot sulphurous acid solution to SO_2Aq , and hot ferrous sulphate solution to ferric sulphate (C) —12 On carbon compounds the action of ICl_2 is to chlorinate, e.g. $\text{C}_2\text{H}_5\text{O}$, even in the dark forms $\text{C}_2\text{H}_5\text{ClO}_2$, with simultaneous production of HCl and ICl

Iodine, cyanides of Better called CYANOGEN IODIDES, v vol n p 313, and CYANURIC IODIDE, v vol n p 320

Iodine, fluoride of IF_5 A colourless, strongly smelling and fuming liquid, does not solidify at -20° , obtained by decomposing AgF by I Attacks glass and Si at red heat, no action on Hg or Pt , decomposed by water to HIO_3 and HF (Gore, C N 24, 291, MacIvor, C N 32, 232)

Iodine, hydride of, v IODHYDRIC ACID, p 11

Iodine, nitride of Better called NITROGEN IODIDE (q v in this vol)

Iodine, oxides of The only oxide of I which has been certainly isolated is I_2O_5 , the existence of IO_2 is probable Other oxides have been suggested, but the proofs of their isolation are very meagre I and O do not combine directly, Wehsarg (B 17, 2896) passed I and O over spongy Pt heated to c 300° without obtaining any compound According to Ogier (C R 86, 722) when a mixture of I vapour and O is submitted to the silent electric discharge all the oxides of I are produced I_2O_5 is the anhydride of HIO_3 , which acid is known as a definite stable solid, IO_2 (if it exists) is not an anhydride, it is said to combine with H_2SO_4 The hypothetical anhydride of periodic acid, I_2O_7 , has not been isolated The heat of formation of I_2O_5 is a large positive quantity = c 45,000 I_2O_5 is a much more stable body than any of the oxides of Cl , no oxide of Br or F has been isolated

IODINE PENTOXIDE I_2O_5 (Iodic anhydride) Mol w. unknown, as oxide has not been gasified SG ρ 4.487 (Ditte, A Ch [4] 21, 5), SG ρ 4.7987 (Kammerer, J pr 79, 94) CE $0^\circ - 51^\circ = -0.00066$ HF from solid $[\text{I}^+, \text{O}^-] = 45,029$, $[\text{I}^+, \text{O}^-] = 43,237$, $[\text{I}^{2+}, \text{O}_2^-] = -1,792$ (Th 2, 164) S = 187.4 at 13° , SG of this solution = 2.1269 (Kammerer, P 138, 390) A white crystalline solid, crystals belong to trimetric system (Schabus, J 1854 310) Produced by slowly heating dry HIO_3 to 170° Melts when heated to c 300° , with decomposition into I and O When HIO_3 crystallises from solutions containing H_2SO_4 , crystals of I_2O_5 are said to accompany the HIO_3 (Rammelsberg) Sol water, insol ether, alcohol, CHCl_3 , CS_2 , or C_2H_5 (Ditte, C R 70, 621) I_2O_5 dissolves in water to form HIO_3 The general reactions of I_2O_5 are those of an oxidiser CO passed over warmed I_2O_5 forms CO_2 and I , SO_2 gives SO_3 , H_2S produces HI , H_2O , S , and I , HCl forms ICl and H_2O , NH_3 on heating gives H_2O , N , and I (Ditte, Lc.). NO

does not react with I_2O_5 (Kammerer, J pr 79, 94)

Combinations —1 According to Kammerer (J pr 83, 72) when dry SO_2 is passed over I , heated to 100° a part of the SO_2 is oxidised to SO_3 , a little I being separated, and then a yellow crystalline compound SiO_2SO_3 is formed As soon as the compound is formed the SO_2 must be stopped The compound is decomposed by moisture, even by exposure to ordinary air Ditte (C R 70, 621) says that in this reaction only I and SO_2 are produced According to Weber (B 20, 87) the compound $\text{I}_2\text{O}_3\text{SO}_3$ is formed by heating the constituents in a sealed tube to c 60° , and pouring off excess of SO_3 , the compound is decomposed above 60° —2 By heating HIO_3 slowly to $30^\circ - 40^\circ$, or more quickly to 130° , the hydrate $3\text{I}_2\text{O}_5 \cdot \text{H}_2\text{O}$ is said to be formed Ditte, however, says this is only a mixture of I_2O_5 and HIO_3

Nitroso derivative(?) By treating perfectly dry finely powdered I with c 10 times its weight of the most conc HNO_3Aq , or a mixture of equal parts of conc HNO_3Aq and H_2SO_4 , a loose flocculent yellow powder is obtained This substance was regarded by Millon (J pr 34, 319) to be a compound of I , O , and HNO_2 , according to Kammerer (J pr 83, 65) it is $\text{I}_2\text{O}_5(\text{NO})_2$, but no analyses are given The substance is extremely unstable, it cannot be dried over CaO without decomposition it is decomposed by water to HNO_3 , HIO_3 , and I Heated in CO , NO_2 and I_2O_5 are formed

IODINE DIOXIDE or TETROXIDE IO_2 , or I_2O_4 This substance probably exists as a definite compound One part of perfectly dry I is rubbed in a basin with 10–12 pts conc HNO_3Aq SG at least 1.486, until a loose flocculent yellow powder is produced (v supra), excess of acid is poured off, the powder is placed in a funnel stopped with asbestos, and is then dried on a porous tile, and finally over lime, HIO_3 and I are then removed by washing with water and then with alcohol (Millon, J pr 34, 319, 337) IO_2 is described as a sulphur yellow powder, unchanged in air, decomposed to I and HIO_3 by heating in air to $170^\circ - 180^\circ$, insol cold water, but decomposed by boiling water giving I and HIO_3 , not acted on by alcohol Warm HNO_3 produces HIO_3 and I , but H_2SO_4 dissolves the substance, and on cooling crystals of $\text{H}_2\text{SO}_4\text{IO}_2$ separate, HClAq forms ICl and Cl , aqueous alkali solutions produce iodates, but alcoholic solutions of alkalis are said to form red liquids containing very unstable compounds

Other compounds of iodine and oxygen have been described, but it is very doubtful whether any definite compound except I_2O_5 , and probably IO , has been isolated By the action of ozone on I , Andrews & Tait obtained a yellow powder, supposed by some chemists to be I_2O (A & T Pr 9, 608, no analyses given) Ogier (C R 85, 957) asserted the formation of I_2O , by the action of ozonised O on I , he described it as a yellow deliquescent powder, decomposing at $125^\circ - 130^\circ$, with evolution of I and O , giving HIO_3 and I with water Millon (J pr 84, 336) supposed he had obtained I_2O_3 by exposing to moist air the product of boiling I_2O_5 with conc H_2SO_4 , until both I and O were coming off. Kammerer, (J pr 88, 78) treated I_2O_5 with SO_2 , and by

washing the product with alcohol and water obtained a brownish yellow powder supposed to be I_2O_5 . These substances were probably mixtures Bengieser [in 1836] (*A* 17, 2nd 4) says that by heating H_2IO_4 (*v* Periodic acid) to 160° it loses water, and that at c 180° O is evolved and I_2O_5 remains, possibly I_2O_4 may be formed at c 160° , but Rammelsberg (*J* pr 193, 278) says that decomposition of H_2IO_4 begins at 133° , and Langlois (*J* pr 56, 36) puts the decomposition temperature at 130° .

Iodine, oxyacids of. The only oxyacids of I which are certainly known are HIO_4 and H_2IO_4 , both are solids. There are indications of the existence of HIO in aqueous solution, but neither this acid nor any of its salts has been actually isolated. The periodic acid corresponding with $HClO_4$ has not been isolated, but many salts of the form MIO_4 are known, the periodates form several complex series of salts whereas all the perchlorates belong to the series $MClO_4$. HIO_4 is a product of the action of many oxidisers on I, *e.g.* HNO_3 , Cl in presence of H_2O , $KOHAq$. This acid or an iodate is also formed by digesting Mg , HgO , or Ag O with I and H_2O , or by shaking up $AgNO_3$ with an alcoholic solution of I. Electrolysis of IAq or $HIAq$ yields HIO_4 . Iodides dissolved in water are oxidised by $KMnO_4$, and by bleaching powder, to iodates. HIO_4 is produced by heating solutions of $HClO_4$, $HClO_3$, $HClO_2$, or $HBrO_3$ with I. If the reactions are compared with those whereby bromic and chloric acids are obtained (vol i p 537, and vol ii p 15) it is seen that it is more easy to pass from less oxidised, or non oxidised, compounds of I to HIO_4 , than from corresponding compounds of Cl or Br to $HClO_4$ or $HBrO_4$. The heat of formation of HIO_4 is much greater than that of HI , whereas the heats of formation of $HClO_4$ and $HBrO_4$ are considerably less than those of HCl and HBr [H , I, Aq] -13,170, [H , I, O^4 , Aq] = 55,800 (*cf* vol ii p 665) Periodic acid and its salts are stable compounds, they are more readily formed by oxidation processes than perchlorates, *e.g.* passing I vapour with O over heated BaO produces Ba periodate, and passing Cl into an alkaline iodate in presence of alkali produces an alkaline periodate. The anhydride of iodic acid, I_2O_5 , is known as a stable solid, but the anhydride of periodic acid, I_2O_7 , has not been isolated (*supra*, Iodine, oxides of).

Detection and estimation of iodates and periodates—Iodates are detected by mixing with an alkaline iodide, adding a little starch paste, and a few drops of a weak acid, *e.g.* tartaric acid, when I is set free and colours the starch blue. Periodates give a brown pp of $AgIO_4$ on addition of $AgNO_3$ in presence of HNO_3 . Periodates may be separated from iodates by ppg both as Ba salts and digesting pp with NH_3 carbonate, when Ba periodate remains unchanged, but the iodate is converted into $BaCO_3$. Iodates may be estimated by reduction to iodides by means of SO_2 or SH_2 , and pppn with $AgNO_3$, or by digesting with KI and a little HCl , when Cl is set free and decomposes the KI , giving free I which is determined volumetrically. Periodates may be estimated by a similar method (*J. Kimmins, O J* 61, 301).

Hydro iodic acid and Hydro-iodic acids Neither the acid nor any of its salts has been isolated,

According to Kōne (*P* 66, 302, Lippmann, *B* 7, 1773) the solution obtained by shaking an alcoholic solution of I with ppg HgO probably contains hypo iodic acid, but this acid quickly decomposes to HIO , and I. When I is added to $KOHAq$ or $NaOHAq$ a yellowish liquid is obtained, which bleaches indigo, gives a blue colour with starch, and is decomposed by H_2O_2 with evolution of O, on heating, iodide and iodate are formed. This solution probably contains KIO or $NaIO$ (Schönbein, *J* pr 84, 337, Berthelot, *B* 10, 900, Van Deventer a Van't Hoff, *C C* 1888 362) Lunge a Schock (*B* 15, 1883), by the action of I on CaO suspended in water, obtained a colourless solution which bleached logwood, litmus, and cochineal, gave no colour with starch, addition of acids separated I, H_2O_2 caused evolution of O, the solution decomposed slowly in the dark, more rapidly in sunlight, but even on boiling for some hours it was only partially decomposed. L a S suppose this solution to contain an I compound, analogous with bleaching powder, probably $CaOI$.

Iodic acid, and Iodates HIO_3 , MIO_3 . The acid seems to have been first obtained by Connel by oxidising I by conc nitric acid (*New Edin Philos Journ* 10, 93, 337, 11, 72, 13, 284).

Occurrence—Sometimes in commercial nitric acid (Pettenkofer, *J* 1857 581).

Formation—1 By oxidising I by very conc HNO_3 , Aq , or by $HBrO_3$, Aq (Kammerer, *J* pr 79, 94), or $HClO_4$, Aq (Davy, *S* 11, 68, 234, 16, 343)—2 By decomposing $Ba(IO_3)_2$ by the proper quantity of H_2SO_4 , Aq —3 By suspending $AgIO_3$ in water, adding an equivalent quantity of I, filtering from AgI , and evaporating to dryness ($10AgIO_3 + 12I + 6H_2O + Aq = 12HIO_4 + 10AgI$, Kammerer, *P* 138, 390). The $AgIO_3$ is prepared by ppg NH_4IO_3 by $AgNO_3$ and the NH_4IO_3 is made by digesting $Ba(IO_3)_2$ with solution of NH_3 carbonate—4 By digesting an aqueous or alcoholic solution of $AgNO_3$ with I ($10AgNO_3 + 12I + 6H_2O = 10HNO_3 + 10AgI + 2HIO_4$, Lassaigne, *J Chim méd* 9, 508, Weltzien, *A* 91, 43)—5 By decomposing KIO_4 , Aq by H_2SiF_6 , Aq , filtering, and evaporating to dryness, the product is impure—6 By leading Cl into water containing I in suspension, the greater the dilution the greater the quantity of Cl required to change all the I into HIO_4 (*cf* Bornemann, *A* 139, 183, Sodini, *B* 9, 1126)—7 By electrolyzing solution of I or HI (Riche, *C R* 46, 348)—8 By the action of Au_2O_3 on I in presence of water ($6I + 5Au_2O_3 + 8H_2O + Aq = 6HIO_4 + 10Au$, Colin, *G A* 48, 280)—9 By moistening ICl with a little water and then shaking with ether or alcohol (Liebig, *P* 24, 363)—10 Alkaline iodates are obtained by acting on I with caustic alkalis, or by oxidising alkaline iodides by $KMnO_4$, Aq or solution of bleaching powder (Hempel, *A* 107, 100, Reinige, *Fv* 9, 39, Reichardt, *Ar Ph* [3] 5, 109 $KIAq + 2KMnO_4 + H_2O = KIO_4 + 2MnO_2 + 2KOH$, Aq , $2KIAq + 5CaOCl_2$, $Aq = Ca(IO_4)_2 + 5CaCl_2$, $Aq + 2KCl$, Aq —11 $Hg(IO_3)_2$ is obtained, along with HgI_2 , by shaking I with ppg HgO suspended in water (Colin, *G A* 48 280).

Preparation—1 About 5-10 grams finely powdered I is placed in a large flask, twice it

weight of conc nitric acid SG 1.5 (not fuming acid) is added, and the bottom of the flask is gently warmed, the I is gradually oxidised, by keeping the upper parts of the flask cool any I which volatilises is prevented from escaping. After a time the acid becomes diluted, it is then poured off, fresh acid is added, and the oxidation is continued until the whole of the I has been converted into white crystals of HIO_3 . The greater part of the acid is poured off, the semi-liquid mass is evaporated to dryness in a basin, and the remaining acid is removed either by repeated evaporations with water or by heating to $100^\circ\text{--}130^\circ$ in an air current. The crystalline mass may be dissolved in water and slowly evaporated to the crystallisation point, and the crystals heated to $170^\circ\text{--}2$. Two pts conc H_2SO_4 and c 8 pts water are added to 9 pts finely powdered $\text{Ba}(\text{IO}_3)_2$, the whole is boiled for half an hour, after settling, BaSO_4 is removed by filtration, and the filtrate is evaporated until HIO_3 separates on cooling, the crystals are dissolved in water, a very little $\text{Ba}(\text{IO}_3)_2$ is added, and the liquid is evaporated and filtered. Stas says that pure HIO_3 cannot be obtained by this method, the crystals always contain either $\text{Ba}(\text{IO}_3)_2$ or H_2SO_4 . The $\text{Ba}(\text{IO}_3)_2$ required may be prepared (1) by adding BaCl_2Aq to NaIO_3Aq obtained by passing Cl into water containing I in suspension, till the I is all dissolved, then adding Na_2CO_3 till neutral, and again passing in Cl (Liebig, *P* 24, 363), (2) by suspending I in hot sat'rated BaO_3Aq and passing Cl into the liquid (Kammerer, *J* 1860 94), (3) by adding rather more than the equivalent quantity of I to hot conc KClO_3Aq , and then a few drops of nitric acid, Cl is freely evolved, and KIO_3 crystallises on cooling, the salt is recrystallised once and decomposed by BaCl_2Aq — 3 NH_4IO_3 is prepared by digesting $\text{Ba}(\text{IO}_3)_2$ with NH_4 carbonate solution, filtering, and crystallising, AgNO_3Aq is added to a solution of the NH_4IO_3 and the AgIO_3 obtained is collected and washed, the AgIO_3 is suspended in water and I is added in the ratio of 1 gram to 1.857 grams of the AgIO_3 , the reaction $10\text{AgIO}_3 + 12\text{I} + 6\text{H}_2\text{O} = 12\text{HIO}_3 + 10\text{AgI}$ proceeds when the liquid is warmed on the water bath, AgI is removed by filtration and the liquid is evaporated to dryness and the residue heated to c 170° (Kammerer, *P* 138, 390).

Properties — HIO_3 forms colourless tetrametric crystals, $a:b:c = 589:1:1903$ (Rammelsberg, *P* 90, 12), $a:b:c = 9388:1:13181$ (Schabus, *J* 1854 310). It is doubtful whether HIO_3 shows dimorphism or not (*v* Rammelsberg, *Handbuch der Krystall-physikal. Chemie*, 1 41). Thomsen gives the following thermal data (*Th* 2, 163) — $[\text{H}, \text{I}, \text{O}^*] = 57,963$, $[\text{HIO}_3^*, \text{Aq}] = -2,166$, $[\text{I}^*, \text{H}^*, \text{O}^*] = 2,540$ SG $\frac{2}{4}$ 629 (Ditte, *C R* 70, 621). Very soluble in water, most conc solution, according to Kammerer, contains 68.51 p.c. HIO_3 , boils at 100° , and has SG 2.1629. Kammerer gives the following table —

SG at 14°	P.c. HIO_3	SG at 14°	P.c. HIO_3
1.0058	1.054	1.4428	36.89
1.0263	5.27	1.5371	42.16
1.0525	10.54	1.6315	46.93
1.1223	15.81	1.7356	52.70
1.2093	21.08	1.8689	57.97
1.2173	26.85	1.9954	63.24
1.3484	31.62	2.1269	68.51

Thomsen has determined the volume change attending the dilution of HIO_3Aq . He expresses

it by the formula $V_a = 18a + 39\left(1 - \frac{a}{a+18}\right)^{13.1}$;

the composition of the solution is represented by $\text{HIO}_3, a\text{H}_2\text{O}$, the volume of one molecular weight of water is taken as 18. For optical properties of HIO_3 crystals *v* Lang, *W A B*. 31. S.H. of $\text{HIO}_3 = 1625$ (Ditte, *A Ch* [4] 21, 52). Electrolysis of HIO_3 produces I and O (Magnus, *P* 102, 1, Buff, *A* 110 257).

Reactions — HIO_3 reacts as an energetic oxidiser. 1 Most non metals are oxidised by HIO_3Aq , e.g. P to H_3PO_4 , As to H_3AsO_4 , B to HBO_3 , Si at 250° to SiO_2 (Ditte, *Bl* 1870 318), S, Se, and C are oxidised by heating in sealed tubes with HIO_3Aq , S to H_2SO_4 , Se to H_2SeO_4 , gas coke at 180° and anthracite at 210° to CO_2 , diamond is not acted on — 2 All metals, except the Pt metals and Au, are oxidised by HIO_3Aq — 3 Sulphurous acid is oxidised to H_2SO_4 , sulphuretted hydrogen to S and HI, the lower oxides of nitrogen to HNO_3 — 4 Hydrochloric acid forms ICl_3 and Cl .

Combinations — 1 With water to form $2\text{HIO}_3, 9\text{H}_2\text{O}$, obtained by cooling saturated HIO_3Aq to -17° — 2 With sulphuric acid, by dissolving HIO_3 in hot conc H_2SO_4 and cooling, crystals of $2\text{HIO}_3, 3\text{H}_2\text{SO}_4$ are said to be obtained, the mother liquor deposits other compounds of the two acids, if $\text{H}_2\text{SO}_4, 2\text{H}_2\text{O}$ is used crystals of $3(\text{H}_2\text{SO}_4, 2\text{H}_2\text{O}), 2\text{HIO}_3, 2\text{H}_2\text{O}$ are formed, these compounds are decomposed by water (Millon, *J pr* 34, 321). Crystalline bodies, supposed to be compounds (Davy), are obtained by mixing conc HIO_3Aq with solutions of H_3PO_4 and HNO_3 .

Basicity and formula of iodic acid — HIO_3 is generally looked on as a monobasic acid, analogous to HClO_3 and HBrO_3 . Besides the normal K salt there exist two acid salts, KHI_2O_6 and KHI_4O_7 , there are no chlorates or bromates similar to these. Iodic acid is easily decomposed by heat to water and the anhydride I_2O_5 , such a reaction does not usually occur with monobasic acids. Iodic acid is isomorphous with the dibasic acids, succinic and itaconic. The chlorates and bromates as a class are easily soluble in water, the iodates are very slightly soluble. The thermal phenomena attending the formation of the three acids, HClO_3 , HBrO_3 , and HIO_3 , mark off HIO_3 from the two others, thus —

$[\text{H}, \text{Cl}, \text{Aq}] = 39,320$	$[\text{H}, \text{Cl}, \text{O}^*, \text{Aq}] = 23,940$
$[\text{H}, \text{Br}, \text{Aq}] = 28,380$	$[\text{H}, \text{Br}, \text{O}^*, \text{Aq}] = 12,420$
$[\text{H}, \text{I}, \text{Aq}] = 13,170$	$[\text{H}, \text{I}, \text{O}^*, \text{Aq}] = 55,800$

The heat of formation of the acid HX decreases as the atomic weight of X increases, whereas in the series of oxyacids the heat of formation decreases from Cl to Br, but then increases very largely from Br to I. A similar variation is shown in the K salts, thus —

$[\text{K}, \text{Cl}] = 105,610$	$[\text{K}, \text{Cl}, \text{O}^*] = 95,840$
$[\text{K}, \text{Br}] = 95,310$	$[\text{K}, \text{Br}, \text{O}^*] = 84,062$
$[\text{K}, \text{I}] = 90,180$	$[\text{K}, \text{I}, \text{O}^*] = 124,489$

The iodates are not generally isomorphous with the chlorates and bromates. Thomsen says the only case of isomorphism is presented by the Ba salts. There is no doubt that the constitution of periodic acid is very different

from that of perchloric acid, but iodic acid shows some fairly marked analogies with periodic acid. Thus, if one molecular proportion of H_5IO_6 (228 grams) is dissolved in $80\text{H}_2\text{O}$ (1,440 grams), the volume of the solution is $1,440 + 60.2 = 1500.2$ cc., if the molecular weight of iodic acid is taken as $\text{H}_5\text{I}_2\text{O}_6$, and this quantity in grams (352) is dissolved in $80\text{H}_2\text{O}$, the volume of the solution is $1440 + 59.9 = 1499.9$ cc. In other words, one molecular proportion of H_5IO_6 , dissolved in 80 molecular proportions of water, produces the same expansion of the liquid as is produced by one molecular proportion of iodic acid, provided the formula of this acid is taken to be $\text{H}_5\text{I}_2\text{O}_6$. The foregoing are the chief arguments brought forward by Thomsen for establishing a difference between the constitutions of iodic acid on the one hand and chloric and bromic acids on the other hand, and for showing that iodic and periodic acids are closely related (*Th* 3, 168, 423).

The heats of neutralisation of HClO_3 , HBrO_3 , and HIO_3 are practically identical (*Th* 1, 242). When NaOHAq is added to NaIO_3 , a very small quantity of heat is developed, about 4 p.c. of the total heat of neutralisation, but the reaction of NaOH with the monosodium salt of an undoubted dibasic acid is always accompanied by the production of as much, or nearly as much, heat as attends the addition of the first molecular weight of NaOH to the acid. If iodic acid is regarded as dibasic, then the salt $\text{KH}_2\text{I}_2\text{O}_6$ must be looked on as either a compound of the normal salt $\text{K}_2\text{I}_2\text{O}_6$ with $2\text{H}_2\text{I}_2\text{O}_6$ ($2\text{KH}_2\text{I}_2\text{O}_6 = \text{K}_2\text{I}_2\text{O}_6 + 2\text{H}_2\text{I}_2\text{O}_6$), or as the acid salt of a hypothetical acid $\text{H}_4\text{I}_2\text{O}_6$.

On the whole there appear to be marked differences between the oxyacids of I and those of Cl and Br. The oxyacids of I form more complex salts than those of the other halogens. The constitution of the iodates and periodates cannot be settled by defining the basicities of the acids HIO_3 and H_5IO_6 . Several series of periodates certainly exist, and there is probably more than a single series of iodates.

The salts $\text{KH}_2\text{I}_2\text{O}_6$ and $\text{KH}_3\text{I}_2\text{O}_6$ may of course be formulated as $\text{K}_2\text{O} \cdot 2\text{I}_2\text{O}_5 \cdot \text{H}_2\text{O}$ and $\text{K}_2\text{O} \cdot 3\text{I}_2\text{O}_5 \cdot 2\text{H}_2\text{O}$ respectively.

Iodates. These salts are formed by neutralising HIO_3 by bases, or in some cases by oxidising I in presence of a base (*v. Iodic acid*, *Formation*, Nos 10 and 11), or by double decomposition from the alkali iodates. The iodates are generally insoluble or slightly sol. in water, the alkali iodates are readily soluble, KIO_3 , however, is only slightly soluble (S at $20^\circ = \text{c } 8$). The iodates are decomposed by heat, generally giving a mixture of metallic iodide and oxide, sometimes iodide only. Solutions of iodates are more easily reduced than chlorates, reduction of KIO_3Aq is brought about by SO_2Aq or SH_2Aq , with HIAq iodates give I and metallic iodide, with HClAq they give ICl , Cl_2 , H_2O , and metallic chloride, dilute H_2SO_4 produces HIO_3 .

The following are the chief memoirs on iodates, they are referred to by numbers in the following descriptions. —(1) Bell, *J* 1871 298, (2) Berthelot, *C R* 84, 1408 (3) Cameron, *J* 1876 284 (4) Clarke, *J* 1877 48, 267 (5) Connell, *S* 62, 498 (6) Ditté, *C R* 70, 622 (7) Flight, *J* 1864 147 (8) Gay-Lussac, *G. A.*

48, 24, 372, 49, 1, 211 (9) Goriach, *Fr* 1869 290 (10) Grosourdy, *J Chim méd* 9, 428 (11) Henry, *B* 3, 893 (12) Kammerer, *J pr* 79, 94 (13) Kremers, *P* 84, 271, 97, 5, 99, 443 (14) Ladenburg, *A* 135, 1 (15) Liebig, *P* 24, 363 (16) Margnac, *J* 1856 296. (17) Melsens, *C C* 1872 552 (18) Millon, *A Ch* [8] 9, 400, 12, 330, 13, 29 (19) Mitscherlich, *P* 11, 162, 17, 481 (20) Naquet, *J* 1860 401 (21) Peilagri, *B* 8, 1357 (22) Penny, *A* 37, 202 (23) Pleischl, *S* 45, 18 (24) Ram melsberg, *P* 44, 545, 46, 159, 62, 416, 90, 12; 115, 584, 125, 147; 134, 368, 499, 137, 305; (25) Schönbein, *J* 1857 63 (26) Serullas, *P* 19, 97, 112, 20, 515 (27) Sonnstadt, *J* 1872 187 (28) Stas, *J* 1867 162.

Ammonium iodate NH_4IO_3 . Lustrous plates, decompose at 150° , S 26 at 15° , 145 at 100° S G 331-334. Formed by action of NH_3Aq on I, or of HIO_3Aq on NH_3Aq or $(\text{NH}_4)_2\text{CO}_3\text{Aq}$ (4, 24, 28).

Barium iodate $\text{Ba}(\text{IO}_3)_2 \cdot \text{H}_2\text{O}$. By dissolving I or ICl_3 in BaOAq , or by ppg NaIO_3Aq by BaCl_2 or $\text{Ba}(\text{NO}_3)_2$. Lustrous monoclinic crystals, which lose H_2O at 130° , and when strongly heated form Ba periodate $\text{Ba}_2\text{I}_2\text{O}_6$. S G 518-528 S 07 at 135° , 15 at 100° (4, 8, 10, 12, 13, 16, 18, 24, 27).

Calcium iodate $\text{Ca}(\text{IO}_3)_2$. Occurs in seawater. Formed by adding CaCl_2Aq to KIO_3Aq , or HIO_3Aq to $\text{Ca}(\text{NO}_3)_2\text{Aq}$. By action of bleaching powder on KIAq crystallises with $6\text{H}_2\text{O}$, and from KIO_3Aq + $\text{Ca}(\text{NO}_3)_2\text{Aq}$ with $4\text{H}_2\text{O}$ (Flight). Efflorescent rhombic crystals, on heating gives mixture of CaO and Ca periodate, v insol. water (6, 16, 18, 24, 27).

Copper iodates — $\text{Cu}(\text{IO}_3)_2 \cdot 2\text{H}_2\text{O}$, greenish-blue pp by mixing conc NaIO_3Aq and CuSO_4Aq , salt with H_2O remains on warming. S 33 cold, 65 at 100° — $2 \cdot 3\text{Cu}(\text{IO}_3)_2 \cdot 3\text{CuO} \cdot 2\text{H}_2\text{O}$, by action of HIO_3Aq on strongly heated CuO . By dissolving (1) in NH_3Aq the compound $\text{Cu}(\text{IO}_3)_2 \cdot 4\text{NH}_3 \cdot 3\text{H}_2\text{O}$ is formed (18).

Mercury iodate $\text{Hg}(\text{IO}_3)_2$. By warming freshly ppg HgO with HIO_3Aq , or adding HIO_3Aq to $\text{Hg}(\text{NO}_3)_2\text{Aq}$ or $\text{Hg}(\text{C}_2\text{H}_3\text{O}_2)_2\text{Aq}$ (not by adding HIO_3 to HgCl_2). White powder, insol water (3).

Potassium iodates — Normal iodate KIO_3 . Prepared by dissolving I in KOHAq , evaporating to dryness, and dissolving out KI by alcohol S G 81, also by passing Cl into I suspended in water, neutralising by KOHAq and evaporating. Stas adds $1\frac{1}{2}$ pts KClO_3 to semi fluid KI in a crucible, dissolves, on cooling, in hot water, and recrystallises the KIO_3 , which separates (8, 16, 17, 28). Crystallises in cubical forms S G 13 3976, KIO_3Aq S G 10741 at 19.5° contains 9.08 KIO_3 to 100 water (13). S at 5° 5.3, at 9.5° 6.7, at 14° 7.7, at 22.2° 9.2, at 45.8° 16.6, at 69.2° 27 (8). Insol alcohol S G 81. Poisonous. Decomposes, at higher temperature than KClO_3 , to KI and O , without production of KIO_4 , heated with MnO_2 forms I_2 and K_2O (8, 24, 25). KIO_3Aq shaken with finely-divided Fe gives KI (21). From solution in hot dilute $\text{H}_2\text{SO}_4\text{Aq}$ rhombic crystals of $2\text{KIO}_3 \cdot \text{H}_2\text{O}$ separate, which lose H_2O at 190° (6). The double salt $\text{KIO}_3 \cdot \text{KH}_2\text{SO}_4$ is obtained by heating KIO_3 in large excess of dilute $\text{H}_2\text{SO}_4\text{Aq}$, evaporating at 25° until $\text{KH}_2\text{I}_2\text{O}_6$ crystallises out, and further

crystallising the mother liquor (16) — 2 Di-iodate or acid iodate $\text{KH}_2\text{I}_2\text{O}_6$ (or $\text{K}_2\text{O} \cdot 2\text{I}_2\text{O}_5 \cdot \text{H}_2\text{O}$) Obtained by half neutralising HIO_3Aq by KOH Aq ; also by acidifying KIO_3Aq by HCl and H_2O by alcohol, also by dissolving KIO_3 in hot dilute HNO_3Aq (6, 18, 26) Crystallises in three forms, one rhombic and two monoclinic (16) S 1.33 at 15° (26) Solution reacts acid Insol alcohol Loses H_2O at 200° (6) Forms a double salt $\text{KH}_2\text{I}_2\text{O}_6 \cdot 2\text{KCl}$, obtained by heating KIO_3 with dilute HCl Aq , or by adding to ICl_3Aq less than enough KOH Aq to saturate it (c ratio $\text{KOH} \cdot 2\text{ICl}_3$) and allowing to evaporate (16, 24, 26) 3 Tri-iodate $\text{KH}_3\text{I}_3\text{O}_9$ (or $\text{K}_2\text{O} \cdot 3\text{I}_2\text{O}_5 \cdot 2\text{H}_2\text{O}$) Formed by adding a large excess of HIO_3Aq to KIO_3Aq and evaporating, also by heating KIO_3 with large excess of dilute $\text{H}_2\text{SO}_4\text{Aq}$ and evaporating at 25° Large transparent triclinic crystals, lose all H_2O at 200° S 4 at 15° (6, 16, 24, 26) Berthelot (2) describes a basic iodate $\text{KIO}_3 \cdot \text{K}_2\text{O}$ obtained by heating KI in O

Silver iodate AgIO_3 Formed by ppg AgNO_3Aq by HIO_3Aq or NaIO_3Aq Crystallises from NH_4Aq in monoclinic crystals S G 5.4 Soluble without decomposition in dilute HNO_3Aq (4, 14, 16, 24, 28)

Sodium iodate NaIO_3 Prepared by saturating 10 pts water holding 1 pt I in suspension, with Cl , neutralising by Na_2CO_3 , again passing Cl , again neutralising by Na_2CO_3 and passing Cl , and so on, finally the solution is evaporated to $\frac{1}{10}$ th its bulk and mixed while warm with half its volume of alcohol, the crystals which separate are pressed and washed with alcohol till free from NaCl (15) Crystallises at under 5° with $2\text{H}_2\text{O}$, above 5° with H_2O , at 70° crystals of NaIO_3 form, hydrates with 8, 6 and 3 H_2O are also described (6, 18, 24) S 2.62 at 0° , 3.39 at 100° (13) Loses O and I when heated (15) According to Rammelsberg (24), a compound of NaI with Na_2O_2 ($6\text{NaI} \cdot \text{Na}_2\text{O}_2$) remains on heating NaIO_3 With conc HCl Aq , Cl is evolved, and a compound of HIO_3 and NaCl remains (6) Double salts with NaI are obtained by evaporating mixtures of NaIO_3Aq with NaIAq , $\text{NaI} \cdot \text{NaIO}_3 \cdot 8\text{H}_2\text{O}$, $\text{NaI} \cdot \text{NaIO}_3 \cdot 10\text{H}_2\text{O}$, and $3\text{NaI} \cdot 2\text{NaIO}_3 \cdot 20\text{H}_2\text{O}$ are described (6, 16, 19, 24) The existence of acid iodates is denied (6, 32, cf 18, 22, 26)

Besides the foregoing iodates, the following have been prepared — $\text{Cd}(\text{IO}_3)_2$ (24), $\text{Co}(\text{IO}_3)_2 \cdot \text{H}_2\text{O}$ and $\text{Co}(\text{IO}_3)_2 \cdot 6\text{H}_2\text{O}$ (4, 24), various Fe iodates (1, 5, 24), $\text{Pb}(\text{IO}_3)_2$ (23, 24), $\text{Mg}(\text{IO}_3)_2 \cdot 4\text{H}_2\text{O}$ (6, 16, 18), $\text{Ni}(\text{IO}_3)_2 \cdot \text{H}_2\text{O}$ and $\text{Ni}(\text{IO}_3)_2 \cdot 6\text{H}_2\text{O}$ (4, 24), $\text{Sr}(\text{IO}_3)_2$ (24), $\text{Zn}(\text{IO}_3)_2 \cdot 2\text{H}_2\text{O}$ Iodates of Al , Bi , Ce , Cr , Di , Au , La , Li , Mn , Sr , Sn , U , Y , and Zn also seem to exist, but they have not been fully investigated

PERIODIC ACID AND PERIODATES — The only acid which has been isolated is H_5IO_6 , but at least five distinct series of periodates are known The anhydride of periodic acid has not been obtained, when the acid is heated it loses H_2O , O , and I , and iodic anhydride, I_2O_5 , remains, Bengieser in 1836 (*A* 17, 254) stated that by heating periodic acid to 160° it lost water of crystallisation, and that O comes off rapidly at 180° , but according to Rammelsberg (*J* pr-103, 278) and Langlois (*J* pr. 56, 86) decomposition begins at $c. 180^\circ$.

Periodic acid was first prepared by Magnus a Ammermüller (*P* 28, 514), they prepared it by reacting on AgIO_3 with cold water, whereby the acid went into solution and another Ag periodate, $\text{Ag}_2\text{I}_2\text{O}_7 \cdot 3\text{H}_2\text{O}$, remained

Formation — By the action of I on conc HClO_4Aq (Kammerer, *P* 138, 406, 410)

Preparation — 1 $\text{Na}_2\text{H}_2\text{IO}_6$ is prepared by saturating a hot solution of equal parts NaOH and NaIO_3 with Cl , concentrating, and allowing to cool (Langlois, *A Ch* 34, 257) This process yields a mixture of $\text{Na}_2\text{H}_2\text{IO}_6$ and $\text{Na}_2\text{H}_4\text{IO}_6$, the latter salt is more soluble and may be removed by long continued washing with cold water (Kimmins, *C* 51, 357), but it is not necessary to do this in preparing H_5IO_6 The Na salt is dissolved in water with addition of just enough HNO_3 to form a clear solution, and AgNO_3Aq is added, a brown pp of $\text{Ag}_2\text{H}_2\text{IO}_6$ is thus obtained (Kimmins, *C* 51, 358, former observers said that Ag_2IO_6 or $\text{Ag}_2\text{H}_4\text{IO}_6$ is produced) The brown Ag salt is suspended in water, and shaken with Br , AgBr ppts, and H_2IO_6 along with HBrO_3 goes into solution, the filtrate is evaporated to the crystallising point, whereby HBrO_3 is decomposed, and is then placed over H_2SO_4 *in vacuo* (Kammerer, *P* 138, 390) — 2 Ag HIO_4 prepared as described in 1 is dissolved in conc HNO_3Aq , and the solution is evaporated at 100° , orange red crystals of $\text{AgIO}_4 \cdot \text{H}_2\text{O}$ separate, by treatment with cold water this salt decomposes to $\text{H}_2\text{IO}_6\text{Aq}$, and $\text{Ag}_2\text{I}_2\text{O}_7 \cdot 3\text{H}_2\text{O}$ which may be again converted into AgIO_4 by solution in conc HNO_3Aq and evaporation (Magnus a Ammermüller, *P* 28, 514) — 3 $\text{Na}_2\text{H}_2\text{IO}_6$ prepared as described in 1, and mixed with $\text{Na}_2\text{H}_4\text{IO}_6$, is dissolved in as little dilute HNO_3Aq as possible, $\text{Pb}(\text{NO}_3)_2\text{Aq}$ is added, the pp of Pb periodate ($\text{Pb}_2(\text{IO}_6)_3$, Kimmins) is digested with rather less dilute $\text{H}_2\text{SO}_4\text{Aq}$ than suffices to decompose it all, and the liquid is filtered and evaporated (Bengieser, *A* 17, 254)

Properties — H_5IO_6 crystallises in transparent, colourless prisms, probably monoclinic (Rammelsberg) M P 133° (Rammelsberg, *J* pr 103, 278), 130° (Langlois, *J* pr 56, 86), melting is accompanied by partial decomposition (Bengieser, *A* 17, 254, put the temperature of decomposition at 180°) H_5IO_6 does not lose weight at 100° , nor by keeping over H_2SO_4 Very deliquescent; fairly sol in alcohol, slightly sol in ether. Thomsen (*B* 7, 71, Th 2, 427) gives the following data for S G and expansion of $\text{H}_5\text{IO}_6\text{Aq}$ —

Ratio of H_5IO_6 , H_2O	S G of solution	Expansion for each formula-weight of acid
H_5IO_6 , $20\text{H}_2\text{O}$	1.4008	59.77
" 40 "	1.2165	59.30
" 80 "	1.1121	59.99
" 160 "	1.0570	60.2
" 320 "	1.0288	60.0

The volume when aH_5IO_6 is present is expressed by the formula $V_a = 18a + 59.8$ The following thermal data are taken from Thomsen (*Th* 2, 166) — $[\text{H}_5\text{I}_2\text{O}_7] = 185,780$, $[\text{H}_5\text{IO}_6\text{Aq}] = -1380$, $[\text{I}_2\text{O}_5\text{Aq}] = 27,000$ $\text{H}_5\text{IO}_6\text{Aq}$ exposed to air becomes yellow, and smells strongly of ozone

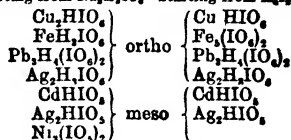
Reactions — $\text{H}_5\text{IO}_6\text{Aq}$ reacts as an energetic oxidiser, with HCl it gives Cl and HIO_4 (*M* a. A., *Lo*), with P and Bi it forms P_2O_5 and Bi_2O_3

Kammerer, *loc*); $C_2H_4O_2$, $C_2H_2O_2$, and many other acids are oxidised to CO_2 , SO_2 , Ag_2SH_2 , and HI are also oxidised, with Zn , Fe , Hg , and Cu it forms ZnO , Fe_2O_3 , HgO , and $Cu(IO_3)_2$ respectively (Bengieser, *loc*)

PERIODATES. Most of the periodates are insoluble in water, they are generally prepared from the Na or K salts by double decomposition from solutions acidified with HNO_3 . Many of them are decomposed by heat giving O and metallic iodides *e.g.* KIO_3 , $AgIO_3$, others evolve O and I , and leave either oxide, *e.g.* salts of Ni and Mg , or a mixture of oxide and iodide, *e.g.* salts of Pb , Cu , Cd . Hg salts give iodide and Hg , NH_4 salts decompose explosively to I , N , O , and H_2O . Very many periodates are known, they may be divided into several classes, the following scheme presents the chief classes and the chief salts in each class. The nomenclature is that adopted in Gmelin Kraut's *Handbuch*. All the acids except H_5IO_6 are hypothetical, formulae of salts are given here without water of crystallisation —

$Ba_3(IO_4)_2$ is obtained, and $Ca(IO_3)_2$ undergoes a similar reaction. The salts obtained by double decomposition from alkali salts of the ortho-series, *c.* of the dimeso-series, usually belong either to the ortho- or meso-series, thus the following salts are obtained —

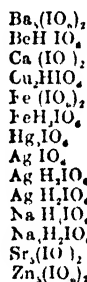
Starting from $Na_2H_2IO_4$. Starting from $K_2I_2O_8$.



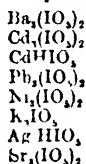
When a salt is obtained by ppn from an acidified solution of an alkali salt, the series to which the ppd salt belongs is conditioned by the relative quantity of acid present, thus when $AgNO_3$ is added to a solution of $Na_2H_2IO_4$, $K_2I_2O_8$ or KIO_3 , acidified by a little HNO_3 , Ag_2HIO_4 is obtained, when more HNO_3 is present the pp is $Ag_2H_2IO_4$, and when conc acid is pre-

PERIODATES

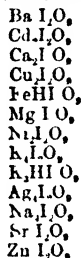
Ortho-periodates
derived from H_2IO_4 .



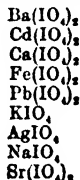
Meso-periodates
derived from H_2IO_4
($H_2IO_4 - H_2O = H_2IO_3$)



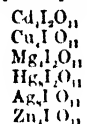
Dimeso-periodates
derived from $H_2I_2O_8$
($2H_2IO_4 - 3H_2O = H_2I_2O_8$)



Meta-periodates
derived from HIO_3
($HIO_3 - 2H_2O = HIO_2$)



Di-periodates
derived from $H_4I_2O_6$
($2H_2IO_4 - H_2O = H_4I_2O_6$)



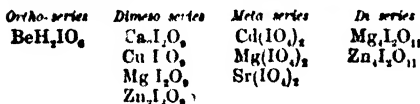
Dimeso-dimeta-periodates
derived from $H_4I_2O_6$
($2H_2I_2O_8 - 3H_2O = H_4I_2O_6$)



Trimeta-periodates
derived from $H_6I_3O_9$
($3H_2IO_3 - 10H_2O = H_6I_3O_9$)



The series to which a periodate belongs seems to depend partly on the nature of the base, and to a large extent on the conditions of preparation. By neutralising H_2IO_4 with $NaOH$ Ag Rammelsberg (P 134, 368, 499) obtained $NaIO_3$, by neutralising the same acid by various carbonates the following salts have been obtained —



By passing Cl into a hot mixture of KIO_3 and KOH , KIO_3 , $K_2I_2O_8$, and $K_2H_2IO_4$ are produced, but the Na salts formed under similar conditions are $Na_2H_2IO_4$, $Na_2H_2IO_4$, $Na_2I_2O_8$, and perhaps $NaIO_3$. By heating $Ba(IO_3)_2$ the salt

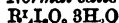
sent the salt $AgIO_3$ is obtained. Boiling salts of the ortho-, meso-, or dimeso-series with HNO_3 Ag as a rule produces salts of the meta-series. Acid salts of the ortho-series are changed to normal salts of the meso- or dimeso-series by heating, and acid salts of the meso-series are changed to normal salts of the dimeso-series, thus $Pb_2H_2(IO_4)_2$ at 275° gives $Pb_2(IO_4)_2$, and H_2O , and Ag_2HIO_4 gives $Ag_2I_2O_8$ and H_2O at 300° . Some of the periodates show very distinctly the difference between so called 'water of crystallisation' and 'water of constitution', thus Ag_2HIO_4 (which might be written $Ag_2I_2H_2O_6$) and $Ag_2I_2H_2O_6$ (which also might be written $Ag_2I_2H_2O_6$) are quite different bodies, the change $2Ag_2HIO_4 = Ag_2I_2O_8 + H_2O$ occurs at 300° , but $Ag_2I_2O_8 \cdot H_2O$ loses H_2O at 130° , again $Ag_2H_2IO_4$ (which might be written $Ag_2I_2H_2O_6$) is a dark red pp, obtained by adding $AgNO_3$ to $Na_2H_2IO_4$ in a slight excess of HNO_3 Ag,

whereas $\text{Ag}_2\text{I}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$ (which also might be written $\text{Ag}_2\text{I}_2\text{H}_2\text{O}_{12}$) forms light yellow crystals, and is obtained by treating $\text{AgIO}_3 \cdot \text{H}_2\text{O}$ with water. As a rule water of crystallisation is removed at 100° , whereas the temperature of acid salts must be raised to $270^\circ\text{--}300^\circ$ before change begins with evolution of water. (For a comparison of the classes of periodates, v. Kimmins, *C J* 55, 162.) Thomsen (*Th* 1, 244) has determined the heat of neutralisation by KOH of H_2IO_6 , with the following results:—

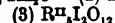
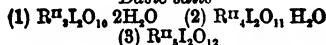
n	$[\text{HIO}^+\text{Aq}, n \text{ KOHAq}]$	ΔH
1	5,150	21,440
2	26,590	8,150
3	29,740	2,800
5	32,040	

From these results Thomsen concludes that H_2IO_6 is dibasic, but that basic salts are produced when more than 2KOH reacts. To explain the existence of many of the periodates, Thomsen prefers to double the formula of the acid and to represent it as normally tetrabasic, thus $\text{H}_4\text{I}_2\text{O}_{10} \cdot 3\text{H}_2\text{O}$. He classifies the chief periodates as follows:—

Normal salts



Basic salts



The existence of undoubted acid salts, i.e. salts in which the H is not combined with O in the form of H_2O , militates against the classification of Thomsen. The reaction of KOH with H_2IO_6 is almost certainly a complex occurrence.

In the following description of periodates only one or two salts in each series are described in detail.

The following are the principal memoirs on periodates, they are referred to in the following paragraphs by numbers. (1) Atterberg, *J* 1873 257, (2) Bengtsson, *A* 17, 250, (3) Blomstrand, *B* 3, 817, (4) Fernlund, *J* 1867 165, (5) Groth, *P* 134, 536, (6) Kimmins, *C J* 51, 356, 55, 148, (7) Langlois, *J* 1852 345, (8) Lautsch, *J pr* 100, 65, (9) Magnus, *A* Ammermuller, *P* 28, 514, (10) Philipp, *B* 3, 4, (11) Rammelsberg, *P* 44, 545, 46, 159, 62, 416, 90, 12, 115, 584, 125, 147, 134, 568, 499, 137, 305, (12) Ritter, *Gm-X* (6th ed.), 1 [2] 303.

Ortho-series, salts derived from H_2IO_6 .

Barium orthoperiodate $\text{Ba}_2(\text{IO}_3)_2$. Prepared by heating $\text{Ba}(\text{IO}_3)_2$ in a current of dry air (Sigura *a* Cross, *C J* 36, 118), by passing I vapour and dry air over heated BaO (S *a* C), by heating BaI_2 in a current of dry air so long as I is given off (S *a* C). Insol water, sol HNO_3 Aq, heated in H gives BaI_2 and BaO .

Silver orthoperiodates—1 AgIO_3 said to be ppd as a brown salt by adding AgNO_3 Aq to a neutral solution of an alkaline periodate, also by shaking $\text{Ag}_2\text{I}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$ with cold AgNO_3 Aq (11), Kimmins (6) failed to obtain this salt, sol in HNO_3 Aq and in NH_3 Aq—2 $\text{Ag}_2\text{H}_2\text{IO}_6$ dark red pp by adding AgNO_3 Aq to $\text{Na}_2\text{H}_2\text{IO}_6$ Aq or $\text{K}_2\text{I}_2\text{O}_3$ Aq in slight excess of HNO_3 (6)—

3 $\text{Ag}_2\text{H}_2\text{IO}_6$, slate coloured pp by adding AgNO_3 Aq to $\text{Na}_2\text{H}_2\text{IO}_6$ Aq or $\text{K}_2\text{I}_2\text{O}_3$ Aq in presence of a fair amount of HNO_3 (6).

Sodium orthoperiodates $\text{Na}_2\text{H}_2\text{IO}_6$ and $\text{Na}_2\text{H}_2\text{IO}_7$, white granular pps, formed by passing Cl into a bq ing solution of equal weights of NaIO_3 and NaOH , the second salt is less soluble than the first, from which it may be separated by long continued washing with small quantities of cold water (6).

For descriptions of the other salts of this series v (1) for Be salt, (6) for Cu and Fe salts, (8) for Hg salt, (11) for Ca, Sr, and Zn salts, (for formulae of salts, v p 23).

Meso series, salts derived from hypothetical H_2IO_5 .

Barium mesoperiodate $\text{Ba}_2(\text{IO}_3)_2 \cdot 6\text{H}_2\text{O}$. By ppg the corresponding K salt by Ba_2NO_3 Aq (11).

Potassium mesoperiodate $\text{K}_2\text{IO}_3 \cdot 4\text{H}_2\text{O}$. K_2IO_3 is prepared by saturating with Cl a hot conc solution of equal weights of KIO_3 and KOH , and repeatedly crystallising, the salt is dissolved in water and the solution is ppd by alcohol (11).

Silver mesoperiodate (acid salt) Ag_2HIO_5 . A dark brown pp by ppg $\text{Na}_2\text{H}_2\text{IO}_6$ or $\text{K}_2\text{I}_2\text{O}_3$ in just enough dilute HNO_3 Aq to form a solution (6).

For description of other salts of this series (formulae given on p 23), v (6) for salts of Pb and Ni, and (1*) for salts of Cd (crystallises with $5\text{H}_2\text{O}$) and Sr.

Dimeso series, salts derived from hypothetical H_2IO_4 .

Barium dimesoperiodate $\text{Ba}_2\text{I}_2\text{O}_6$. Obtained by ppg from alkali periodates in presence of HNO_3 . By neutralising H_2IO_6 Aq by BaO Aq, a salt with $7\text{H}_2\text{O}$ is obtained, which loses $4\text{H}_2\text{O}$ at 100° , and on strongly heating goes to BaIO_3 (11, 7, 8).

Ferric dimesoperiodate (acid salt) $\text{FeH}_2\text{I}_2\text{O}_6$. By ppg solution of $\text{Na}_2\text{H}_2\text{IO}_6$ or $\text{K}_2\text{I}_2\text{O}_3$ by FeCl_3 Aq and drying at 100° . Boiling with dilute HNO_3 Aq does not change this salt (6).

Potassium dimesoperiodates $\text{K}_2\text{I}_2\text{O}_6$ and $\text{K}_2\text{HI}_2\text{O}_6$. Both salts are obtained by passing Cl into KIO_3 mixed with KOH Aq (6). If equal weights of the KIO_3 and KOH are used, and the KIO_3 which separates is removed by filtration, the filtrate on concentration gives $\text{K}_2\text{I}_2\text{O}_6$, if this filtrate is exactly neutralised by HNO_3 the salt $\text{K}_2\text{I}_2\text{O}_6 \cdot 3\text{H}_2\text{O}$ (v also 11) crystallises out without concentration, if excess of HNO_3 is used, $\text{K}_2\text{HI}_2\text{O}_6$ is obtained. A solution of KIO_3 to which KOH is added is said to give triclinic crystals of $\text{K}_2\text{I}_2\text{O}_6 \cdot 9\text{H}_2\text{O}$ on concentration (5, 9). $\text{K}_2\text{I}_2\text{O}_6$ is soluble in KOH Aq, S 103 cold water, solution has alkaline reaction, Cl led into boiling $\text{K}_2\text{I}_2\text{O}_6$ Aq produces KIO_3 , with I, KI and KIO_3 are produced strongly heated leaves 2KI K_2O (11).

Silver dimesoperiodate $\text{Ag}_2\text{I}_2\text{O}_6 \cdot 3\text{H}_2\text{O}$. Light yellow crystals, obtained by treating $\text{AgIO}_3 \cdot \text{H}_2\text{O}$ with cold water (6, 9). Heated to 100° for 12 hours claret coloured crystals of $\text{Ag}_2\text{I}_2\text{O}_6 \cdot \text{H}_2\text{O}$ are formed, and when the temperature is raised to 180° and kept there for many hours a chocolate coloured powder, $\text{Ag}_2\text{I}_2\text{O}_6$, remains (6). When $\text{Ag}_2\text{I}_2\text{O}_6$ is treated with NH_3 Aq, $\text{Ag}_2\text{I}_2\text{O}_{11}$ is produced (11).

For descriptions of other salts of this series

(formulae given on p 23) v (2) for salt of Ca (crystallises with 7 and 9 H₂O), (6) for salts of Ni and Zn, (7) for Sr salt (crystallises with 3H₂O), (8) and (9) for Na salt (crystallises with 3H₂O), and (11) for salts of Cd (crystallises with 9H₂O), Cu (with 6H₂O), Mg (with 1 and 15 H₂O)

Meta series, salts derived from hypothetical HIO,

Ferric metaperiodate Fe(IO₃)₃. Bright yellow, produced by boiling FeH₂IO₆ for some time with conc HNO₃ Aq (6)

Potassium metaperiodate KIO₄. Prepared by saturating with Cl a boiling conc solution of equal parts KIO₃ and KOH, cooling, and crystallising repeatedly from water (6, 7, 9, 11) S 35 cold water, almost insol KOH Aq KIO₄ shows acid reaction At 300° forms KIO₃, decomposed by I at 160° Reduced in solution by KI to KIO₃, and I, which is then oxidised to KIO₄ (8, 9, 10)

Silver metaperiodate AgIO₄. By dissolving Na₂H₂IO₆ or Na₂H₂IO₆ or K₂H₂IO₆ in conc HNO₃ Aq and adding AgNO₃, also by treating Ag₂H₂IO₆, Ag₂H₂IO₆, or Ag₂H₂IO₆ with conc HNO₃ Aq and evaporating on steam bath Orange coloured crystals Crystals are AgIO₄, H₂O, heated to 130° for 6 hours AgIO₄ remains as a bright yellow powder (6) Decomposed by water giving Ag₂IO₄, 3H₂O (11)

For descriptions of other salts of this series (formulae given on p 23) v (11) for salts of Ba, Cd, Ca, and Sr (crystallises with 6H₂O), (5), (7), (8), (9), and (11) for salt of Na (crystallises with 2 and 3 H₂O)

Di series, salts derived from hypothetical H₂I₂O₁₁

Cadmium diperiodate Cd₂I₂O₁₁, 3H₂O By prep a slightly acidulated solution of NaIO₃ by a soluble Cd salt (11)

Silver diperiodate Ag₂I₂O₁₁. By treating Ag₂I₂O₁₁ with NH₃ Aq (8, 11)

For descriptions of the other salts of this series (formulae given on p 23) v (11) for salt of Cu (crystallises with H₂O and 7H₂O), (11) for salt of Mg (with 6 and 9 H₂O), (11) for salt of Hg (cf 8), (7) for salt of Zn (cf 11)

Dimeso di and Tri meso series, salts derived from hypothetical H₂I₂O₁₁, and H₃I₃O₁₅, respectively, v (8) and (11)

Iodine phosphides of, better called *Phosphorus iodides*, v PHOSPHORUS

Iodine selenides of, better called *Selenium iodides*, v SELENIUM

Iodine silicides of, better called *Silicon iodides*, v SILICON

Iodine sulphides of, better called *Sulphur iodides*, v SULPHUR

Iodine tellurides of, better called *Tellurium iodides*, v TELLURIUM M M P M

iodo acetic acid C₂H₃IO₂, s.e.

CH₂IC₂H₃ fol w 186 [82°]

Formation—1 By digesting bromo acetic ether with KI and alcohol, for about two hours in the dark, distilling off the resulting iodo-acetic ether and decomposing it with baryta-water (Perkin A Duppa, P M [4] 18, 54)—2 By boiling Ac₂O with iodine and iodic acid (Schützenberger, J pr 107, 108)—3. By the oxidation of iodo-acetic aldehyde (Chautard, A Ch [6] 16, 162)

Properties—Trimetric plates, decomposed by heat

Reactions—1 Moist silver oxide yields glycolic acid—2 Resolved by HI, even in the cold, into acetic acid and free iodine (Kekulé, C J 17, 207)

Salts The NH₄ and K salts are very soluble, crystalline, and non deliquescent—BaA', crystalline, m sol water, ppd from aqueous solution by alcohol—PbA', prisms In solution it easily splits up into PbI₂ and glycolic acid

Methyl ether MeA' (170°) Pungent liquid (Aronstein & Kramps, B 14, 604)

Ethyl ether EtA' (180°) From bromo-acetic ether, KI, and alcohol (P a D) Also from chloro acetic acid, KI, and alcohol (96 p a) by heating on the water bath in the dark Easily obtained by heating chloro acetic ether with CaI₂ 3 aq at 75° (Spindler, A 231, 272) Pungent heavy oil Decomposed by EtI at 230° giving acetic ether and ethylene iodide (Aronstein & Kramps, B 13, 489, 14, 604) With Me₂S it forms MeS CH₂ CO₂H and, finally, S(CH₂CO₂H)₂

Chloro ethyl ether ClCH₂CH₂OCOCH₃ S G 15 1954 From chloro ethyl chloro acetate by gently heating with an alcoholic solution of NaI (Henry, C R 97, 1308) Very pungent oil Turns brown in daylight

Propyl ether PrA' (198°) S G 15 1679 From propyl chloro acetate and alcoholic NaI (Henry, C R 100, 114) Pungent oil

Amide CH₂IC₂ONH₂ [158°] Formed from methyl iodo acetate and alcoholic NH₃ (Henry) Also from chloro acetamide and alcoholic KI, by standing a few days in the cold (Menschutkin a Fersolajeff, Z 1571, 5) Small needles or prisms

Nitrile CH₂IC₂NC *Iodo acetonitrile* (187°) S G 15 2307 From chloro acetonitrile and alcoholic NaI (Henry, C R 103, 413) Colourless, very pungent oil, which becomes brown in daylight V sol alcohol and ether Slightly decomposed on distillation, with liberation of some iodine and HCN AgOAc forms CH₂(OAc)CN

Di iodo acetic acid CHL₂CO₂H Formed by heating di bromo acetic ether with KI, converting the resulting di iodo acetic ether into a Ca salt with milk of lime, and adding HCl (Perkin A Duppa, C J 13, 1) Crystalline sulphur yellow compound, volatile in the cold, s sol water Does not blister the skin The K and Na salts are deliquescent—BaA', trimetric crystals, giving off iodine when heated—CaA', silky yellow needles—PbA', crystalline pp—AgA' yellow crystalline powder

Ethyl ether EtA' Formed by treating di chloro acetic ether with CaI₂ 3 aq at 75° (Spindler, A 231, 278) Formed also by adding iodine to a solution of diazo acetic ether in ether (Curtius, J pr [2] 38, 433) Oil, volatile with steam, reddens on exposure to air

Amide CH₂IC₂CONH₂ [202°] Formed by the action of iodine upon a cold alcoholic solution of diazo acetamide (Curtius, B 15, 1286) Formed also by the action of conc NH₃ Aq upon di-iodo-acetic ether Greenish-white prisms (from hot water), v sol water. Very stable towards acids and alkalis.

IODO ACETIC ALDEHYDE $\text{CH}_2\text{I CHO}$ S G

12 2 14

Formation—1 By boiling an aqueous solution of chloroacetic aldehyde with KI, allowing to stand for 12 hours, and then extracting with ether (Glinksky, *Z* [2] 4, 618)—2 By the oxidation of a solution of iodine in alcohol

Preparation—By mixing iodine (50 g), iodic acid (20 g), aldehyde (30 cc) and water (60 cc) and leaving the closed flask to stand for about a week (in summer) Water (500 cc) is then added, when the iodoacetic aldehyde separates as an oil (Chautard, *A Ch* [6] 16, 145, *C R* 102, 118, *C N* 54, 87, cf W P Bloxam & E F Herroun, *C N* 53, 301, 54, 133)

Properties—Limpid colourless oil, uninflammable. Blackens rapidly when exposed to light. Its vapour is excessively irritating. Cannot be distilled even *in vacuo*. Decomposes at 80°. Miscible with alcohol, ether, chloroform, and CS_2 . Soluble in water. Forms a crystalline hydrate. Forms a crystalline compound with NaHSO_4 .

Reactions—1 Dilute aqueous KOH forms iodoform. Conc KOH aq resinifies it.—2 *Ammonia* either aqueous or alcoholic forms ammonium iodide and various bases, e.g. oxytriethylamine ($\text{C}_2\text{H}_5\text{O}$), $\text{N C}_2\text{H}_5\text{OH}$ —3 *Chlorine and bromine* at ordinary temperatures displace iodine giving chloro and bromoacetic aldehydes and resins.—4 All mineral acids decompose iodoacetic aldehyde.—5 *Reducing agents* (Na amalgam, Zn and HCl) remove the iodine.—6 *Oxidation* with HNO_3 yields $\text{CH}_2\text{I CO}_2\text{H}$ [82°]—7 NaOEt and KOEt yield only resinous matters.—8 *Silver acetate* yields acetic ether (74°)—9 AgCN gives $\text{CH}_2(\text{CN})\text{CO}_2\text{H}$ —10 AgSCN similarly gives $\text{CH}_2(\text{SCN})\text{CO}_2\text{H}$ —11 *The amines* form crystalline bases with elimination of water. Thus aniline gives $\text{CH}_2\text{I CH}(\text{NH}_2\text{C}_6\text{H}_5)$, and *p*-toluidine gives $\text{CH}_2\text{I CH}(\text{NH}_2\text{C}_6\text{H}_4\text{CH}_3)$.

Tri-iodoacetic aldehyde CHI_3CHO *Iodal* (above 200°). From chloral or bromal and HI (Bertrand, *J* 1881, 588)

IODO-ACETO ACETIC ETHER

$\text{CH}_3\text{CO CH}_2\text{CO}_2\text{Et}$ S G 14 17053 From cupric acetoacetic ether and iodine in ether (Schönbrodt, *A* 253, 178). Yellowish oil, v sol ether. Begins to decompose at 25°. AgCl gives chloroacetoacetic ether. AgCy gives di-oxyterephthalic ether dihydride. AgNO_3 forms nitroacetoacetic ether. Reduced silver gives diacetyl-fumaric ether. Sodium acetoacetic ether in benzene forms diacetyl succinic ether.

IODO ACETONE $\text{C}_2\text{H}_5\text{IO}$ i.e. $\text{CH}_3\text{CO CH}_2\text{I}$ S G, 12 2 17 Formed by the action of KI on chloroacetone (Glutz & E Fischer, *J pr* [2] 4, 52). Prepared by mixing acetone (200 cc) with iodine (100 g) and iodic acid (10 g), and, after a week, boiling for three hours, adding water (500 cc), drying the separated oil over CaCl_2 , and distilling *in vacuo* in the dark (De Clermont & Chautard, *C R* 100, 745). Very pungent oil. Not inflammable. Decomposes slowly in sunlight, but more rapidly when heated. Reduces Fehling's solution. Mineral acids convert it into $\text{CH}_2\text{I CO CH}_2\text{I}$. Silver chloride forms chloroacetone. LiOAc gives $\text{CH}_3\text{CO CH}_2\text{OAc}$.

Di iodoacetone $\text{C}_2\text{H}_4\text{I}_2\text{O}$ i.e. $\text{CH}_2\text{I CO CH}_2\text{I}$ [62°] (M Simpson, *J pr* 102, 380, O Völker, *A* 192, 90).

Formation—1 From acetone (48 g), water (1000 g), and ICl_3 (96 g) at 68°. As soon as reaction begins the liquid is cooled. The oil that separates is exposed to the air till crystals separate (eight weeks), these are recrystallised from alcohol.—2 From $\text{CH}_2\text{I CO CH}_2\text{I}$ and aqueous KI.

Properties—White needles. Violently attacks the mucous membrane. Decomposed when heated. Not very soluble in chloroform, CS_2 , or alcohol. Soluble in benzene, v sol ether and acetone. Cannot be distilled undecomposed.

Reactions—1 *Silver chloride* gives solid $\text{CH}_2\text{I CO CH}_2\text{I}$ [43°]—2 Decomposed by cold KOH, and by boiling K_2CO_3 —3 Not converted into acrolein or acrylic acid by Ag_2O , AgCy , or HgO —4 HgI_2 forms acetone, iodic acetone, HI, and acetic acid, but no acrolein—5 *Zinc* and *acetic acid* reduce it to acetone.

p IODO ACETOPHENONE

[14] $\text{C}_6\text{H}_4\text{I CO CH}_3$ [79°] Obtained by diazotising *p*-amidoacetophenone, and heating the diazo-chloride with excess of HI (Klingel, *B* 18, 2692). White flat needles or plates. Easily sol alcohol and ether, and to a certain extent in hot water. By CrO_3 it is easily oxidised to *p*-iodo benzoic acid [266°].

IODO-ACETOTHENONE v Iodo THIENYL**MPHTHYL KETONE**

IODO ACETYLENE HC CI Formed by passing a current of steam through a solution of barium iodo-propargylate ($\text{C}_3\text{H}_3\text{O}$) $_2\text{Ba}$ (Bayer, *B* 18, 2271). Crystalline solid. Sol water. Volatile with steam. Very poisonous. It gives a purple red pp with ammoniacal Cu_2Cl_2 , which is soon converted by an excess of the copper solution into copper acetylene and cuprous iodide. It soon polymerises on keeping.

Di iodoacetylene I C CI [78°] Formed by the action of iodine upon acetylene silver. By treatment with an excess of ammoniacal Cu_2Cl_2 it is converted into copper acetylene and cuprous iodide. It readily polymerises (Bayer, *B* 18, 2275).

Di iodo diacetylene I C C C I [101°] Colourless crystals. Odour resembling iodoform. Formed by the action of a solution of iodine in aqueous KI upon silver diacetylene. By treatment with an excess of ammoniacal Cu_2Cl_2 it yields copper diacetylene and cuprous iodide. It explodes violently on heating, with a brilliant red flash. It soon polymerises on keeping (Bayer, *B* 18, 2276).

IODO ACETYLENE CARBOXYLIC ACID**IODO PROPIONIC ACID****IODO ACIDS** v IODO COMPOUNDS

IODO ACRYLIC ACID $\text{C}_3\text{H}_3\text{IO}_2$ Large, colourless, four sided prisms [65°], or plates [140°]. Easily soluble in water, alcohol, and ether. Formed by the addition of HI to propiolic acid (Bandrowski, *B* 15, 2703, Stolz, *B* 19, 542). The Pb and Ag salts are white crystalline pps.

Di iodo-acrylic acid $\text{CHI CI CO}_2\text{H}$ [106°]. Formed by treating propiolic acid with an ethereal solution of iodine (Homolka & Stolz, *B* 18, 2284). Colourless prisms, or long needles. Volatile with steam. Easily soluble in alcohol, ether, and hot water.

Di iodo-acrylic acid $\text{I C CH CO}_2\text{H}$ [133°]. Formed by addition of HI to iodo propiolic acid.

$\text{IO}_2\text{C.CO}_2\text{H}$ (Homolka & Stolz, *B* 18, 2284)
Prisms Easily soluble in alcohol and ether, tolerably in hot water, very sparingly in cold Not volatile with steam

Tri-iodo-acrylic acid $\text{Cl}_3\text{C.CO}_2\text{H}$ [207°]
Formed by the action of an ethereal solution of iodine upon iodo-propionic acid $\text{ClC.CO}_2\text{H}$ (Homolka & Stolz, *B* 18, 2286) Large colourless prisms Easily soluble in alcohol and ether, insoluble in cold water

iodo-di allyl-acetic acid $\text{C}_6\text{H}_9\text{IO}_2$ *i.e.* $\text{Cl}(\text{C}_6\text{H}_7)\text{CO}_2\text{H}$ Iodo octenoic acid From $\text{C}(\text{OH})(\text{C}_6\text{H}_7)\text{CO}_2\text{H}$ and fuming HIAq (Schatzky, *J R* 17, 78) Crystals, insol water, v sol alcohol and ether Easily decomposes

iodo allyl alcohol $\text{C}_3\text{H}_5\text{IOH}$ [160°]
Produced by the action of aqueous Na_2CO_3 on di iodo propyl alcohol (Hübner & Lellmann, *B* 18, 460) Needles Highly volatile with steam Insol water, sol alcohol, chloroform, and HOAc Not attacked by Ac_2O

iodo allylene $\text{C}_3\text{H}_4\text{I}_2$ *i.e.* $\text{CH}_2\text{C}(\text{Cl})_2$ (?) (98°) SG 17 From silver allylene and iodine in KIAq (Liebermann, *A* 135, 270) Pungent oil Almost insol alcohol With zinc and HCl it gives off allylene Combines with iodine (1 mol)

Iodo allylene $\text{CH}_2\text{I.CCH}$ Propargyl iodide (115°) SG 2 20177 From propargyl bromide and alcoholic NaI (Henry, *B* 17, 1132) Liquid V e sol alcohol and ether, m sol cooling water Readily combines with iodine, forming colourless needles of $\text{CHI.CI.CH}_2\text{I}$ [41°] Combines with mercury

iodo allylene iodide v **Tri-iodo propylene**

Tri-iodo-allyl ethyl oxide $\text{C}_6\text{H}_9\text{IO}_3$ *i.e.* $\text{Cl}_3\text{C.CI.CH}_2\text{OEt}$ From the silver derivative of propargyl ethyl oxide and iodine in ethereal solution (Liebermann, *A* 135, 285) Oil

Di iodo allyl iodide v **Tri iodo propylene**

Tetra-iodo-di-allyl phosphite
 $(\text{CHI.CI.CH}_2\text{O})_2\text{P}(\text{OH})$ [49°] From propargyl alcohol, iodine, and red phosphorus (Henry, *B* 8, 398, 17, 1133) Long slender needles (from alcohol) Very pungent

iodo-amido-benzoic acid $\text{C}_6\text{H}_4\text{INO}_2$ *i.e.* $\text{C}_6\text{H}_4(\text{NH}_2)\text{CO}_2\text{H}$ [3 2or6 1] [137°] Formed by reducing iodo nitro benzoic acid [235°] dissolved in glacial acetic acid (Grothe, *J pr* [2] 18, 326) Dark brown crystals V sol water By further reduction it forms o-amido benzoic acid [143°] — HA'Cl — BaA' , aq

Iodo-amido-benzoic acid $\text{C}_6\text{H}_4(\text{NH}_2)\text{CO}_2\text{H}$ [8 6or2 1] [209°] From iodo nitro benzoic acid [174°] Needles, sl sol water May be reduced to o-amido benzoic acid — CaA' , 2aq — BaA' , — BaA' .

Di-iodo-m-amido benzoic acid
 $\text{C}_6\text{H}_3\text{I}_2(\text{NH}_2)\text{CO}_2\text{H}$ Formed, together with $\text{N}(\text{C}_6\text{H}_4\text{I.CO}_2\text{H})_2$, by treating an alcoholic solution of m-amido benzoic acid [173°] with iodine and mercuric oxide (Beydick, *B* 8, 384) The acids are separated by fractional ppn of their alcoholic solution by lead acetate Long needles Decomposed by heat Insol water, v sol alcohol, ether, aqueous acids, alkalis, and Na_2CO_3 — KA' long silky needles, obtained by mixing alcoholic solutions of the acid and KOH .

Di-iodo-p-amido-benzoic acid
 $\text{C}_6\text{H}_3\text{I}_2(\text{NH}_2)\text{CO}_2\text{H}$ [above 300°] Obtained by dissolving p-amido benzoic acid in dilute HCl and passing in the vapour of ICl (2 mols) (Michael & Norton, *Am* 1, 264) Tables Insol water, alcohol, and HOAc , sol nitro-benzene Does not combine with acids — NaA' , 5aq long white needles, sol hot water. — BaA' , 4aq needles, sol hot water — AgA'

DI IODO-p AMIDO PHENOL
 $\text{C}_6\text{H}_3\text{I}_2(\text{NH}_2)\text{OH}$ [222°] Formed by reducing the corresponding nitro phenol with SnCl_2 and HCl (R Seifert, *J pr* [2] 28, 437) Needles (from alcohol) or plates (from MeOH) With HCl and bleaching powder it gives di-iodo quinone chloride (*q v*) With H_2SO_4 and $\text{K}_2\text{Cr}_2\text{O}_7$ gives di-iodo quinone (*q v*) — B'Cl Decomposed by water

iodo AMIDO THYMOL
 $\text{C}_9\text{HIMEPr}(\text{OH})(\text{NH})$ [6 5 2 1 4] From the oxim of iodo thymoquinone by reduction with SnCl_2 (Kehrmann, *J pr* [2] 39, 392) Its stannous chloride forms large colourless prisms

iodo AMIDO-TOLUENE SULPHONIC ACID
 $\text{C}_6\text{H}_4\text{MeI}(\text{NH}_2)\text{SO}_3\text{H}$ [1 4 2 5] Formed from the corresponding nitro toluidine sulphonic acid by diazotisation and treatment of the product with conc HIAq at 135° (Foth, *A* 230, 308) Slender silky needles (from water) V sl sol cold water

iodo AMYL ALCOHOL $\text{C}_5\text{H}_{11}\text{IO}$? Amylene glycol iodhydrin Formed by shaking un-amylene with iodine, water, and HgO (Lippmann, *Z* 1867, 17, 4 *Suppl* 5, 124) Heavy oil, decomposed by distillation

iodo AMYLIDENE ANILINE
 $\text{C}_6\text{H}_4\text{ICH}=\text{C}_6\text{H}_5$ Formed by heating iodo iso valeric aldehyde with aniline (Chautard, *A Ch* [6] 16, 168) Yellow prismatic needles or rectangular tables Decomposed by heat V sol alcohol, sol water and ether, v sl sol benzene and chloroform With HCl it forms an uncrystallisable salt

o IODO ANILINE $\text{C}_6\text{H}_4\text{IN}$ *i.e.* $\text{C}_6\text{H}_3\text{I}(\text{NH}_2)$ [1 2] Mol w 219 [56 5°] Formed by heating at 100° a mixture of o-iodo nitro benzene (25 pts) with ferrous sulphate (250 pts) and a moderate excess of dilute ammonia The product is extracted with ether, the extract distilled with steam, the product dissolved in dilute H_2SO_4 , freed from iodo nitro benzene by shaking with ether, ppd by ammonia and the iodo aniline again distilled with steam (Korner & Wender, *G* 17, 486) Long silky needles, with an odour resembling pyridine Sl sol hot water, v sol other solvents Exhibits an alkaline reaction. Turns brown on exposure to air and light Decomposes when heated, evolving iodine Iodine converts it into di-iodo aniline [96°]

Salts — B'Cl aq small prisms or cubes, turning opaque on exposure, with loss of water of crystallisation — B' , $(\text{H}_2\text{SO}_4)_2$, silky needles, m sol water — Nitrate small four sided tables

Acetyl derivative $\text{C}_6\text{H}_3\text{INHAc}$ [110°] Prisms or hexagonal tables, m sol hot water, v sol alcohol

m-Iodo-aniline $\text{C}_6\text{H}_4(\text{NH}_2)$ [1 3] [27°] Formed by reducing m-iodo nitro-benzene (Gress, *Z* 1866, 218) Silvery plates

Acetyl derivative $\text{C}_6\text{H}_4(\text{NHAc})$ [119 5°]

Glistening needles; more stable than the *o*-isomeride (Körner & Wender, *G* 17, 486)

p-Iodo-aniline $C_6H_4I(NH_2)$ [14] [83°]
Formed by reducing *p* iodo-nitro-benzene [172°] (Grüss, *C* J 20, 85) Formed also by adding iodine (3 pts) to aniline (2 pts) and mixing the solution with aqueous HCl (S G 111) when *p*-iodo aniline hydrochloride is ppt (Hofmann, *A* 67, 64) Prisms or needles, heavier than water Has no action on litmus Sl sol cold water, v sol other solvents Bromine converts it into tri-bromo-aniline — B'HCl thin laminae or needles (from hot water), sl sol cold water, almost insol HClAq, sol alcohol, insol ether — $B'_2H_4PtCl_6$, orange pp — $B'_2H_4C_2O_4$, long needles, sl sol water and alcohol, insol ether — $B'_2H_4SO_4$, scales

Acetyl derivative $C_6H_4I(NHAc)$ [183°]
S (alcohol of 95 p c) 6.4 at 20.5° Small glistening tables or trimetric prisms (K & W) Formed by dissolving acetanilide in glacial acetic acid, and passing in vapour of iodine chloride, the yield being 85 p c (Michael & Norton, *Am* 1, 255)

Benzoyl derivative $C_6H_4I(NHBz)$ [180°]
Long needles (Hubner, *B* 10, 1717) An isomeride [210°] is formed from benzanilide and ICl

Di iodo aniline $C_6H_3I_2NH_2$ [4 2 1] [96°]
Formed by the action of iodine on phenylated 'white precipitate' NHPH HgCl (Rudolph, *B* 11, 78) Formed also by passing ICl (2 mols) into a solution of aniline in HOAc (Michael & Norton, *Am* 1, 255, *B* 11, 109) Needles Sl sol boiling water and ligroin, v sol ether, chloroform, CS₂, acetic ether, and hot alcohol Volatile with steam It is slightly basic, but its hydrochloride is decomposed by cold water — B'HCl: long white needles, decomposed at 50°. — $B'_2H_4PtCl_6$ — B'_2HNO_3 — $B'_2(H_2SO_4)_2$

Benzoyl derivative $C_6H_3I_2NHBz$ [181°]
Slender needles (Rudolph)

Tri-iodo-aniline $C_6H_2I_3(NH_2)$ [6 4 2 1] [185.5°]
Prepared by the action of ICl (3 mols) on a solution of aniline in HClAq (Michael & Norton, *Am* 1, 255) Long white needles, sol CS₂ and acetic ether, insol water

IODO-ANISIC ACID *v* *Methyl derivative of Iodo oxy-benzoic acid*

IODO-BENZENE C_6H_4I *Phenyl iodide*
Mol w 204 (188°) S G § 18606, $\frac{15.2}{1}$ 18380 (Young, *C* J 55, 486), $\frac{2}{1}$ 18578, $\frac{1}{2}$ 18403, $\frac{3}{2}$ 18321 S V 130 55 (R Schiff, *B* 19, 564) $n_D^{20} = 1.6189$ (Seubert, *B* 22, 2520).

Formation — 1 By treating phenol with iodine and phosphorus The yield is bad (Williamson & Scruggam, *C* J 18, 244) — 2 By the action of ICl on sodium benzoate (Schützenberger, *C. R* 52, 963) — 3 By heating benzene for some time with iodic acid (Peltzer, *A* 186, 194) — 4 By the action of HI on diazobenzene salts (Grüss, *J* 1866, 447) — 5 By heating benzene (20 g) with iodine (15 g) and iodic acid (10 g), at 220° (Kekulé, *A* 187, 157) — 6 From benzene, iodine and FeCl₃ (Lothar Meyer, *A* 281, 195) — 7 By the action of excess of iodine on phenyl hydrazine (E v Meyer, *J. pr.* [2] 86, 115)

Preparation — By allowing chloride of iodine to drop slowly into a large excess of benzene

containing a small quantity of aluminium chloride (Greene, *C. R* 9f, 40)

Properties — Colourless oil, insol water Solidifies when cooled by solid CO₂. Easily reduced by sodium amalgam to benzene Aqueous HIAq at 250° reduces it to benzene Not affected by solid KOH at 250°, nor by alcoholic KOH or NH₃ (Kekulé)

Reactions — 1 H₂SO at 100° forms a mixture of iodo-benzene sulphonic acid, di iodo-benzene, and benzene sulphonic acid, in proportions varying with proportions of materials employed, strength of the acid, temperature, and duration of the experiment (Neumann, *A* 241, 47) — 2 *Silver nitrate* does not act below 135°, but between 140° and 150° reaction takes place forming tri-nitro phenol, AgI, and metallic silver (Geuther, *A* 245, 99) — 3 By heating with *aluminium chloride* there is formed benzoene, di-iodo benzenes, HCl, and iodine (Dumreicher, *B* 15, 1868)

Dichloride $C_6H_4I_2Cl_2$. Formed by passing chlorine into liquid iodobenzene Formed also by passing chlorine into a solution of iodobenzene (5 g) in chloroform (15 g) (Willgerodt, *J. pr* [2] 83, 154) Begins to decompose at 80°, and at 120° it is all broken up into Cl₂ and C₆H₅I It may be kept in glass bottles, but if placed over H₂SO, it gives off chlorine It is decomposed by solution in alcohol, but it may be crystallised from chloroform as yellow needles It dissolves in benzene, glacial acetic acid, light petroleum, CS₂, and ether As a reagent it displaces iodine by chlorine, turning KI, PbI₂, &c, into KCl PbCl₂, &c, with separation of iodine May therefore be used as a test for iodides It also converts alcoholic iodides (e.g MeI) into chlorides It is hardly attacked by cold aqueous NH₃ or NaOH

o Di iodo benzene $C_6H_3I_2$ [1 2] [27°] (287°)

Preparation — Iodo aniline (10 g) is dissolved in hot water (70 g) containing H₂SO₄ (9 g), KNO₃ (4.5 g) dissolved in a little water is added, and then an excess of solution of KI in aqueous HI (S G 11) After standing a few hours the oil that has separated is washed with KOHAq and distilled with steam (Körner & Wender, *G* 17, 486, cf Körner, *G* 4, 305) Long prisms or hexagonal tables Volatile with steam Sl sol water, sol alcohol

m Di iodo benzene $C_6H_3I_2$ [1 3] [40°] (K), [37°] (R) (285°) From *m* iodo aniline by displacing NH₂ by I (Körner) Also from di iodo aniline by elimination of NH₂ (Rudolph) Trimetric tables (from ether-alcohol)

p-Di-iodo-benzene $C_6H_3I_2$ [1 4] [127°] (Kekulé), [129°] (Körner) A product of the action of iodine chloride on NaOBz (Schützenberger) and of iodine and iodic acid on benzene (Kekulé) Formed also from *p* iodo-aniline by the diazo-reaction (Kekulé, *Z* 1866, 688). Nacreous laminae Readily sublimed

Tri-iodo-benzene $C_6H_2I_3$ [1 2 4] Mol w 456. [76°]. A product of the action of iodine and iodic acid on benzene (Kekulé). Small needles May be sublimed

IODO-BENZENE-AZOXY-COMPOUNDS *v* **AZOXY-COMPOUNDS.**

IODO-BENZENE *o*-SULPHONIC ACID
 $C_6H_3I_2SO_3H$ [1:2] From *o*-amido-benzene sulphonic acid by the diazo-reaction (Bahlmann, *A*.

186, 825) — KA' :aq: crystals, sl. sol. water — BA'_2 , needles, sl. sol. cold, v. sol. hot, water
Chloride $\text{C}_6\text{H}_4\text{I SO}_2\text{Cl}$ [51°] Thick prisms (from ether)

Amide $\text{C}_6\text{H}_4\text{I SO}_2\text{NH}_2$. [170°]. White laminae, sl. sol. water

Iodo-benzene p sulphonic acid $\text{C}_6\text{H}_4\text{ISO}_3\text{H}$ [14] From iodo benzene and fuming H_2SO_4 (Körner a Paterno, *G* 2, 448) Formed also from amido benzene *p* sulphonic acid by displacing NH_2 by *I* through the diazo-reaction (Lenz, *B* 10, 1135) Deliquescent needles — $\text{NH}_4\text{A}'$. minute needles — KA' needles — CaA'_2 — BaA'_2 . minute plates, sl. sol. water — PbA'_2 .

Chloride $\text{C}_6\text{H}_4\text{ISO}_2\text{Cl}$ [87°] Laminae

Amide $\text{C}_6\text{H}_4\text{I SO}_2\text{NH}_2$. [183°]. Crystalline powder, sl. sol. water, v. so. alcohol

o IODO-BENZOIC ACID $\text{C}_6\text{H}_4\text{I CO}_2\text{H}$ [157°]

Formation — 1 From *o* amido benzoic acid by the diazo reaction (Griess, *C J* 24, 702) — 2 By oxidising *o* iodo toluene with dilute HNO_3 (Kekulé, *B* 7, 1007) — 3 From *m* iodo nitro benzene and alcoholic KCy at 200° (Richter, *B* 4, 554)

Properties — Long needles, may be readily sublimed. Sl. sol. hot water, v. e. sol. ether and alcohol. Gives salicylic acid when fused with potash

Salts — CaA'_2 , 2aq — BaA'_2 , 6aq

m Iodo benzoic acid $\text{C}_6\text{H}_4\text{I CO}_2\text{H}$ [187°]

Formation — 1 From *m* amido benzoic acid by the diazo reaction (Griess, *A* 113, 334, 117, 1, Cunze a Hübner, *A* 135, 108, Grothe, *J pr* [2] 18, 324) — 2 By heating benzoic acid (1 pt) with KIO_3 (2 pts) and dilute H_2SO_4 (Peltzer, *A* 136, 201) — 3 By oxidising *o* iodo toluene with chromic acid mixture (Körner, *Z* [2] 5, 637) — 4 By heating dry silver benzoate with iodine at 150°–180° (Birnbäum a Reinherz, *B* 15, 456)

Properties — Needles, sl. sol. water, v. sol. alcohol. May be sublimed. Gives *p* oxy benzoic acid on oxidation. Ammonia forms amido benzoic acid

Salts — NaA' , aq — MgA'_2 , 4aq — CaA'_2 , 2aq scales — CaA'_2 , nodules — BaA'_2 , 4aq needles, sol. alcohol. — MgA'_2 , 4aq

Ethyl ether EtA' Oil

Nitrile $\text{C}_6\text{H}_4\text{I CN}$ [41°] From *m*-amido benzonitrile by the diazo reaction. Needles

p Iodo-benzoic acid $\text{C}_6\text{H}_4\text{I CO}_2\text{H}$ [266°] (Beran, *B* 18, 137, Klingel, *B* 18, 2693) Formed by oxidation with chromic acid mixture from *p* ic lo-toluene (Körner, *Z* [2] 5, 327), from *p* iodo-phenyl acetic acid (Jackson a Mabery, *Am* 2, 253), or from *p* iodo acetophenone (Klingel) Nacreous scales (from alcohol) Nearly insol. boiling water. Converted by potash fusion into *p* oxy benzoic acid

Salts (Glassner, *B* 8, 562) — NaA' , aq colourless needles, v. sol. water — KA' — BaA'_2 , 1½aq. long trimetric plates — CaA' , aq — SrA'_2 , aq nacreous laminae. — ZnA'_2 , 4aq cubes

Methyl ether MeA' [114°] Long needles (Schmidt a Schultz, *A*, 207, 883)

Ethyl ether EtA' Oil

o IODO-BENZOIC ALDEHYDE $\text{C}_6\text{H}_4\text{I CHO}$ [87°] Formed from *o*-nitro cinnamic acid by successive conversion into amido-, diazo-, and iodo cinnamic acid, and oxidation of the latter with KMnO_4 (Stuart, *C J*, 53, 140).

p-Iodo-benzoic aldehyde $\text{C}_6\text{H}_4\text{I CHO}$. [73°]. Needles (Jackson a. White, *B* 11, 1042, *P Am A* 15, 269)

p-IODO-BENZYL ALCOHOL $\text{C}_6\text{H}_4\text{I CH}_2\text{OH}$ [72°] Prepared by heating *p*- iodo benzyl bromide with NaOAc and aqueous NH_3 at 160° Formed also by heating *p* iodo benzyl bromide with water for a long time (Jackson, *P Am A* 13, 202, Jackson a. Mabery, *Am* 2, 251, *B* 11, 56) Silky scales (from alcohol or CS_2) or long needles (from water) Sl. sol. cold water, v. sol. alcohol, ether, benzene, and CS_2 .

o IODO-BENZYLAMINE $\text{C}_6\text{H}_4\text{I NH}_2$

[21] $\text{C}_6\text{H}_4\text{I CH}_2\text{NH}_2$ From *o* iodo benzyl bromide and alcoholic NH_3 (Mabery a Robinson, *Am* 4, 103) Liquid. Absorbs CO from the air — $\text{B}'\text{H}_2\text{PtCl}_6$ minute yellow prisms

p Iodo benzylamine [41] $\text{C}_6\text{H}_4\text{I CH}_2\text{NH}_2$

Formed by heating *p* iodo benzyl bromide with alcoholic NH_3 at 120° (Jackson a. Mabery, *Am* 2, 257) Oil. Absorbs CO from the air forming a carbonate [113°] — $\text{B}'\text{HCl}$ slender white needles [240°], sol. water and alcohol, sl. sol. ether — $\text{B}'\text{H}_2\text{PtCl}_6$

Di-p-iodo-di benzyl amine $(\text{C}_6\text{H}_4\text{I CH}_2)_2\text{NH}$

[76°] Formed, together with tri *p* iodo tri benzylamine, by boiling *p* iodo benzyl bromide with alcoholic NH_3 (Jackson a. Mabery, *Am* 2, 256, *B* 11, 58, *P Am A* 13, 209) White needles, insol. water, v. sol. hot alcohol, ether, benzene, and CS_2

Salts — $\text{B}'\text{HCl}$ thick white plates, sol. CS_2 and HOAc , sl. sol. alcohol and benzene — $\text{B}'\text{HBr}$ thick pearly prisms, insol. water, sl. sol. alcohol sol. ether, benzene, and CS_2 — $\text{B}'\text{H}_2\text{PtCl}_6$ minute yellow needles, almost insol. water and alcohol — $\text{B}'\text{H}_2\text{CO}$, [113°] Crystalline

Tri-*p* iodo-tri-benzyl-amine $(\text{C}_6\text{H}_4\text{I CH}_2)_3\text{N}$

[115°] Formed as above (J. a. M.) White needles (from ether), v. sl. sol. hot alcohol, v. sol. ether, benzene, and CS_2 . — $\text{B}'\text{H}_2\text{PtCl}_6$ yellow needles, nearly insol. water and alcohol

o IODO BENZYL BROMIDE $\text{C}_6\text{H}_4\text{I CH}_2\text{Br}$

[53°] Prepared by dropping bromine into *o* iodo toluene heated to 190°–200° (Mabery a Robinson, *Am* 4, 102, *P Am A* 17, 103) Thick flattened prisms (from ligroin), v. sol. ether, hot alcohol, benzene, CS_2 , and chloroform, insol. water. Its vapour is very pungent. Gives *o* iodo benzoic acid on oxidation with dilute HNO_3 .

p Iodo benzyl bromide $\text{C}_6\text{H}_4\text{I CH}_2\text{Br}$ [79°]

Prepared by heating *p* iodo toluene in bromine vapour at 115°–150° (Jackson, *Am* 1, 93) Straw coloured needles, somewhat pungent. Insol. water and cold alcohol, sol. hot alcohol, v. sol. ether. Hardly attacked by CrO_3 .

IODO-BENZYL CYANIDE *o* Nitrile of Iodo-benzyl-acetic acid

o IODO BENZYLIDENE-MALONIC ACID

$\text{C}_6\text{H}_4\text{IO}$, *o* $\text{C}_6\text{H}_4\text{I CH}(\text{CO}_2\text{H})_2$ [204°] Formed by heating equal weights of malonic acid and iodo benzoic aldehyde for several hours at 100° with half their weight of HOAc (Stuart, *C J* 53, 142) Decomposed on melting into CO_2 and *o*-iodo cinnamic acid

o-IODO-BENZYL-MALONIC ETHER

$\text{C}_6\text{H}_4\text{CH}_2\text{Cl}(\text{CO}_2\text{Et})_2$ From sodium benzyl malonic ether and iodine (Bischoff a. Hausdöfler, *A*, 239, 110). Oil. Decomposes on hydrolysis

into benzoic aldehyde, alcohol, acetic acid, and CO_2 .

***p*-IODO BENZYL SULPHOCYANIDE**

$\text{C}_6\text{H}_4\text{I} \cdot \text{SCy}$ [40°] Formed by boiling *p* iodo-benzyl bromide with an alcoholic solution of potassium sulphocyanide (Jackson, *P Am A* 18, 207, *B* 11, 58) Long white plates (from alcohol) *M* sol hot alcohol, *v* sol ether, benzene, CS_2 , and HOAc

IODO BETORCIN $\text{C}_8\text{HIME}(\text{OH})_2$ [93°]

Formed by the action of PbO and *I* (2 pts) on an ethereal solution (50 pts) of betorcin (1 pt) Crystallised from light petroleum (Stenhouse & Groves, *C J* 37, 404) *V* sol ether, CS_2 , and benzene

IODO BROMO-*v* BROMO IODO-

IODO BUTANE *v* BUTYL IODIDES

Di iodo butane $\text{C}_4\text{H}_2\text{I}_2$ *se*

$\text{CH}_3\text{CH}_2\text{CHI}_2\text{CH}_3$ (116° *in vacuo*) *SG* 2 291 From the corresponding di oxy butane and *HI* (Wurtz, *Bl* [2] 41, 362)

***p* IODO-ISOBUTYL BENZENE $\text{C}_6\text{H}_5(\text{C}_4\text{H}_9)\text{I}$**

(256° *cor*) Formed by the action of *HI* on the diazo compound from *p* amido phenyl butane (Pahl, *B* 17, 1232) Colourless oil By HNO_3 , it is oxidised to *p* iodo benzoic acid

IODO BUTYLENE $\text{C}_4\text{H}_6\text{I}$ Crotyl iodide

(132°) From tri oxy butane (butenyl glycerin), iodine, and phosphorus (Lieben & Ziesel, *M* 1, 836) Pungent liquid Combines with mercury forming an unstable crystalline compound

IODO-ISOBUTYL TOLUENE

$\text{C}_6\text{H}_5(\text{CH}_3)(\text{C}_4\text{H}_9)\text{I}$ [136] [*c* 35°] (264°) Formed by diazotising amido tolyl isobutane and treating the product with *HI* (Effront, *B* 17, 2325) Long white needles *V* sol alcohol and ether On oxidation with HNO_3 at 200° it gives nitro tolyl isobutyric acid

a* IODO *n* BUTYRIC ACID $\text{C}_4\text{H}_9\text{IO}_2$ *se

$\text{CH}_3\text{CH}_2\text{CHI CO}_2\text{H}$ [110°] (Fittig, *B* 9, 1194) From *a* crotonic acid and fuming *HI* at 100° (Hemilian, *A* 174, 324, *v infia*) Monoclinic crystals (Haushofer, *Z K* 6, 135)

Ethyl ether EtA' (191°) Formed by boiling *a* bromo butyric ether with alcoholic *KI* (Hell, *B* 6, 29)

***β* Iodo-butyric acid $\text{CH}_3\text{CHI CH}_2\text{CO}_2\text{H}$**

Formed, in small quantity, in the preparation of the preceding Liquid

According to Michael and Freer (*J pr* [2] 40, 95) *HI* unites with solid crotonic acid forming, contrary to the statement of Hemilian, only *β* iodo-butyric acid, of low melting point The same acid is formed from liquid crotonic acid and *HI* Boiling water converts it into *β* oxy-butyric acid Hot NaOH aq gives solid crotonic acid

***γ*-Iodo-butyric acid $\text{CH}_3\text{I CH}_2\text{CH}_2\text{CO}_2\text{H}$**

[41°] From the lactone of *γ* oxy butyric acid and *HI* (Henry, *C R* 102, 868, A Saytzeff, *B* 14, 2826) Colourless plates, which become yellow in sunlight *Sl* sol warm water (difference from *β* isomeride), *v* sol. methyl alcohol, ether, and CS_2 .

Methyl ether MeA' (199°) *SG* 1 666

Oil, with pleasant odour

Iodo-isobutyric acid $(\text{CH}_3)_2\text{CI CO}_2\text{H}$ (?)

[36°] From methacrylic acid and fuming *HI* at 0° (Fittig & Paul, *A* 188, 58, 200, 67) Tables (from CS_2) or large prisms (from conc. HIA aq) *Sl* sol. water.

Di iodo isobutyric acid $\text{O}_2\text{H}_2\text{IO}_2$ [127°].

Formed by treating tri oxy isobutyric acid with *HI* and *P* and extracting the product with ether (E Fischer & Tafel, *B* 22, 108) Long needles, *v* *e* sol alcohol and ether

IODO ISOBUTYRIC ALDEHYDE

$(\text{CH}_3)_2\text{CI CH CO H}$ or $(\text{CH}_3)_2\text{CI CO H}$ *SG* 2 229

Preparation—By treating isobutyric aldehyde (22.5 cc) dissolved in 40 cc of alcohol, with iodine (25 g) and iodic acid (10 g) It is very difficult to obtain pure, 130 g of the crude body yielding not more than 1 g sufficiently pure for analysis Reduced silver removes the excess of iodine with the least loss of substance (Chautard, *A Ch* [6] 14, 160)

Properties—Heavy liquid, colourless when pure Its vapour is, like its homologues, exceedingly irritating It is completely decomposed at 100°, and cannot be distilled even *in vacuo* Forms a crystalline compound with NaHSO_4 Is rapidly resinified by solutions of alkalis or mineral acids Combines with aniline, with elimination of water, giving a di amine $\text{Ag}(\text{C}_4\text{H}_9\text{O}_2)$ at 100° yields isobutyl acetate (114°) AgCN and AgSCN yield the corresponding cyano and sulphocyno derivatives

IODO CAMPHOR *v* CAMPHOR

IODO CAPROIC ACID *v* IODO HEXOIC ACID

IODO-CAPRYL-BENZENE *v* IODO OCTYL-

BENZENE

IODO CARBOSTYRIL *v* IODO OXY-QUINOLINE

IODO CHELIDONIC ACID *v* CHELIDONIC ACID

IODO CHRYSIN *v* CHRYSIN

o* IODO CINNAMIC ACID $\text{C}_6\text{H}_4\text{IO}_2$ *se

$\text{C}_6\text{H}_4\text{I CH CO}_2\text{H}$ [214°] (*G a H*), [207°] (*S*) Formed by boiling *o* diazo cinnamic acid with aqueous *HI* (Gabriel & Herzberg, *B* 14, 2037) Formed also by heating *o* iodo benzyl idene malonic acid to its melting point (Stuart, *C J* 53, 142) Crystals (from dilute alcohol)

***m*-Iodo-cinnamic acid $\text{C}_6\text{H}_4\text{I C}_6\text{H}_4\text{CO}_2\text{H}$**

[182°] Formed by boiling *m* diazo cinnamic acid with *HI* (*G a H*) Sol benzene, petroleum ether, and hot alcohol, *sl* sol water

***p* Iodo-cinnamic acid $\text{C}_6\text{H}_4\text{I C}_6\text{H}_4\text{CO}_2\text{H}$**

[*c* 255°] Formed by boiling *p* diazo cinnamic acid with *HI* (Gabriel & Herzberg, *B* 16, 2040) Crystalline mass

IODO CODEINE *v* CODEINE

IODO-COMPOUNDS Organic compounds containing iodine attached to carbon

Formation—1 Unlike chlorine an 'bromine, iodine is not capable of directly displacing hydrogen, inasmuch as the *HI* produced would cause a reverse reaction But if iodic acid, mercuric oxide, or some other substance capable of removing *HI* be present, the substitution may take place (Kekulé, *A* 131, 281) In the case of aniline, the free *HI* combines with excess of aniline and thus iodo aniline may be formed Mercuric oxide will not induce the iodination of fatty compounds, but is especially useful in the case of phenols and oxy acids (Wesselsky, *A* 174, 99). The operation is performed in boiling alcoholic solution, and an intermediate body appears to be mercurioiodate (Lippmann, *B* 7, 1773) Aromatic hydrocarbons may be iodated by heating with iodine and dry FeCl_3 (L Meyer, *A* 231, 195) — 2 By the action of *HI* on hydroxylic compounds,

The alkyl iodides may be prepared in this way by the action of HI or, better, of iodine and phosphorus on the corresponding alcohols, at 100°. The polyhydric alcohols give rise, in this way, to secondary alkyl iodides, the iodine displacing hydrogen attached to a penultimate atom of carbon thus glycerin gives isopropyl iodide. Iodides of acid radicles are formed by the action of iodide of phosphorus on the dry alkaline salts (Canours, *A* 104, 111) — 3 By the union of HI with unsaturated compounds. This combination takes place more readily than in the case of HCl and HBr. The iodine attaches itself to that one of the involved atoms of carbon that is combined with the smaller amount of hydrogen, e.g. $\text{CH}_3\text{CH}=\text{CH}_2 + \text{HI} = \text{CH}_3\text{CHI}-\text{CH}_3$. 4 From aromatic amido compounds by the diazo reaction. This may be performed by heating the diazo compounds with HIAq, or with aqueous KI, e.g. $\text{C}_6\text{H}_5\text{N}_2\text{SO}_3\text{H} + \text{KI} = \text{C}_6\text{H}_5\text{I} + \text{N}_2 + \text{KHSO}_4$. Also by Sandmeyer's reaction which consists in boiling the diazo compound with cuprous iodide, or Diazo compounds — 5 By gradually adding HNO_3 to a hot solution of the amine in HIAq (Losanitsch, *B* 18, 39) — 6 By displacing Cl or Br by I by treatment with a metallic iodide. The elements K, Mg, Ca, Sr, Ba, Al, Mn, and Co prefer chlorine and bromine to iodine and hence the iodides of these metals when heated with chloro or bromo compounds produce iodo compounds (Köhnlcin, *A* 225, 194) Thus propyl chloride is converted into propyl iodide by CaI_2 , SrI_2 , MnI_2 , and CoI_2 , while it is but slightly affected by NiI_2 and FeI_2 , is not affected by SnI_2 , and is wholly decomposed by PI_3 . The conversion of benzyl chloride into benzyl iodide is partially effected by BaI_2 at 15°, by ZnI_2 at 10°, by CdI_2 at 100°, and by PbI_2 at 160°. On the other hand isobutyl chloride is not converted into iodide at 70° by BaI_2 , by ZnI_2 or by CdI_2 , while CdI_2 at 135° decomposes it. The substitution of Cl in chloroacetic acid by I is partly effected by CdI_2 or TI_2 at 100°, and may be effected by BaI_2 or ZnI_2 at 100° (Brix *A* 225, 166). Potassium iodide is a very convenient reagent for the purpose (Perkin & Duppa, *A* 112, 125) but cannot be always depended upon, for though it displaces Cl by I in the case of epichlorhydrin, chloro lactic acid, dichlorhydrin, and mono- and di-chloro acetone, it does not act on di-chlorinated ether, and it decomposes chloral into chloroform and CO_2 . Calcium iodide $\text{CaI}_2 \cdot 3\text{aq}$ is also an excellent reagent for converting chloro and bromo compounds into the corresponding iodo-derivatives (Spindler, *A* 231, 257). Thus it converts $\text{CH}_3\text{CH}_2\text{CH}_2\text{Br}$ into ethylene iodide, and $\text{CH}_3\text{CH}_2\text{CH}_2\text{Br}$ into CH_3CHI , but it fails with $\text{CH}_3\text{CH}_2\text{CH}_2\text{Cl}$, with $\text{CHCl}_2\text{CH}_2\text{Cl}$, with $\text{CCl}_2\text{CH}_2\text{Cl}$, with CCl_3 , with chloro benzene, and with chloral. Dry CaI_2 requires a higher temperature (120°) before it reacts, and this decomposes many of the iodo compounds that should be formed — 7 Iodine chloride also acts as an iodating substance, especially upon aromatic amines. Thus it converts aniline (dissolved in HOAc) into di-iodo aniline (when 2 mols ICl are used) and tri-iodo-aniline (when 3 mols ICl are used). It converts acetanilide into acetyl p-iodo aniline, m nitro aniline into di-iodo m nitro-aniline, p-nitro aniline chiefly into iodo-p-nitro-aniline, and p-toluidine into di-iodo-p-toluidine: but it

does not act on acetyl p toluidine (*B* 11, 107) — 8 By heating chloro- compounds with fuming HIAq (Lieben, *Z* 1868, 712) — 9 Nitrogen iodide, acting on dilute alcoholic solutions of the potassium derivatives of phenols, forms iodo phenols and ammonia (Willgerodt, *J pr* [2] 37, 446).

Reactions — 1 The displacement of iodine by chlorine or bromine may be effected by heating with the chloride or bromide of Hg, Cu, Ag, Sn, Pb, As, or Sb (v **Chloro compounds and Bromo compounds**) — 2 Chlorine and bromine displace iodine directly — 3 Sodium frequently removes iodine with ease, e.g. $2\text{C}_6\text{H}_5\text{I} + \text{Na}_2 = \text{C}_6\text{H}_5 + 2\text{NaI}$. In like manner (by what is known as Fittig's reaction) $\text{C}_6\text{H}_5\text{I} + \text{C}_6\text{H}_5\text{I} + \text{Na}_2 = \text{C}_6\text{H}_5\text{C}_6\text{H}_5 + 2\text{NaI}$. 4 Water readily decomposes tertiary alkyl iodides, forming HI and a tertiary alcohol. Methyl and ethyl alcohol at 100° act in like manner, forming a tertiary alcohol and MeI or EtI — 5 Silver obtained by reduction sometimes removes iodine e.g. $2\text{CH}_3\text{I} + \text{CH}_3\text{CO}_2\text{H} + \text{Ag}_2 = 2\text{AgI} + \text{CO}_2\text{H} + \text{CH}_3\text{CH}_2\text{CH}_2\text{CO}_2\text{H}$.

Some aromatic iodo compounds can take up chlorine, e.g. PhI gives PhICl_2 (Willgerodt, *J pr* [2] 33, 154), $\text{C}_6\text{H}_5\text{BrI}$ gives $\text{C}_6\text{H}_5\text{BrICl}_2$, while $\text{C}_6\text{H}_5\text{BrI}$ gives $\text{C}_6\text{H}_5\text{BrICl}_2$, and $\text{C}_6\text{H}_5(\text{NO}_2)\text{I}$ gives $\text{C}_6\text{H}_5(\text{NO})\text{ICl}_2$. $\text{C}_6\text{H}_5(\text{NO})\text{I}$ has no action.

iodo m CRESOL $\text{C}_6\text{H}_4\text{MeI}(\text{OH})$ [1 x 3] Oil. Formed by adding iodine to a solution of m-cresol in ammonia and alcohol (Willgerodt & Kornblum, *J pr* [2] 39, 289).

Iodo p cresol $\text{C}_6\text{H}_4\text{MeI}(\text{OH})$ [1 3 4] Formed by the action of dry iodine on sodium m-cresol suspended in CS (Schall & Dralle, *B* 17, 2533). Liquid. Volatile with steam.

Methyl ether $\text{C}_6\text{H}_4\text{MeI}(\text{OMe})$ (238°), oil. **Di iodo o cresol** $\text{C}_6\text{H}_3\text{MeI}_2(\text{OH})$ [1 3 4 2] [70°] Formed by the action of iodide of nitrogen upon the sodium compound of o-cresol (Willgerodt *J pr* [2] 37, 448, 39, 289).

Acetyl derivative $\text{C}_6\text{H}_3\text{MeI}_2(\text{OAc})$ [56°] Tables.

Picrate. [204°] Yellow crystalline aggregates.

Di iodo m cresol $\text{C}_6\text{H}_3\text{MeI}_2(\text{OH})$ [76°] Formed together with iodo m cresol (W & K). Long needles, sol. usual menstrua.

Di iodo p cresol $\text{C}_6\text{H}_3\text{MeI}_2(\text{OH})$ [1 3 5 4] [61°] Formed together with the preceding (Schall & Dralle, *B* 17, 2534). White tables. Easily soluble in alcohol, sparingly in water and ligroin. Volatile with steam.

Acetyl derivative $\text{C}_6\text{H}_3\text{MeI}_2(\text{OAc})$ [62°], white tables.

Benzoyl derivative $\text{C}_6\text{H}_3\text{MeI}_2(\text{OBz})$ [130°], glimmering white prisms.

Ethyl ether $\text{C}_6\text{H}_3\text{MeI}_2(\text{OEt})$ [77°] White solid sol. organic solvents (Willgerodt & Kornblum, *J pr* [2] 39, 289).

iodo CRESOL SULPHONIC ACID $\text{C}_6\text{H}_4\text{MeI}(\text{OH})\text{SO}_3\text{H}$ [3 1 2 5] [155°] Formed together with an isomeride, by the action of iodine on the potassium salt of o-cresol sulphonic acid (Kehrmann, *J pr* [2] 37, 338). Colourless plates (containing 3aq). Converted by nitric acid into dinitro cresol [86°] Chromic acid forms iodo toluquinone [115°].

Salts — A'K fine needles — A', Ba 4aq. **Di-iodo m cresol sulphonic acid** $\text{C}_6\text{H}_3\text{MeI}_2(\text{OH})(\text{SO}_3\text{H})$ [3 6 2 1 4] [70°] From potassium m-cresol sulphionate by treatment with

Iodine (Kehrmann, *J pr* [2] 39, 392) Small needles Oxidised by CrO_3 to di-iodo-toluquinone

IODO- ψ -CUMENE $\text{C}_9\text{H}_7\text{IO}_2$ [1245] *Iodo-tri-methyl-benzene* [37⁵] (258^o) Formed by decomposing ψ -cumene diazopiperidine with HIAq (SG 173) (Wallach & Heusler, A 243, 233) Formed also from ψ -cumidine by diazotising and heating the product, dissolved in H_2SO_4 with KI (Kurzel, B 22, 1586) Scales Treatment with H_2SO_4 forms two isomeric di-iodo ψ -cumene sulphonic acids ψ -cumene sulphonic acid and iodo ψ -cumene sulphonic acid One of the di-iodo ψ -cumenes melts at 74^o, the other is an oil

IODO ψ -CUMENE SULPHONIC ACID

$\text{C}_9\text{H}_6\text{IO}_2\text{SO}_3\text{H}$ Formed as above Scales, m sol water

Salts — NaA'aq plates — BaA' aq needles

IODO CYANETHINE v **CYANTHINE**

IODO ETHANE v **ETHYL IODIDE**

Di-iodo-ethane v **ETHYLENE IODIDE** and **ETHYLDENE IODIDE**

Tri-iodo-ethane $\text{C}_2\text{H}_3\text{I}_3$, ρ CH_3Cl , [95^o]

Formed by the action of aluminium iodide upon CH_3CCl_3 (De Boissieu, B [2] 49, 16) Yellow octahedra V e sol CS, benzene, and ether, v sol petroleum ether, m sol alcohol Smells faintly of iodoform

IODO-ETHYL ALCOHOL v **GLYCOLIODHYDRIN**
 β -IODO ETHYL AMINE $\text{CH}_3\text{CH}_2\text{NH}_2$

Hydriodide B'HI [194⁵] From vinylamine and conc HIAq (Gabriel, B 21, 1049) Trim⁴ crystals (from hot alcohol) V sol water When treated with KOH it gives a liquid, soluble in water, with a disagreeable odour

Picrate $\text{B}'\text{C}_6\text{H}_3(\text{NO}_2)_3\text{OH} \frac{1}{2}\text{aq}$ [c 105^o] (hydrated), [131⁵] (anhydrous) Short compact yellow prisms

Di-iodo ethyl-amine EtNI_2 , v *Ethyl di-iodo amine*, vol 11 p 474

Di-iodo ETHYL BENZENE $\text{C}_8\text{H}_7\text{I}_2$, ρ $\text{C}_6\text{H}_5\text{CH}_2\text{CH}_2\text{I}$ *Styrene di iodide* From styrene and a concentrated solution of iodine in KIAq (Berthelot, B 6, 295, 7, 277) Crystals, v sol ether Rapidly decomposes into iodine and metastyrene

IODO-ETHYLENE $\text{C}_2\text{H}_2\text{I}_2$, ρ CH_2CHI *Vinyl iodide* (56^o) SG 2 208 Formed by distilling ethylene iodide with conc alcoholic potash, the receiver being kept very cool, and the distillate finally mixed with water (Regnault, A 15, 69, E Kopp, C R 18, 871) Also from ethyldene iodide (Gustavson, B 7, 731) Formed also by the union of HI with acetylene (Berthelot, A 132, 122, Semenov, Z 186⁵, 725) Oil, v sol alcohol and ether, with alliacious odour

Di-iodo-ethylene $\text{C}_2\text{H}_2\text{I}_2$, ρ CHI CHI *Acetylene di iodide* [73⁵] (192^o cor) SG 2 3302 Formed by passing acetylene over iodine moistened with alcohol (Sabanejeff, A 178, 109, 216, 275, Plimpton, C J 41, 392) Elastic needles (from alcohol) It appears to be accompanied by a small quantity of a liquid isomeride SG 2 2942 With alcoholic AgNO_3 , it forms long needles of a compound $\text{C}_2\text{H}_2\text{I}_4\text{AgNO}_3$, which is decomposed by HCl, giving off acetylene

Tetra-iodo-ethylene Cl_2Cl_2 *Di carb t-tetra-iodide* [c 165⁵] Prisms Formed by the action of a solution of iodine in aqueous KI upon the cuprous compound of potassium propargylate. Formed also by the action of the

iodine solution upon acetylene copper (Hemcolls & Stolz, B 18, 2283)

IODO-ETHYLDENE DI-ETHYL DI-SULPHONE $\text{CH}_3\text{CH}(\text{SO}_2\text{Et})_2$ [109⁵] From ethyldene diethyl di sulphone by boiling with iodine (Fromm, A 253, 146) Needles

IODO-DI-ETHYL OXIDE $\text{C}_4\text{H}_{10}\text{IO}$, ρ $\text{CH}_3\text{CH}_2\text{CH}_2\text{OEt}$ (155^o) SG 2 1692 VD 691 (obs and calc) Formed by heating ethylene iodide with alcohol at 75^o (Baumstark, B 7, 1172) Formed also from the mono ethyl derivative of glycol by treatment with iodide of phosphorus (Demole, B 9, 746) Pungent oil, not decomposed by water Alcoholic potash attacks it energetically, yielding $\text{CH}_3\text{CH}_2\text{OEt}$ (35^o), and $\text{CH}_3(\text{OEt})\text{CH}_2(\text{OEt})$ (17^o) Boiling with reduced silver does not affect it

IODO-(β)-ETHYL-THIOPHENE

$\text{C}_6\text{H}_4(\text{C}_2\text{H}_5)\text{IS}$ Oil Formed by the action of iodine upon (β) ethyl thiophene in presence of HgO (Bonz, B 18, 551)

IODO-FLUORO-BENZENE v **FLUORO IODO BENZENE**

IODOFORM CHI_3 , *Tri-iodo methane* Mol w 394 [119⁵] SG 2

Depression of freezing point of benzene by iodoform Raoult, A Ch [6] 2, 88, Paterno, B 22, 465

Formation —1 By heating alcohol with iodine and caustic or carbonated alkali (Strullus A Ch [2] 20, 314) Iodoform is formed in like manner by the action of iodine and potash or aldehyde, acetone, lactic acid, and many other substances (cf vol 1 p 96) —2 From chloroform and HI (Baeyer, B 5, 1091) —3 By heating chloroform with CaI_2 at 75^o (Spindler, A 231, 263) —4 Together with carbamic ether, by the action of iodine on sodium carbamic ether (Mulder, B T C 7, 333)

Preparation —1 Alcohol containing about 20 p.c. aldehyde is mixed with ten times its weight of aqueous NaOH , iodine is added, and the mixture stirred from time to time The iodoform quickly separates in the cold (Günther, Ar Ph [3] 25, 373) —2 To prepare iodoform R. Rother (Ph [3] 4, 593) heats 32 pts iodine, 32 potassium carbonate, 16 alcohol of 9 per cent and 80 water, till the mixture becomes colourless, then decants the clear liquid, collects the separated crystals of iodoform, and mixes the filtered liquid with 16 to 24 pts hydrochloric acid and 2 to 3 pts potassium dichromate The liquid is then neutralised with potassium carbonate, and 32 pts more of that salt are added, together with 6 pts iodine and 16 alcohol The reaction having been completed at the heat of the water bath, the product is treated as above, whereby a fresh quantity of iodoform is obtained —3 By the action of excess of sodium hypochlorite on an aqueous solution of acetone (6 pts), potassium iodide (50 pts), and NaOH (2 pts) (Suillot & Raynaud, B [3] 1, 3)

Properties —Hexagonal yellow tables, $\alpha = 1.9015$ (Rammelsberg & Kokscharoff, C C 1857, 524, cf Dogiel, N Peter's Acad Bull 20, 337) Above its melting-point it partly decomposes, giving off iodine and HI Has a peculiar persistent odour Volatile with steam Insol water, acids, and alkalis, v sol⁵ alcohol, ether, and fixed and volatile oils An alcoholic solution of iodoform is decomposed under the influence of

light. The decomposition is accelerated by dry oxalic acid, iodine and methylene iodide are produced (Mulder, *R T C* 7, 316, cf. Humbert, *J Ph* [8] 29, 352). Does not in the least hinder the putrefaction of pancreas at 35° (Boillat, *r pr* 183, 308).

Detection—An alcoholic solution of iodoform warmed with aqueous KOH and a little phenol forms a red substance, which dissolves in dilute alcohol, giving it a crimson colour (Lustgarten, *M* 3, 717).

Reactions—1 Heated in a sealed tube at 150°, either alone or with iodine, it yields methylene iodide and brown products (Hofmann, *C J* 13, 65).—2 Bromine gives bromoform (Löschner, *B* 21, 410), not an excess of Br at high temperatures forms CB_3 .—3 Distilled with $HgCl_2$ it yields C^2Cl_2I (Schlagdenhaufen, *J Ph* [8] 29, 247).—4 Mercuric oxide forms CO, formic acid, HgI_2 , and water.—5 Boiling aqueous KOH forms a little potassium formate.—6 Boiling alcoholic potash forms methylene iodide (Bruning, *A* 104, 187, Butlerow, *C R* 47, 595).—7 NaOEt forms methylene iodide, NaI, aldehyde, acrylic acid, and $CH_3CH(OEt)CO_2H$ (Butlerow, *A* 107, 110, 114, 204, 118, 325).—8 K_2SO_4 forms $CH_3(SO_4K)$, (Streeker, *A* 148, 90).—9 By the copper zinc couple iodoform may be reduced to methane (Gladstone & Tribe, *C J* 28, 508).—10 Unites with PEt_3 forming $C_3H_7P_3I_4$ (Hofmann, *Pr* 10, 189).—11 PCl_5 forms chloroform. 12 When made into a paste with moist finely-divided silver, acetylene is at once given off. A mixture of ppt silver and copper is even more effective (Cazeneuve, *C R* 97, 1371).—13 Silver nitrate acts readily at ordinary temperatures, forming AgI, nitric acid, and CO, so that by nitrating the liberated acid an estimation of the quantity of iodoform present may be formed (Greshoff, *R T C* 7, 342).—14 Iodoform exposed to direct sunlight is completely oxidised by air, forming CO_2 , iodine, and water (Dacomo, *G* 16, 247).—15 Mercuric acetate is reduced by iodoform to mercurous acetate, CO_2 being evolved. It does not reduce mercuric nitrate (although chloral forms, in this case, mercurous chloride) (Cotton, *J Ph* [5] 16, 481).—16 Isobutyl alcohol (200 g) mixed with sodium (20 g) and iodoform (100 g) gives a violent reaction, the products being formic acid, isobutyric acid, pentenoic acid, OMe , CH_3CO_2H [70°], isobutyl oxy isobutyric acid $C_4H_9O_2$, CO_2H , a lactone of an oxy octoic acid, isobutylene, methyl isobutyl oxide, methylene di isobutyloxyde, a compound $C_7H_{14}O_2$, and other bodies (Gorboff & Kessler, *J R* 1887, 428).

iodo-fumaric acid $C_2HI(CO_2H)_2$. Iodo-maleic acid [184°]. Formed by dissolving acetylene di carboxylic acid in strong aqueous HI (Baudrowski, *B* 15, 2697). Fine glistening crystals. Very soluble in water, alcohol, and ether.

Salts—A'HK small sparingly soluble crystals—A'Ag, crystalline pp—A'Pb 2aq crystalline pp.

iodo-hen-decoic acid $C_{11}H_{21}IO_2$. $CH_3(CH_2)_9CO_2H$ [24°]. From hen-decenoic acid and HI (Ph Brunner, *B* 19, 2224). Crystalline.

iodo heptioic aldehyde $C_6H_{11}IO$. S.G. 12 281. Formed by treating heptioic

(α -nanthic) aldehyde (75 g) diluted with alcohol (100 cc) with iodine (50 g) and iodic acid (20 g). The crude product is purified by treatment with reduced silver. The yield is very bad (Chautard, *A Ch* [6] 16, 170). Very unstable liquid with irritating odour. Decomposed by heat. Easily decomposed by KOH, NaOH, ammonia, and mineral acids.

Reactions—1 Nitric acid gives heptioic aldehyde and heptioic acid.—2 Silver acetate forms octyl acetate (208°).—3 By heating iodo heptioic aldehyde (120 g) dissolved in alcohol with silver cyanide (67 g) there is formed $C_6H_{13}CyCHO$ (177°) S.G. 12 913. This is a colourless liquid which reduces Fehling's solution and ammoniacal $AgNO_3$, but does not form crystalline compounds with aniline or phenyl hydrazine.—4 Silver sulphocyanide forms the corresponding sulphocyanic heptioic aldehyde.—5 Heating with aniline forms $C_6H_5I(NHC_6H_5)$.

iodo heptylene $C_6H_{11}I$. $Pr CH_2Cl CHMe$ or $Pr CH Cl CH_2Me$ (140°–150°). From heptene (*q v*) and HI (Morris, *C J* 41, 179).

iodo hexane $C_6H_{13}I$. Di iodo hexane $C_6H_{11}I_2$. $CH_3CHICH_2CH_2CHICH_3$. **Hexylene iodide** S.G. 12 024. Formed by heating di allyl in a sealed tube for 5 hours with excess of conc HIAq at 100° (Wurtz, *A Ch* [4] 3, 129), or better, by passing gaseous HI into strongly cooled diallyl (Sorokin, *J pr* [2] 23, 18). Oil, gives off iodine when heated. With sodium it gives hexylene and hexenyl iodide. Alcoholic potash forms diallyl. Silver acetate forms hexylene acetate $C_6H_{11}(OAc)$, whence baryta forms di oxy hexane.

Tetra iodo hexane $C_6H_8I_4$ [above 100°]. From diallyl and iodine (Berthelot & De Luca, *A* 100, 363).

TETRA IODO-HEXINENE *v* **Tetra iodide** of DIPROPARGYL.

iodo-hexoic acid $C_6H_{11}IO_2$. Formed by mixing hydrosorbic acid with conc HIAq at 0° (Fittig, *A* 200, 44). Oil, turning yellow on exposure to light. May be reduced by sodium amalgam to *n*-hexoic acid.

iodo hexylene $C_6H_{11}I$. $CH_2CHCH_2CHICH_2CH_3$. **Hexenyl iodide**. **Diallyl hydronolide** (165°) S.G. 12 1497. Formed, together with di iodo hexane, by the action of HI on diallyl (Wurtz, *A Ch* [4] 3, 155). Formed also by the action of alcoholic potash or of an alloy of tin and sodium on di iodo hexane $CH_2CHICH_2CH_2CHICH_2CH_3$. When treated for 24 hours with moist Ag_2O it forms diallyl, hexylene, and two liquids $C_6H_{12}O$ (d 135°) and $C_{12}H_{24}O$ (180°).

Iodo hexylene $C_6H_{11}I$ (131°) S.G. 12 192. Formed from the hexenyl alcohol obtained by distilling glycerin with lime (Destrem, *A Ch* [5] 27, 58).

Iodo hexylene $C_6H_{11}I$ (142°–145°). From pinacone hydrate and HI at 160° (Bouchardat, *Z* 1871, 699). Reduced by tin and HCl to hexylene. Bromine gives crystalline $C_6H_8Br_2$.

iodo hippuric acid $C_9H_{11}NO_3$. $NH_2CH(CO_2H)(?)$. Formed by treating a boiling alcoholic solution of hippuric acid with iodine (Maier, *Z* 1865, 415). White needles. Decomposes at 90°. All its salts are soluble in water excepting the silver salt.

Iodo-hippuric acid $C_6H_4I.CO.NH.OH.CO_2H$
Formed by the action of aqueous HI on the sulphate of diazo hippuric acid (Griess, *Z* [2] 4, 725, *B* 1, 190) Laminæ, v sol cold alcohol and ether, m sol hot water, nearly insol cold water The iodine is not easily removed

IODO-HYDROCINNAMIC ACID v Iodo-
PHENYL PROPIONIC ACID

DI-IODO-HYDROQUINONE $C_6H_2I_2(OH)_2$ [5841] [145°] Formed by boiling di-iodo-quinone [180°] with a dilute solution of $SnCl_2$ (Seifert, *J pr* [2] 28, 438, Kehrman, *J pr* [2] 37, 337) Long colourless needles (from hot water) Volatile with steam

Di-iodo hydroquinone $C_6H_2I_2(OH)_2$ [142°] From di-iodo quinone [159°] and aqueous SO_2 (Metzeler, *B* 21, 2555) White needles, v sol hot water, alcohol, ether, and $HOAc$, m sol benzene Reoxidised by $FeCl_3$ to di-iodo quinone

Di-acetyl derivative $C_6H_2I_2(OAc)_2$ [148°] Needles or plates (from alcohol), insol water, v sol alcohol and ether, sl sol cold $HOAc$

IODO HYDROTHYMOQUINONE
 $C_8H_8MePr(OH)_2$ [52641] [74°] Obtained by reducing iodo thymoquinone with $SnCl_2$ (Kehrman, *J pr* [2] 39, 392) Colourless prisms, v sol organic solvents

IODO HYDRO TOLUQUINONE
 $C_8H_8Me(OH)_2$ [6241] [111°] From the corresponding quinone, by reduction with $SnCl_2$ (Kehrman, *J pr* [2] 39, 392) Soft silky aggregates of needles, sol organic solvents

IODOL v **TETRA IODO PYROLE**

IODO-LACTIC ACID v **IODO OXY PROPIONIC ACID**

IODO-MALEIC ACID v **IODO FUMARIC ACID**

IODO MELLILIC ACID *Methyl derivative*
 $C_8H_8IO_2$, v $C_8H_8(OMe)_2$ $C_6H_4I.CO_2H$ From the methyl derivatives of (a) and of (β) coumaric acid and fuming aqueous HI in the cold (Perkin, *C J* 39, 429) Aqueous Na_2CO_3 forms $C_8H_8(OMe)CH_2CH_2$ and CO_2

IODO METHANE v **METHYL IODIDE**

Di-iodo-methane v **METHYLENE IODIDE**

Tri-iodo methane v **IODOFORM**

Tetra-iodo-methane CI_4 *Carbon tetra iodide*
SG 22 432 Formed by mixing CCl_4 with aluminium iodide dissolved in CS_2 (Gustavson, *A* 172, 173, *C R* 78, 882) Formed also by heating CCl_4 with CaI_2 at 75° (Spindler, *A* 231, 264) Dark red regular octahedra (Friedel, *Bl* [3] 21, 482) Sol CS_2 , alcohol, ether, and MeI May be kept for some days, but gradually decomposes The decomposition is hastened by both CS_2 and CO_2 Above 80° it quickly splits up into carbon and iodine Converted into iodo form by boiling with aqueous HI, and even by boiling with water Scarcely attacked by aqueous potash or H_2SO_4 Chlorine converts it into CCl_4 Br gives CBr_4

IODO-DI METHYL AMINE v *Di methyl-iodo-amine* under **DI METHYL AMINE**

IODO-DI-METHYL-ANILINE C_6H_4IN v $C_6H_4INMe_2$ [79°] Formed by adding iodine to a solution of di-methyl aniline in CS_2 (Weber, *B* 10, 765) Formed also by the action of cyanogen iodide on di-methyl aniline in the cold, heat being evolved and HCN given off (Merz & Weith, *B* 10, 757). White laminæ, sol alcohol

and $HClAq$ Not attacked by caustic soda When heated above 80° it gives off HI and leaves a residue which forms a deep violet solution in alcohol — $B' F.PtCl_2$

IODO-TRI METHYL-BENZENE v. Iodo ψ

CUMENE

IODO METHYL-BUTYL-BENZENE v Iodo-BUTYL-TOLUENE

DI IODO METHYLENE **DI ETHYL DI-SULPHONE** $Cl_2(SO_2Et)_2$ [177°] Formed from methylene di ethyl di sulphone I, and $KIAq$, by exposure to sunlight (Fromm, *A* 253, 161) Needles, sl sol boiling water

DI-iodo-METHYLENE OXIDE $C_2H_2I_2O$ v $(CH_2I)_2O$ (219°) From dry 'oxymethylene' (formic paraldehyde) and gaseous HI (Tishchenko, *J R* 1887, 464) Oil

IODO TRI METHYL PIPERIDINE $C_8H_{11}IN$ [60°] Formed by heating ethyldiene diacetamide with HI (E Fischer, *B* 17, 1796) Colourless rectangular prisms Nearly insol water — $B'HI$ very sparingly soluble

Iodo-tetra-methyl-piperidine $C_8H_{11}NI$ [90°] Formed by heating triacetone with HI at 150° (E Fischer, *B* 17, 1791) Easily soluble in alcohol and ether, insol water Colourless tables (from ether)

IODO DI METHYL THIOPHENE C_6H_4IS v

CI CMe }
 CH CMe } S (97°) From di-methyl thiophene (from coal tar), iodine, and HgO (Messinger, *B* 18, 1638) Oil

Iodo - tri - methyl - thiophene C_6H_3IS v

CMe CMe }
 CI CMe } S Formed by the action of iodine (48 g) and HgO (21 g) on tri-methyl thiophene (12 g) dissolved in ligroin (24 g) (Zelinsky, *B* 21, 1837) Non volatile oil

(a) **IODO NAPHTHALENE** $C_{10}H_7I$ (305°)

Formation — 1 By adding iodine to a solution of mercury dinaphthyl $Hg(C_{10}H_7)_2 + 2I_2 = HgI_2 + 2C_{10}H_7I$ (Otto, *A* 147, 173) — 2 From diazo naphthalene sulphate and Li (Nolting, *B* 19, 135)

Properties Thick yellowish oil, not solidified at — 17° Miscible with alcohol, ether, CS_2 , and benzene Alcoholic potash at 160° forms some naphthalene Boiling $HIAq$ (127°) completely converts it into naphthalene When a solution in CS_2 is boiled with aluminium chloride there is formed naphthalene, iodine, and tarry matter, but no (β) iodo naphthalene (Roux, *Bl* [2] 45, 517)

Picric acid compound [127°] Golden yellow needles

(β) **Iodo-naphthalene** $C_{10}H_7I$ [55°] Prepared by the action of HI on (β) diazo naphthalene sulphate (P Jacobson, *B* 14, 804) Colourless plates, sol ether, alcohol, and $HOAc$ Volatile with steam

(α, α) **Di-iodo-naphthalene** $C_{10}H_6I_2$ [14] [110°] From (α) iodo (α) nitro naphthalene by reduction, and treatment of the resulting iodo naphthylamine with H_2SO_4 and $NaNO_2$, followed by HI (Meldola, *C J* 47, 522) Needles (from alcohol)

(α, β) **Di-iodo-naphthalene** $C_{10}H_6I_2$ [12] [81°] From (β)-iodo-(α)-nitro-naphthalene in

the same way as the preceding (Meldola) Scales (from alcohol)

(*β*) - IODO - NAPHTHALENE SULPHONIC ACIDS $C_{10}H_7I(SO_3H)$ Two isomerides are formed by sulphonation (*β*) iodo-naphthalene (Armstrong & Wynne, *C J Proc* 3, 22) One of the isomerides forms the salts KA' and BaA' , 4 $\frac{1}{2}$ aq, both crystallising in hexagonal scales, the other forms the salts KA' & Ca' crystallising in laminæ, and BaA' , 2aq in flat needles

(1, 5) Iodo naphthalene sulphonic acid $C_{10}H_7I(SO_3H)$ [129°] Formed from (a) diazo naphthalene sulphonic acid and warm $HIAq$ (40 °c) (Mauzelus, *B* 22, 2820) Colourless tables (containing 2aq) v sol. water

Salts— $A'K$ aq plates m sol water— $A'Na$ aq indistinct needles— $A'Ag$ plates S 66— $A'Ca$ 2aq scales, sl sol water— $A'Ba$ aq scales S 23 (in the cold) S 1 at 00°— $A'Mg$ needles sl sol water— $A'Zn$ 6aq scales, m sol water— $A'Cu$ 4aq greenish white needles, sl sol water— $A'Pb$ 4aq sl sol. water— $A'Mn$ 4aq indistinct crystals sl sol water

Methyl ether MeA' [60°] Prisms v sol Et_2O , sl sol cold $EtOH$

Ethyl ether EtA' [74°] Six sided tables v sol Et_2O

n Propyl ether PrA' [67°] Colourless tables

Isopropyl ether PrA' [90°] Prisms Chloride $C_{10}H_7ISO_2Cl$ [114°] Monoclinic prisms v sol hot $AcOH$, sl sol ligroin

Bromide [153°] Prisms

Amide $C_{10}H_7I(SO_2NH_2)$ [239°] Scales

(a) IODO (*β*) NAPHTHOL $C_{10}H_7I(OH)$ [12] [94.5°] (M), [c 100°] (W) Prepared by dissolving equal parts (20 g) of (*β*) naphthol, lead acetate, and dried sodium acetate in $HOAc$, cooling, and gradually adding iodine (35.2 g) dissolved in hot $HOAc$ The product is pptd by water and recrystallised from dilute alcohol (Meldola, *C J* 47, 527) Formed also by the action of iodide of nitrogen on a dilute alcoholic solution of sodium (*β*) naphthol (Willgerodt, *J pr* [2] 37, '48)

Properties—Long colourless needles, sl sol boiling water, v sol organic solvents Volatile with steam Sol cold dilute $KOHAq$, decomposed by hot alkalis Nitric acid liberates iodine $KMnO_4$ gives phthalic acid

(a) IODO (a) NAPHTHYLAMINE

$C_{10}H_7I(NH_2)$ [41] From (a) iodo (a) nitro-naphthalene, zinc dust, and $HOAc$ (Meldola, *C J* 47, 521) Its solution rapidly oxidises in the air forming a red colouring matter— $B'H_2SO_4$, tolerably stable white needles

Acetyl derivative $C_{10}H_7I(NHAc)$ [196°] From the preceding by boiling with Ac_2O (M) Needles (from alcohol) Gives a nitro derivative [286°]

IODO-*o* NITRO ANILINE $C_6H_4I(NO_2)(NH_2)$ [52.1] Formed by heating *o* iodo nitro-benzene [168°] with alcoholic NH_3 at 170° for several days (Körner, *G* 4, 386) Large steel blue laminæ (from alcohol in sealed tubes) yellow by transmitted light Does not melt at 220° Not attacked by nitrous ether

Iodo *o*-nitro-aniline $C_6H_4I(NO_2)(NH_2)$ [42.1] [122°] Formed by warming a solution of the acetyl derivative of *p* iodo-aniline in $HOAc$ with

HNO_3 (Michael & Norton, *B* 11, 109) Long orange-yellow needles (from alcohol)

Iodo-*p*-nitro-aniline $C_6H_4I(NO_2)(NH_2)$ [24.1] [105.5°] Prepared by the action of iodine chloride (2 pts) on *p* nitro aniline (1 pt) dissolved in dilute $HClAq$ (Michael & Norton, *m* 1, 2f2, *B* 11, 113) Long yellow needles— $B'HCl$ [163°]

Di iodo-*m*-nitro-aniline $C_6H_3I_2(NO_2)_2(NH_2)$ [42.3.1] [145.5°] Formed by passing iodine chloride (3 pts) into a solution of *m* nitro-aniline (1 pt) in $HClAq$ (Michael & Norton, *Am* 1, 255, *B* 11, 112) Long slender yellow needles, v sol cold alcohol, sl sol acetic ether, insol water Its hydrochloride is not decomposed by water

Di-iodo-*p*-nitro-aniline $C_6H_3I_2(NO_2)_2(NH_2)$ [62.4.1] [244°] Prepared by the action of iodine chloride (2 mols) on *p* nitro aniline dissolved in chloroform (M & N) Yellow prisms with blue fluorescence (from alcohol) Sl sol hot alcohol, sol $CHCl_3$ and $EtOAc$, v sol hot nitro benzene It is not basic

o IODO NITRO BENZENE $C_6H_4I(NO_2)$ [12] [49°] Formed, together with the *p* isomeride, by treating iodo benzene with HNO_3 (Körner, *G* 4, 305) The *p* isomeride crystallises first from alcohol Long flat lemon yellow needles Gives a little *o* iodo aniline and much aniline on reduction

m Iodo-nitro benzene $C_6H_4I(NO_2)$ [1.3] [36°] (c 280°) Formed from *m* nitro aniline by displacing NH_3 by *I* through the diazo reaction (Griess, *Z* 1866, 218) Monoclinic plates, *a b c* = 2.296 1.1129 (Panebianco *G* 9, 360) The same body is perhaps formed by the action of ICl on sodium *m* nitro benzoate (Schutzenberger & Segenwald, *Rép Chim pure*, 1862, 144)

p Iodo-nitro-benzene $C_6H_4I(NO_2)$ [1.4] [171°] The chief product of the action of fuming HNO_3 upon iodo benzene (Kukulé, *A* 137, 129) Obtained also from *p* nitro aniline by the diazo reaction Pale yellow needles

Di chloride $C_6H_3I_2(NO)_2Cl_2$ Formed by passing Cl_2 into a solution of iodo nitro benzene in chloroform (C Willgerodt, *J pr* [2] 33, 160) Hardly sol CS_2 , Et_2O , or light petroleum Sol. $CHCl_3$ and benzene At 150° it gives off Cl_2 , leaving $C_6H_3I_2(NO_2)_2$ With alcohol it reacts forming aldehyde, HCl and $C_6H_3I_2(NO_2)_2$

Iodo-*di*-nitro-benzene $C_6H_3I_2(NO_2)_2$ [12.4] [89°] Obtained by nitrating *o* or *p*-iodo nitro-benzene with H_2SO_4 and HNO_3 (Körner, *J* 1875, 322) Yellow triclinic crystals, *a b c* = 1.634 1.940 (La Valle, *G* 10, 3) V sl sol cold alcohol. Hot dilute KOH forms *p* nitro phenol Alcoholic NH_3 gives *di* nitro aniline, though the decomposition is incomplete in the cold

Iodo *di*-nitro-benzene $C_6H_3I_2(NO_2)_2$ [12.6] [114°] Formed in small quantity, together with the preceding, by treating *o* iodo nitro benzene with HNO_3 mixed with H_2SO_4 (Körner, *G* 4, 305) Orange triclinic tables (from alcohol) More sol alcohol and ether than the preceding Converted by heating with alcoholic NH_3 into *di* nitro-aniline [188] May be reduced to phenylene diamine.

Iodo-*tri*-nitro-benzene $C_6H_2I_3(NO_2)_3$ [12.4.6] [164°] From chloro *tri* nitro-benzene and alcoholic KI (Hepp, *A* 215, 361) Golden needles

Converted by boiling aqueous KOH into picric acid

Di-iodo nitro-benzene $C_6H_4I_2(NO_2)$ [134] [168°] Obtained from *m* di-iodo benzene by dissolving in hot fuming HNO_3 (Körner). Tri-methyl octahedra, $a, b, c = 647, 1458$ (La Valle, *G* 10, 2) Sl sol ether, v sl sol cold alcohol By heating for a few days with alcoholic NH_3 at 170° it forms iodo nitro aniline

Di-iodo-nitro-benzene $C_6H_3I_2(NO_2)$ [124?] [1125°] Formed by dissolving *o*-di-iodo benzene in fuming HNO_3 and, after a few minutes, diluting with water (Körner a Wender, *G* 17, 491) Lemon yellow needles or small prisms (from alcohol)

(α)-IODO-NITRO-BENZOIC ACID $C_6H_3INO_3$, *to* $C_6H_4I(NO_2)(CO_2H)$ [32or61] [235°] Formed, together with the two following, by heating *m*-iodo-benzoic acid with conc HNO_3 (Grothe, *J pr* [2] 18, 324, Cunze a Hubner, *A* 135, 106) Sl sol water, especially if HCl be present — Gives *o*-amido benzoic acid on reduction $NH_4A'aq$ — $NaA'3aq$ — $CaA'2aq$ — $SrA'4aq$ — $BaA'3aq$

Ethyl ether EtA' [84°]

(β)-Iodo nitro-benzoic acid $C_6H_3I(NO_2)(CO_2H)$ [36or21] [174°] Formed as above V sol water, especially if HCl be present Melts under water May be reduced to *o*-amido benzoic acid — $NH_4A'aq$ — $LiA'aq$ — $NaA'4aq$ — $KA'3aq$ — CaA' — SrA' — $BaA'6aq$

Ethyl ether EtA' [64°] Yellow plates

(γ)-Iodo-nitro-benzoic acid $C_6H_3I(NO_2)(CO_2H)$ [3x1] [192°] Formed as above V sol water, but does not melt under it — $NaA'aq$ silky yellow needles — $CaA'3aq$ — $SrA'4aq$ — $BaA'3aq$

Iodo-nitro-benzoic acid $C_6H_3I(NO_2)(CO_2H)$ [431] [210°] Formed by heating *p*-iodo benzoic acid with fuming HNO_3 (Glassner, *B* 8, 562) Nearly insol water, v sol alcohol — $NaA'aq$ — $KA'aq$ — $CaA'1\frac{1}{2}aq$

(α, α)-IODO-NITRO-NAPHTHALENE

$C_{10}H_7I_2(NO_2)$ [41] [123°] Formed from acetyl-(α)-naphthylamine by nitration, saponification, diazotisation, and treatment with HI (Meldola, *C J* 47, 519) Minute white needles (from hot alcohol) Sol benzene, HOAc, and acetone May be reduced to (α) naphthylamine

(α, β)-Iodo nitro-naphthalene $C_{10}H_7I_2(NO_2)$ [12] [1085°] Formed from the mother liquor of the preceding (Meldola) Yellow scales May be reduced to (β)-naphthylamine

(β, α)-Iodo-nitro-naphthalene $C_{10}H_7I_2(NO_2)$ [21] [885°] From acetyl (α) nitro (β) naphthylamine by saponifying with hot dilute H_2SO_4 and subsequent displacement of NH_2 by I (Meldola, *C J* 47, 520)

(α)-IODO (β) NITRO-(α) NAPHTHOL

$C_{10}H_7I(NO_2)(OH)$ [421] [146°] Formed by boiling acetyl (4, 2, 1)-iodo nitro-naphthylamine with strong caustic alkali (Meldola, *C J* 47, 524) Yellow needles (from alcohol) Insol cold water, sl sol hot water and hot benzene, v sol alcohol, acetic acid, and acetone Yields phthalic acid on oxidation Its K and Na derivatives form minute orange needles, m sol cold water — $\{C_{10}H_7I(NO_2)\}_2Ba$ 8aq bright red amorphous powder, almost insol. boiling water, deflagrates on ignition

(α)-IODO-(β)-NITRO-(α)-NAPHTHYL-AMINE Acetyl derivative

$C_{10}H_7I(NO_2)(NH_2Ac)$ [421], [286°] From acetyl-iodo naphthylamine by warming its solution in HOAc with HNO_3 at 75° (Meldola, *C J* 47, 524) Stray coloured needles (from alcohol) Sublimes while melting More difficult to saponify by H_2SO_4 than the bromo-compound

IODO NITRO-*o*-OXY BENZOIC ACID

$C_6H_3INO_3$, *ie* $C_6H_4I(NO_2)(OH)(CO_2H)$ From (5, 2, 1) nitro oxy benzoic acid by treatment in alcoholic solution with I and HgO (Weselsky, *A* 174, 108) Slender yellow needles — $KA'2aq$ — $C_6H_5KNO_3$ 3aq — $BaA'6aq$

Iodo nitro *o* oxy benzoic acid

$C_6H_3I(NO_2)(OH)(CO_2H)$ [204°] From (5, 2, 1)-iodo oxy benzoic acid by nitration (Hubner, *B* 12, 1347)

Iodo nitro *m* oxy benzoic acid

$C_6H_3I(NO_2)(OH)(CO_2H)$ Formed by treating an alcoholic solution of nitro *m* oxy benzoic acid with I and HgO (Weselsky, *A* 174, 100) Lemon-yellow crystals, sl sol water, v sol cold alcohol — $BaA'6aq$ minute red needles

Iodo nitro *p* oxy benzoic acid

$C_6H_3I(NO_2)(OH)(CO_2H)$ From nitro *p* oxy-benzoic acid, iodine, and HgO (W) Lemon-yellow needles — $BaA'4aq$ flat red needles $BaC_6H_4INO_2$ 2aq short dark red needles

DI-IODO DI NITRO DI-OXY-DI-PHENYL SULPHONE $C_{12}H_8I_2N_2O_4$, *ie*

$SO_2(C_6H_4I(NO_2)OH)_2$ [205°] Formed by heating iodine (50 g) dissolved in alcohol with a mixture of di-nitro-di-oxy-di-phenyl sulphone (34 g) and HgO (21 g) at 100° (Annaheim, *B* 9, 660) Needles (from HOAc), insol water and alcohol Decomposes alkaline carbonates — $Na_2C_{12}H_8I_2N_2O_4$ 2aq

IODO *o* NITRO PHENOL $C_6H_4INO_3$, *ie*

$C_6H_3I(NO_2)(OH)$ [110°] (Armstrong, *Watts' Dict Ed 1 Suppl* 2, 917, *B* 6, 649) Formed by the action of iodine and HgO on *o*-nitro phenol dissolved in HOAc (Busch, *B* 7, 462, cf Armstrong) Short yellow needles, v sol hot water, alcohol, and ether Moderately volatile with steam — $KA'aq$ garnet red plates with green lustre (A), v e sol hot water and alcohol

Iodo-*p* nitro phenol $C_6H_4I(NO_2)(OH)$ [241],

[93°] (Körner, Armstrong) Formed by the action of iodine and HgO on *p*-nitro phenol dissolved in HOAc (Körner, *Bull Acad Belg* [2] 24, 166, Busch, *B* 7, 462) Also from nitro-amido phenol by the diazo reaction (K) Yellow needles — $NaA'2\frac{1}{2}aq$ (K) — $KA'1\frac{1}{2}aq$ (B) Nitric acid converts it into iodo di-nitro phenol [108°].

Iodo di-nitro phenol

$C_6H_3I(NO_2)_2(OH)$ [6421] [106°] (A), [108°] (W) Formed from (4, 2, 1) di-nitro phenol by treatment with iodine, potassium iodate, and potash (Körner, *G* 4, 397) or with iodine and HgO (Armstrong, *B* 6, 649, Weselsky, *A* 174, 111) Formed also from di-nitro-amido phenol by the diazo-reaction (A), and by the nitration of the preceding iodo nitro phenol (A) Long slender lemon yellow needles (from hot water) Sl sol hot water — KA deep orange needles, v sol hot, sl sol cold, water — $CaA'5aq$ orange red plates, v sol, hot water

Iodo-di-nitro-phenol

$C_6H_3I(NO_2)_2(OH)$ [4621] [113°] Formed from (6, 2, 1)-di-nitro phenol by treatment with iodine

and iodic acid (Körner, *G* 4, 897) or with iodine and HgO (Armstrong, *B* 6, 649) Long, lemon-yellow needles (from water), v sol water and alcohol The potassium derivative forms magnificent crimson needles with green and golden lustre, sl sol water The silver derivative forms small brownish red needles

Di iodo-*o*-nitro-phenol $C_6H_3I_2(NO_2)(OH)$ [6 4 2 1] [98°] From *o* nitro-phenol, iodine, and iodic acid in alkaline solution (Körner) Slender dark yellow needles (from ether alcohol), sl sol water, v sol boiling alcohol and ether—NaA' aq dark brown prisms, with golden lustre—KA' very soluble reddish brown needles

Di iodo *p* nitro phenol $C_6H_3I_2(NO_2)(OH)$ [6 2 4 1] [157°] (K) From *p* nitro phenol, iodine, KIO, and potash (K., cf Seifert, *J pr* [2] 28, 437) Also from *p* nitro phenol *o* sulphonic acid iodine in alcoholic solution, and HgO (Post a Brackebusch, *A* 205, 91) Formed also by treating nitro-*o* oxy benzic acid with iodine and HgO (Weselsky, *A* 174, 177) Large colourless prisms (from ether) turning yellow on exposure to air Decomposes a little above its melting point—KA' golden needles with violet iridescence—NaA' aq efflorescent orange laminae

Di iodo nitro phenols appear also to have been obtained by Piria (*A* 198, 268)

iodo NITRO PHENOL SULPHONIC ACID $C_6H_3I_2NSO_3$ *ie* $C_6H_3I_2(NO_2)(OH)(SO_3H)$ [2 6 1 4] Formed when iodine and HgO are added to an alcoholic solution of *o* nitro phenol *p* sulphonic acid (Armstrong a Brown, *C J* 25, 869)—KA' yellow scales, composed of short needles, sl sol water—K₂C₂H₂INSO₃ red crystals—BaA' aq pale yellow silky needles, sl sol water

Iodo nitro phenol sulphonic acid $C_6H_3I_2(NO_2)(OH)(SO_3H)$ [6 4 1 2] Formed, together with di iodo *p* nitro phenol, by treating an alcoholic solution of *p* nitro phenol *o* sulphonic acid with iodine and mercuric oxide (Post a Brackebusch, *A* 205, 88)—Salts CaC₂H₂INSO₃ 3aq coarse yellow needles, m sol water—BaC₂H₂INSO₃ 3aq long yellow needles, m sol water—Pb₂(OH)₂C₂H₂INSO₃ 2½aq yellow crystalline pp

DI IODO NITRO-RESORCIN $C_6H_2I_2(NO_2)(OH)_2$ From nitro resorcin in alcoholic solution by alternate addition of iodine and mercuric oxide (Weselsky, *A* 174, 111) Golden needles

iodo NITRO-THIOPHENE $C_4H_2I_2(NO_2)S$ [74°] Formed by nitration of iodo thiophene (Kreis, *B* 17, 2073) Glistering yellow prisms

iodo NITRO-TOLUENE $C_7H_4I_2(NO_2)$ *ie* $C_7H_4MeI_2(NO_2)$ [1 4 2] [61°] (286°) From (4,2,1) di-nitro toluene (Heynemann, *A* 158, 397) Pale yellowish crystals V e sol CS₂ and ether

Iodo-nitro-toluene $C_7H_4MeI_2(NO_2)$ [1 4 3] [56°] From *p* toluidine by nitration and displacement of NH₂ by I (Beilstein a. Kuhlberg, *A* 158, 844) Large flat needles, v sol boiling alcohol

Iodo-nitro-toluene $C_7H_4MeI_2(NO_2)$ [1 2 2] [104°] Formed by nitrating *o*-iodo-toluene (B a. K) Minute needles

Iodo nitro-toluene (?) [109°] Formed by nitrating *m*-iodo-toluene (B. a. K.). Small needles.

Iodo di-nitro-toluene $C_7H_3MeI_2(NO_2)_2$ [1 4 5 3] [138°] Formed by nitrating *p*-iodo toluene (Glassner, *B* 8, 561) Crystals

iodo-OCTINOIC ACID $C_{11}H_{11}IO_4$ *ie* $(C_6H_4)ClCO_2H$ Iodo-di-allyl acetic acid from $(C_6H_4)ClCO(OH)CO_2H$ (17 g) and cold fuming HIAq (45 g) (Schatzky, *J R* 17 78) Crystals, insol water, v sol alcohol and ether Easily decomposed

***p* IODO *n*-OCTYL-BENZENE** $C_{18}H_{35}I$ *ie* $C_6H_4(C_6H_4)I$ [1 4] (319°) Heavy oil. Formed by diazotising *p* amido phenyl octane, and treating the diazo-octyl benzene with HI (Beran, *B* 18, 136) On oxidation with CrO₃ it gives *p*-iodo benzoic acid Iodo octyl benzene obtained by the action of iodine and HgO upon *n*-octylbenzene is described by Ahrens (*B* 19, 2720) as an oil, solidifying at -4°, and completely decomposed by distillation

***p*-Iodo sec-octyl-benzene** $C_{18}H_{35}(C_6H_4)I$ [1 4] *p* Iodo capryl-benzene (305° uncor) Formed by diazotising *p* amido phenyl sec octane and treating the diazo capryl benzene with HI Heavy oil. V sol ether and acetic acid, sl sol alcohol It is oxidised by CrO₃ to *p* iodo benzoic acid (Beran, *B* 18, 142)

iodo (a) OCTYL THIOPHENE $C_{11}H_{11}IS$ *ie* $C_6H_4SI(C_6H_4)$ [c 0°] SG $\frac{2}{3}$ 1.2614 From octyl thiophene (10 g) in ligroin by treatment with iodine (10 g) and HgO (11 g) (Schweinitz, *B* 19, 644) V sol ether, cannot be distilled

iodo OLEIC ACID *v* IODO-STEARIDENIC ACID

iodo ORCIN $C_8H_7IO_2$ *ie* $C_8H_7MeI_2(OH)_2$ [86 5°] Formed by dissolving orcin (1 pt) and iodine (2 pts) in ether (6 pts), and gradually adding finely powdered litharge (9 pts) The ether is distilled off, and the residue recrystallised from benzene (Stenhouse, *Pr* 22, 53) Prisms, decomposing when heated alone or with water Sl sol cold water, v sol hot water, alcohol, and ether Has not the astringent sweet taste of orcin

Tri iodo-orcin $C_8H_5I_3O_2$ *ie* $C_8MeI_3(OH)_2$ Formed by adding to a dilute aqueous solution of orcin a quantity of ICl₃, not quite sufficient to ppt all the orcin, and crystallising the pp first from CS₂ and then from boiling alcohol (Stenhouse, *C J* 17, 327) Large transparent plates, tinged with brown V sol CS₂, v e sol ether, m sol alcohol, insol water Becomes brown at 100° Dissolves with decomposition in aqueous KOH Decomposed by hot H₂SO₄ and by hot HNO₃

Iodo-(8)-orcin *v* Iodo BETORCIN

DI-iodo-ORSELLIC ACID Methyl ether $C_8H_7I_2O_4$ *ie* $C_8H_7I_2O_4Me$ Methyl di-iodo-di-oxytoluate Formed by adding a dilute solution of chloride of iodine, containing excess of iodine, to a cold saturated aqueous solution of methyl orsellate (Stenhouse, *A* 149, 295) Needles from CS₂, sol benzene, CS₂, alcohol, and boiling water Decomposed on fusion

Ethyl ether $C_8H_7I_2O_4Et$ Formed in like manner Small needles, almost insol cold water, m sol benzene, CS₂, and boiling alcohol Decomposed on fusion

iodo-*o* OXY-BENZOIC ACID $C_7H_4IO_4$ *ie* $C_7H_3I(OH)(CO_2H)$ [3 2 1] Iodo-salicylic acid. [198°] Formed, together with the following compound, by boiling a solution of salicylic acid

and iodine in alcohol, and separated by fractionally crystallising the product from water (A Miller, *C J* 41, 406, A 220, 124). Long needles, v sol water. Gives a violet colouration with FeCl_3 . Gives the corresponding di oxy benzoic acid on potash fusion— $\text{BaA}'_2 3\frac{1}{2}$ g concentrically grouped needles S 5 at 8° .

Iodo-o-oxy benzoic acid $\text{C}_6\text{H}_4\text{I}(\text{OH})(\text{CO}_2\text{H})$ [5 2 1] [196°] (G), [193 5°] (F), [197°] (M) S 11 at 20° , 1 at 100° (L).

Formation—1 As above—2 Together with di- and tri-iodo oxy-benzoic acids, by fusing salicylic acid (1 mol) with iodine (1 mol), and treating the product with aqueous KOH (Kolbe & Lautemann, A 115, 157)—3 By dissolving equal weights of iodine and salicylic acid in 80 p c alcohol, boiling for 3 hours, evaporating, and dissolving in aqueous Na_2CO_3 (Lautemann, A 120, 299)—4 From salicylic acid, iodine, and HgO (Hlasiwetz & Weselsky, B 5, 380, A 171, 99)—5 By agitating a hot solution of salicylic acid (1 pt) in water (25 pts) with a mixture of iodine (1 pt) and iodic acid ($\frac{1}{3}$ pt), keeping the liquid hot for some time. The resulting acid is converted into sodium salt, and the satiny-needles of the salt of the di iodo acid separated, if necessary by hand picking, from the lancet shaped laminae of the mono iodo acid (Liechti, A Suppl 7, 129, H Fischer, A 180, 346)—6 From the corresponding nitro oxy benzoic acid by reduction to oxy amido benzoic acid and exchange of NH_2 for I by the diazo reaction (Goldberg, J pr [2] 19, 368, P F Frankland, C J 37, 749)—7 By warming dry silver salicylate with iodine (Birnbau & Reinherz, B 15, 458).

Properties—Long needles (from water), sl sol water, v sol alcohol and ether. Gives a violet colour with FeCl_3 . On potash fusion at 200° it gives di oxy benzoic acid [197°]. When suddenly heated it splits up into CO_2 and iodo phenol.

Salts— $\text{NaA}'\text{aq}$ S 77 at 20° (L)— $\text{KA}'_3\text{aq}$ laminae S 19 at 20° (L)— $\text{NH}_4\text{A}'_3\text{aq}$ needles S 95 at 20° (L)— $\text{BaA}'_2\text{aq}$ warty scales S (of BaA'_2) 8 at 8° (M), 13 at 20° (L)— $\text{BaC}_6\text{H}_4\text{IO}_2\text{aq}$ tufts of small needles (L)— $\text{CaA}'_2\text{aq}$ — $\text{MgA}'_2\text{aq}$ — PbA'_2 — AgA' yellowish pp.

Ethyl ether EtA' [71°] (Schmitt, Z 1864, 321).

Iodo-m-oxy-benzoic acid $\text{C}_6\text{H}_3\text{I}(\text{OH})(\text{CO}_2\text{H})$ [x 3 1] Formed, together with di iodo phenol, by alternately adding iodine and HgO to oxy benzoic acid dissolved in 90 p c alcohol, and separated by solution in boiling water (Weselsky, A 174, 105). Slender needles, sl sol cold water.

Iodo-p-oxy-benzoic acid $\text{C}_6\text{H}_3\text{I}(\text{OH})(\text{CO}_2\text{H})$ [3 4 1] [160°] (P) S 18 in the cold.

Formation—1 By boiling a solution of p-oxy-benzoic acid for a few minutes with iodine and iodic acid (Peltzer, A 146, 288)—2 From p oxy benzoic acid dissolved in alcohol by alternate addition of iodine and HgO (Weselsky, A 174, 99).

Properties—Small needles (from water). Decomposes at 152° . May be sublimed below its melting-point V e sol alcohol and ether. Gives with FeCl_3 a dingy brown pp.

Salts— $\text{NaA}'_2\text{aq}$ monochrome efflorescent crystals.— $\text{Na}_2\text{C}_6\text{H}_3\text{IO}_2\text{aq}$. hygroscopic silky

needles (from alcohol)— $\text{BaA}'_2 7\text{aq}$ trimetric tables— AgA' white pp.

Methyl derivative $\text{C}_6\text{H}_3\text{I}(\text{OMe})(\text{CO}_2\text{H})$
Iodo anisic acid [235°] S (cold ether) 7. Formed by heating anisic acid with iodine and iodic acid at 150° (P). Formed also by oxidation of the methyl ether of iodo p cresol (Schall & Dralle, B 17, 2538). The same, or an isomeric acid, is formed from amido anisic acid by the diazo reaction (Griess, *Pr*-10, 309). Plates (S & D) or needles (G). Nearly insol boiling water, m sol alcohol— $\text{NaA}'_2\text{aq}$ concentrically grouped needles— $\text{BaA}'_2 3\text{aq}$ vitreous prisms— $\text{CaA}'_2 3\text{aq}$ nacreous laminae— PbA'_2 (at 100°): curdy pp— AgA' micro-crystalline laminae.

Di-iodo-o oxy-benzoic acid $\text{C}_6\text{H}_2\text{I}_2(\text{OH})(\text{CO}_2\text{H})$
Di iodo salicylic acid S 07 at 15° , 15 at 100° (L). Formed, together with iodo o-oxy-benzoic acid, by the action of iodine and aqueous potash, of iodine and iodic acid, or of iodine and HgO upon salicylic acid (Lautemann, A 120, 304, Liechti, A Suppl 7, 141, Demole, B 7, 1439, Weselsky, A 174, 103). Formed also by heating dry silver salicylate with iodine (Birnbau & Reinherz, B 15, 459). White felted mass (from hot water). V sol alcohol and ether. At 197° (L) or 220° (W) it begins to decompose, giving off iodine. It gives a violet colouration with FeCl_3 . On potash fusion it gives tri oxy benz. ic (? gallic) acid and pyrogallol.

Salts— $\text{NaA}'_2\text{aq}$ long flat needles, mostly grouped in druses S 2 at 20° , much less soluble in water than iodo o oxy benzoic acid— $\text{KA}'_3\text{aq}$ minute thick prisms S 553 at 20° . V sol alcohol, sl sol ether— $\text{NH}_4\text{A}'_3\text{aq}$ arborescent groups of small needles S 32 at 20° — $\text{BaA}'_2 3\text{aq}$ needles S 074 at 18° . V sl sol alcohol— $\text{BaC}_6\text{H}_2\text{I}_2\text{O}_2\text{aq}$ small silky tablets, v sl sol water, forming an alkaline solution— $\text{CaA}'_2\text{aq}$ needles S 086 at 18° .

Di iodo-p oxy benzoic acid
 $\text{C}_6\text{H}_2\text{I}_2(\text{OH})(\text{CO}_2\text{H})$ Formed, together with iodo p oxy benzoic acid, by iodation of p oxy benzoic acid (Peltzer). Small needles (from dilute alcohol). Nearly insol boiling water, v e sol alcohol and ether. Decomposes when heated without previous fusion. Cannot be sublimed— $\text{NaA}'_7\text{aq}$ tufts of delicate, iridescent, efflorescent needles, v sol water— $\text{NaC}_6\text{H}_2\text{I}_2\text{O}_2\text{aq}$ trimetric tables— BaA'_2 gelatinous pp got by adding alcohol to its aqueous solution— $\text{CaA}'_2 2\text{aq}$ nacreous laminae— PbA'_2 bulky pp Swells up like mercuric sulphocyanide when heated— AgA' — $\text{Ag}_2\text{C}_6\text{H}_2\text{I}_2\text{O}_2$.

Tri iodo o oxy benzoic acid
 $\text{C}_6\text{H}_2\text{I}_3(\text{OH})(\text{CO}_2\text{H})$ [c 157°] Obtained in small quantity in preparing mono, and di iodo o oxy benzoic acids by the action of iodine and KOH on salicylic acid (Lautemann). Weselsky (A 174, 104) could not obtain it by treating salicylic acid with iodine and HgO . Tufts of needles (from alcohol). Insol water, sol alcohol and ether. Decomposed by alkalis into CO_2 and a red body $\text{C}_6\text{H}_2\text{I}_3\text{O}$ (?). Its sodium salt is a grey-green mass, v sl sol water.

IODO p-OXY-BENZOIC ALDEHYDE $\text{C}_6\text{H}_3\text{IO}_2$, *i.e.* $\text{C}_6\text{H}_3\text{I}(\text{OH})\text{CHO}$ [199°] Formed by heating the dilute alcoholic solution of p oxy benzoic aldehyde with iodine for some hours (Herzfeld, B 10, 2196). Sl sol. water and benzene, v sol

alcohol and ether. Converted by potash fusion into protocatechuic acid.

Iodo di-oxy-benzoic aldehyde *Methyl derivative* $C_6H_4IO_2$, i.e. $C_6H_4I(OH)(OMe)CHO$ *Iodo vanillin* [174°] Formed by warming vanillin with an alcoholic solution of iodine (Carles, *Bl* [2] 17, 12). Needles. Sol. cold alcohol and ether.

Di iodo-di-oxy-benzoic aldehyde *Methyl derivative* $C_6H_3I_2(OL)(OMe)CHO$ Formed like the preceding, using more iodine (C). Crystals, insol. chloroform, sol. alcohol and ether.

α IODO- β -OXY-PHENYL PROPIONIC ACID $C_9H_7IO_3$, i.e. $C_6H_4CH(OH)CHI CO_2H$ [138°] Formed by treating cinnamic acid with an aqueous solution of iodine chloride, the compound $C_6H_4CHCl CHI CO_2H$ being probably first formed (Erlenmeyer & Rosenhek, *B* 19, 2464). Large crystals (from benzene). With HCl it gives the compound C_6H_4CHClO , which may possibly be $C_6H_4CH(OH)C(OH)C(OH)CHClO$.

Ezo Iodo- α oxy phenyl-propionic acid *Methyl derivative* $C_6H_4(OMe)C_2H_4I CO_2H$ Formed by the combination of the methyl derivative of (a) or (b) coumaric acid with HI (Perkin, *C* J 39, 429). Decomposed by aqueous Na_2CO_3 into CO_2 , HI, and $C_6H_4(OMe)C_2H_4$.

TETRA IODO-DI OXY DI PHENYL SULPHONE $C_{12}H_4I_4SO_2$, i.e. $SO_2(C_6H_4I)_2$ [260°-270°] SG 1.2797. From di oxy di phenyl sulphone by treatment in alcoholic solution with iodine and HgO (Annaheim, *B* 9, 1150). Minute needles (from $HOAc$). Decomposes on fusion. Insol. cold alcohol, nearly insol. boiling alcohol and $HOAc$.

β IODO- α OXY-PROPIONIC ACID $C_6H_4IO_3$, i.e. $CH_2I CH(OH) CO_2H$ *Iodo lactic acid* [101°] (M), [85°] (G). Formed by treating β chloro α oxy propionic acid with KI at 10° (Ginsky, *B* 6, 1257). Prepared by the addition of HI to sodium glycidate (Melikoff, *B* 14, 917). Large prisms, v. sol. water, alcohol, and ether. Converted by alcoholic KOH into glycidic acid— AgA' white unstable pp— ZnA' crystalline powder (M) or tables (G)— CaA' 3sq. amorphous (M).

DI IODO-OXY-PYRIDINE $C_5NH_4I_2(OH)$ [245°] (?) [259°] Formed by heating an alkaline solution of pyridine di carboxylic acid (quinolinic acid) with iodine and KI at 180°-200°. Small flat needles. Sol. hot acetic acid and amyl alcohol, v. sol. boiling water, alcohol, ether, or chloroform. Dissolves in dilute alkalis— $C_5NH_4I_2(ONa)3sq$ colourless glistening scales, sl. sol. aqueous $NaOH$ (Pfeiffer, *B* 20, 1352).

IODO-(Py 3)-OXY QUINOLINE $C_8H_6NOI_2$, i.e. $C_8H_4\begin{matrix} Cl & CH \\ \diagdown & / \\ N & C(OH) \end{matrix}$ or $C_8H_4\begin{matrix} CH & Cl \\ / & \diagdown \\ N & C(OH) \end{matrix}$ *Iodo carbostyryl* [76°] Formed by boiling α amido phenyl propionic acid $C_6H_4(NH_2)C(CO_2H)$ with dilute HI (Bayer & Bloem, *B* 15, 2149). Sublimable.

DI-iodo-DI-OXY TOLUIC ACID v. **DI-iodo-ORSELLIC ACID**

IODO-PENTANE v. **AMYL IODIDE**
Di-iodo-pentane $C_5H_{12}I_2$, i.e. $(CE_2CHI)_2CH_3$ (c 183°). Formed by heating methylene di-

methyl diketone $(CH_3CO)_2CH_2$ with HI at about 90° (Combes, *A Ch* [6] 12, 235). Liquid, begins to decompose at its boiling point.

IODO-PENTINENE $C_5H_4I_2$, i.e. $(CH_2)_2CH C OI$ (140°) Formed from the silver derivative of $(CH_2)_2CH C O H$ by treatment with a solution of iodine in KI (Eltzoff, *J R* 9, 225). Liquid. Gives $(CH_2)_2CH C O H$ when heated with $NaOEt$.

DI-iodo DIPHENIC ACID v. **DI-iodo-DIPHENYL DI CARBOXYLIC ACID**
 α IODO-PHENOL $C_6H_4IO_2$ [2] $C_6H_4I(OH)$ [43°]

Formation—1 Probably occurs among the products of the iodation of phenol (Schutzenberger & Sengenwald, *C R* 54, 197, Körner, *A* 137, 197, Hlasivetz & Weselsky, *Sitz W* 60 [2] 290, Lobanoff, *B* 6, 1251)—2 By the action of iodine on sodium phenol suspended in CS_2 (Schall, *B* 16, 1897)—3 By heating the chloride or sulphate of α diazo phenol with KI, distilling the product with steam, and recrystallising from water (Nöling & Wrzesinsky, *B* 8, 820, Nöling & Stricker, *B* 20, 3018, *Bl* [2] 49, 659, Neumann, *A* 241, 68)—4 By adding powdered iodine to a dilute alcoholic solution of phenol, mixed with ammonia (Willgerodt, *J pr* [2] 37, 446).

Properties—White needles, v. sol. water and other ordinary menstrua. According to Schall (*B* 20, 3363) it forms in crystalline branches, melting at 43° or needles melting at 40°. Nitric acid attacks it, setting iodine free, but chlorine does not do so, but forms chloroiodo phenol. Potash fusion at 165°-250° yields pyrocatechin. Not affected by air and light. Cold conc. H_2SO_4 produces di iodo phenol (Neumann, *B* 20, 581).

m-Iodo phenol [3] $C_6H_4I(OH)$ [40°] Prepared from *m* iodo aniline by diazo reaction. The product is distilled with steam, extracted with ether, and recrystallised from ligroin (N & S). It may be prepared in like manner from *m* amido phenol, by diazotising and heating the product with conc. KI. Needles (from ligroin). May be sublimed. V. sol. alcohol and ether. It does not give off iodine when treated either with HNO_3 or with chlorine. It yields resorcin when fused with KOH.

p Iodo phenol [4] $C_6H_4I(OH)$ [94°]

Formation—1 In the iodation of phenol—2 From *p* amido phenol by diazotising and heating the product with $KIAq$ (N & S)—3 A by product, together with di and tri iodo-phenol, in the formation of *o* iodo phenol from iodine, ammonia, and an alcoholic solution of phenol (Willgerodt, *J pr* [2] 37, 446).

Properties—Long needles. Nitric acid sets iodine free, but chlorine does not do so. When heated with conc. H_2SO_4 it gives (1, 3, 6) di iodo-phenol [72°]. Potash fusion gives resorcin at high temperatures.

Di-iodo-phenol $C_6H_3I_2(OH)$ [421] [72°] Formed by mixing *o* iodo phenol (10 g) with H_2SO_4 (30 g) at -10° (Neumann, *A* 241, 71). Formed also in like manner from *p*-iodo phenol. White needles (from water). Volatile with steam. Boiling HNO_3 converts it into picric acid.

Acetyl derivative $C_6H_3I_2(OAc)$ [76°]. Long flat trimetric prisms (from dilute alcohol); *a.b.c.* = 751 : 1 : 832.

Di-iodo-phenol $C_6H_4I_2(OH)$ [68°] Formed as a by product in the action of iodine upon phenol-sodium suspended in CS_2 (Schall, B 16, 1899, 1902, 20, 3364) Apparently formed also in the action of iodine chloride on phenol (S & S) Wh'rs glistening plates Volatile with steam — $C_6H_4I_2ONa$ needles

Acetyl derivative $C_6H_4I_2(OAc)$ [107° uncor], small prisms

Benzoyl derivative $C_6H_4I_2(OBz)$ [96° uncor]

Di iodo phenol $C_6H_4I_2(OH)$ [160°] Formed by the action of iodine and HgO on an alcoholic solution of phenol (Hlasiwetz & Weselsky, *Satz W* 60 [2] 290, C C 1870, 63) Silky felted needles (from dilute alcohol), v sol alcohol, ether, and CS_2 . May be sublimed Not affected by boiling alcoholic KOH

Tri iodo-phenol $C_6H_3I_3(OH)$ [6 4 2 1] [155°]

Formation—1 By treating phenol with a considerable quantity of ICl , and exhausting the product with boiling alcohol in which tri iodo-phenol is not very soluble (Schutzenberger, *Bl* [2] 4, 102)—2 From phenol, iodine, and KOH (Lautemann, A 120, 307)—3 From phenol, iodine, and iodic acid (Körner, A 137, 213)—4. From salicylic acid, iodine and iodic acid (Kekulé, A 131, 231)—5 Formed as a by product from the action of iodine on phenol sodium suspended in CS_2 (Schall, B 16, 1899)—6 By the action of iodine of nitrogen on a dilute alcoholic solution of sodium phenolate (Willgerodt, *J pr* [2] 37, 447)

Properties—Needles (from dilute alcohol) Decomposed on fusion M sol alcohol Converted by excess of ICl into per chloro-phenol HNO_3 gives picric acid HCl and $KClO_3$ gave chloranil

DI-IODO-PHENOL SULPHONIC ACID

$C_6H_3I_2(OH)(SO_3H)$ [6 2 1 4] [120°] Formed by adding a solution of KI and KIO_3 to one of phenol *p*-sulphonic acid in aqueous HCl (Kehrmann, *J pr* [2] 37, 11, 334, 359, Osteryayer, *J pr* [2] 37, 215) Monoclinic prisms c sol water, but ppp by HCl or H_2SO_4 , decomposes at 190° Converted by nitric acid to picric acid Chromic acid oxidises it to di iodo quinone

Salts— KA 2aq long needles or thick prisms— NaA 3aq sol water, commercially known as 'sodazole'— $C_6H_3I_2K_2SO_3$ dimetric prisms— BaA , 3aq shining needles, v sl sol water— ZnA , 6aq long needles The copper salt is greenish white and v sol water

DI-p-IODO-DIPHENYL $C_{12}H_8I_2$

$C_{12}H_8I_2$ [202°] Formed by the action of HI on the diazo-compound derived from benzidine (Schmidt & Schultz, B 12, 489) Yellow leaflets

o-IODO-PHENYL-ACETIC ACID $C_8H_7IO_2$

$C_8H_7IO_2$, CO_2H [96°] Prepared by digesting its nitrile with fuming HCl for four hours at 126° The nitrile is formed by treating o-iodo-benzyl bromide with KCy (Mabery & Robinson, *Am* 4, 102) Slender needles (from water), sol hot water, alcohol, ether, CS_2 , and ligroin— AgA curdy pp; sl sol water

p-Iodo-phenyl-acetic acid $C_8H_7IO_2$, CO_2H [186°] Prepared by heating its nitrile with fuming $HClAq$ in sealed tubes at 100° (Jackson, *Mabery, Am* 2, 253, *P Am* A 18, 205, B 11, 56). Narrow white plates (from water) with

agreeable odour like sweet allysum, v sol hot water, alcohol, ether, benzene, and CS_2 . Gives *p*-iodo benzoic acid on oxidation— BaA , aq minute white needles, v sol water— AgA glistening plates, sl sol boiling water

Nitrile $C_8H_6I_2CH_2CN$ *p* Iodo benzyl cyanide [50 5°] Prepared by boiling *p*-iodo benzyl bromide with alcoholic KCy Pearly plates insol water, v sol alcohol, ether, and CS_2

IODO PHENYL-ACETYLENE C_8H_5Cl Obtained from phenyl acetylene (Holleman, B 20, 3080) Brownish-yellow liquid

IODO-PHENYL-ACRYLIC ACID v IODO CRYNACIC ACID

Ezo-IODO-DI-PHENYL-AMYLIDENE-DIAMINE $C_{12}H_{11}N_2I$ & $C_{12}H_9I(NHPh)_2$ Formed by heating iodo isovalic c aldehyde with aniline (Chautard, A Ch [6] 16, 169) Large brownish yellow needles, v sol water and alcohol, m sol other solvents Decomposed on fusion Forms uncrystallisable salts

DI-IODO-DIPHENYL-DI-CARBOXYLIC ACID $C_{12}H_8I_2O_4$ & $C_{12}H_8I_2(CO_2H)_2$ $C_{12}H_8I_2CO_2H$ [262°] Prepared by the action of HI on the diazo-derivative of di amido di phenyl dicarboxylic acid (Schultz, B 11, 217, A 196, 21, 203, 95) Amorphous, v sl sol boiling water, v sol alcohol, acetone, and ether

DI IODO PHENYLENE OXIDE so called $C_{12}H_8I_2O$ Tetra iodo diphenylene quinone Formed in the action of iodine and KOH upon *o*, *m*, or *p* oxy benzoic acid and on phenol (Lautemann, A 120, 309, Kammerer & Benzing, B 11, 557) Formed also by boiling tri iodo phenol with aqueous $Na_2CO_3(L)$, and by distilling di iodo diphenyl dicarboxylic acid with lime (Schultz, B 11, 217) Reddish brown powder, insol water, alcohol, ether, chloroform, and benzene Dissolves in CS_2 with deep red colour Boiling $KOHAq$ has little action on it Conc HNO_3 forms picric acid It decomposes at 200° Aqueous SO_2 at 100° forms colourless $C_{12}H_8I_2O_2$

DI-EZO IODO-DI-PHENYL-ETHYLENE

$C_{12}H_{10}I_2$ & $C_{12}H_8I_2$ CIPh CIPh Formed by heating di-phenyl acetylene (tolane) with dry iodine and crystallising from chloroform (E Fischer, A 211, 238) Rose coloured plates, v sl sol alcohol Decomposed by heat

DI IODO DI-PHENYL-ETHYLIDENE-DIAMINE $C_{12}H_{11}IN_2$ & $C_{12}H_9I_2(NHPh)_2$ Obtained by heating iodo acetic aldehyde with excess of aniline (Chautard, A Ch [6] 16, 155) Yellow needles or tables Cannot be melted without decomposition V sol alcohol, sol ether, benzene and CS_2 With strong acids it forms uncrystallisable salts, v sol water and alcohol

DI-p-IODO DI-PHENYL-GUANIDINE

$C_{12}H_{11}I_2N_3$ & $CN_2H_2(C_6H_4I)_2$? From *p*-iodo-aniline in ethereal solution and $CyCl$ (Hofmann, A 67, 148) Crystalline— $B_2H_3PtCl_4$

Tri-p-Iodo-tri-phenyl-guanidine $C_{18}H_{15}I_3N_3$ From di *p*-iodo di phenyl thio-urea and iodine (Losanitsch, B 5, 158)

Ezo-IODO-DI-PHENYL-HEPTYLIDENE-DIAMINE $C_{18}H_{19}I_2(NHOC_2H_5)_2$ Formed by heating iodo heptioic aldehyde with aniline (Chautard, A Ch [6] 16, 178) Lemon yellow deliquescent tables, v sol alcohol, sl sol benzene and $CHCl_3$ Decomposes on heating Does not give crystalline salts

p-iodo-phenyl-hydrazine $C_6H_4I.NH_2$, ϵ $C_6H_4I.NH.NH_2$ [10°] Formed, by reducing the diazo-derivative of p-iodo aniline [60°] with an excess of $SnCl_2$ (Neufeld, A 248, 98) Silky needles (from hot water), v sol alcohol, ether, chloroform, benzene, and dilute H_2OAc With acetone it forms $C_6H_4I.N_2H.CMe_2$ crystallising from petroleum ether in white plates [114°] With acetic aldehyde it gives $C_6H_4I.N_2H.CHMe$ [107°] crystallising from petroleum ether in yellow needles

Di-iodo-phenyl-hydrazine $C_6H_3I_2(NH_2)_2$ [4 2 1] [112°] From di-iodo aniline by diazotising and reducing with $SnCl_2$ (Neufeld, A 248, 99) White silky needles, v sol alcohol, ether, and benzene, sl sol hot water and petroleum ether The hydrochloride [162°] crystallises from water in white needles

Di-iodo-s di-phenyl hydrazine $C_6H_4I_2.NH_2$, ϵ [8 1] $C_6H_4I.NH.NH.C_6H_4I$ [1 3] *Hydrazo di-iodo benzene* [90°] Formed by heating the azoxy compound $C_6H_4I.N_2O.C_6H_4I$ with alcoholic ammonium sulphide at 100° (Gabriel, B 9, 1408) V sol ordinary solvents

Di-iodo-s di-phenyl hydrazine [4 1] $C_6H_4I.NH.NH.C_6H_4I$ [1 4] Formed by heating the corresponding azoxy compound with alcoholic ammonium sulphide at 100° (G) Flat needles, decomposed before melting

p-iodo-phenyl oxamic acid $C_6H_4I.NH.CO.CO_2H$ [c 200°] S 072 at 25° Formed by the action of alcoholic potash on the di-iodo di-anilide of oxalic acid (Dyer a Mixer, Am 8, 357) Long white fibres Sol alcohol and ether — BaA' white pp — AgA' white pp — KA' sol hot water

p-iodo-phenyl phthalimide
(1) $C=O$
 $C_6H_4 \begin{matrix} \diagup \\ \diagdown \end{matrix} \begin{matrix} O \\ O \end{matrix}$
(2) $C=N.C_6H_4I$ [227°–228°] Prepared by heating p-iodo aniline with phthalic anhydride (Gabriel, B 11, 2261) Needles V sol C_6H_6 , less sol ether

o-iodo-phenyl propionic acid $C_6H_4IO_2$, ϵ $C_6H_4I.C_2H_4.CO_2H$ *o-iodo hydrocinnamic acid* [103°] Plates (from water) Formed by reduction of o-iodo cinnamic acid (Gabriel a Herzberg, B 16, 2037)

m-iodo-phenyl propionic acid $C_6H_4I.C_2H_4.CO_2H$ *m-iodohydrocinnamic acid* [68°] Colourless plates Formed by reduction of m-iodo-cinnamic acid (Gabriel a Herzberg, B 16, 2039)

p-iodo-phenyl-propionic acid $C_6H_4I.C_2H_4.CO_2H$ *p-iodo hydrocinnamic acid* [141°] White prisms Formed by reduction of p-iodo-cinnamic acid with HI and P (Gabriel a Herzberg, B 16, 2040)

β -iodo-phenyl-propionic acid $C_6H_4I.CH_2.CO_2H$ [120°] Formed by adding fuming $HIaq$ to a concentrated aqueous solution of β oxy phenyl propionic acid (Glaser, A 147, 97) Formed also from cinnamic acid and conc $HIaq$ in the cold (Fittig a Binder, A 195, 188) Small crystals (from CS_2) Decomposed on fusion Boiling with water converts it into HI and cinnamic acid Aqueous Na_2CO_3 gives HI, styrene, and CO_2

o-iodo-di-phenyl-propylidene-di-amine $C_6H_4I_2.N_2$, ϵ $CH_2I.CH_2.CH(NHPh)_2$

Formed by heating β -iodo propionic aldehyde with excess of aniline (Chautard, A Ch [6] 16, 159) Brownish yellow needles or tables, v sol all ordinary solvents Decomposed by heat without previous fusion Gives no crystalline salts

Di-iodo-di-phenyl sulphide $C_6H_4I_2S$, ϵ $(C_6H_4I)_2S$ [139°] Formed by heating di-phenyl sulphide with iodine and iodic acid in sealed tubes Got also from di-amido di-phenyl sulphide by the diazo reaction (Krafft, B 7, 1165) Lamine

p-iodo-phenyl thiocarbimide $C_6H_4I.NCS$ [65°] Formed by the action of iodine on an alcoholic solution of di-p-iodo di-phenyl thio urea (Losantsch, B 5, 158) Crystals

Di-p-iodo-di-phenyl-thio urea $C_6H_4I_2.N_2S$, ϵ $CS(NH.C_6H_4I)_2$ [173°] From p-iodo aniline, alcohol, and CS_2 (Losantsch, B 5, 157) V sol sol alcohol and ether

iodo-iso-phthalic acid $C_6H_4I(CO_2H)_2$ [6 3 1] [204°] Formed by oxidation of iodo-tolyl methyl ketone $C_6H_4MeI.CO.CH_3$ [1 2 5] with CrO_3 (Klingel, B 18, 2701) Fine white needles Sublimable V sol alcohol, acetic acid, and ether, insol cold water

Salts — $A'Ba$ very sparingly soluble fine white needles — $A'Ca$ microscopic needles — $A'Ag$ white crystalline pp — $A'Cu$ green crystalline pp

iodo propane v propyl iodide

Di-iodo-propane $CH_2I.CH_2.CH_2I$ *Propylene iodide* SG 2 249 From propylene and iodine (Berthelot a De Luca, C R 39, 744) From allyl iodide and gaseous HI at -18° (Malbot, C R 107, 114, B [2] 50, 449) Liquid Split up by alcoholic potash into iodine and propylene

Di-iodo-propane $CH_2I.CH_2.CH_2I$ *Tri-methylene iodide* (227°), (169° at 170 mm) SG 2 2681, d_4^{20} 2 5962, n_D^{20} 2 5614 Obtained by heating s di oxy propane (trimethylene glycol) with fuming $HIaq$ at 100° (Freund, M 2, 640) Also from trimethylene bromide, alcohol, and KI (Perkin, jun, C J 51, 12)

Di-iodo-propane $CH_2I.CH_2.CH_2I$ *Allylene dihydro-di-iodide* (148°) SG 2 215 (O), 2 45 (S) Obtained by direct combination of allylene with HI (in conc solution) (Oppenheim, Bl [2] 4, 434, Semenoff, Bl [2] 5, 446) Heavy oil Partly decomposed by distillation, but may be distilled with steam or with any indifferent gas Sl sol alcohol, v sol ether Becomes coloured in air and light Alcoholic potash forms $CH_2I.CI.CH_2$ Ag_2O gives acetone (Sorokin, Z 1871, 264)

iodo-PROPIOLIC ACID $C_3H_4IO_2$, ϵ $ICCO_2H$ *Iodo propargylic acid Iodo-acetylenic carboxylic acid* [140°] Small colourless prisms or glistening needles

Formation — 1 By saponification of the ethyl ether — 2 By the action of a solution of iodine in KI upon the cuprous compound of calcium or barium propargylate

Reactions — It combines with HBr to form brom-iodo acrylic acid [96°], with HI it yields di-iodo acrylic acid [133°], with bromine dissolved in chloroform it yields di-bromo-iodo-acrylic acid [147°], with an ethereal solution of iodine, tri-iodo-acrylic acid [207°], and with iodine bromide, bromo-di-iodo acrylic acid [182°].

On the other hand, an aqueous solution of bromine converts it into di-bromo iodo ethylene IBrC CHBr with evolution of CO_2 .

Salts— A^+K^- small glistening very hygroscopic needles— A^+Ag^- white pp, decomposed by warming with water forming AgI — A^+Ba^- easily soluble amorphous solid— A^+Cu^-

Ethyl ester A^+Et^- [68°], large colourless prisms Formed by the action of a solution of iodine in KI upon the cuprous compound of propargylic ether (Baeyer, *B* 18, 2274, Homolka a. Stolz, *B* 18, 3282, 19, 536)

α -IODO PROPIONIC ACID $\text{C}_3\text{H}_5\text{IO}_2$, \pm

$\text{CH}_3\text{CHI CO}_2\text{H}$ Prepared by digesting syrupy lactic acid (1 mol) with PI_2 (1 mol), pouring the product into water and extracting with ether (Wichelhaus, *A* 144, 352) Oil, nearly insol water

β -Iodo-propionic acid $\text{CH}_3\text{I CH}_2\text{CO}_2\text{H}$ [82°]

Formation—1 By the action of iodide of phosphorus on glyceric acid (Beilstein, *A* 120, 226, 122, 366, Erlenmeyer, *A* 191, 284)—2 From acrylic acid and conc HIAg (Wislicenus, *A* 166, 1)

Preparation—The product of the oxidation of glycerin with nitric acid is evaporated on the water-bath, and the syrupy residue diluted with water until it possesses S.G. 1.26 This solution (30 c c) is then poured into a flask containing iodine of phosphorus prepared from iodine (50 g) and yellow phosphorus (6.5 g) After the vigorous action has taken place, the solid residue is recrystallised from water (V Meyer, *B* 19, 3294, 21, 24)

Properties—Colourless laminae, v sol hot, v sl sol cold, water, v e sol alcohol and ether May be converted into propionic acid by HI or by sodium amalgam (Moldenhauer, *A* 131, 328) By boiling with water it is converted into hydroacrylic acid and a little acrylic acid Boiling with water and Ag_2O forms hydroacrylic acid, para adipic acid $\text{C}_6\text{H}_{10}\text{O}_4$, dihydroacrylic acid $\text{C}_4\text{H}_6\text{O}_3$, and paracrylic acid $\text{C}_4\text{H}_6\text{O}_2$ [69°] (Beilstein, Wislicenus, *Z* [2] 4, 683, Socloff, Klumenko, *Bl* [2] 34, 321, v also ACRYLIC ACID) β -Iodo propionic acid is converted into adipic acid by heating with reduced silver (W) AgNO_3 gives in an aqueous solution of β iodo propionic acid an immediate pp of AgI

Methyl ether MeA^+ (188°) S.G. 1.841 (Henry, *C* R 100, 114) Colourless oil, turning brown in light Is not pungent Formed from the acid, alcohol, and H_2SO_4

Ethyl ether EtA^+ (202°) (Fittig a. Wolff, *A* 216, 128) S.G. \pm 1.707 (Henry) Formed by heating an alcoholic solution of the acid after adding a few drops of H_2SO_4 Formed also by dissolving the acid (1 pt) in alcohol (3 pts) and saturating with HCl , the yield is only 50 p c, for much EtI is evolved (Lewkowitsch, *J pr* [2] 20, 167; Wichelhaus, *B* 1, 25, Wislicenus, *A* 192, 129) Oil Partly decomposed on distillation

Amide $\text{CH}_3\text{I CH}_2\text{CONH}_2$ [101°] Formed by the action of aqueous NH_3 on the methyl ether in the cold (Henry) Colourless tables, turning yellow in light V sol water Its solution gives a pp. with AgNO_3

β -IODO PROPIONIC ALDEHYDE

$\text{CH}_3\text{I CH}_2\text{CO H}$ S.G. 1.221 Prepared by the action of iodine (25 g.) and iodic acid (10 g) on propionic aldehyde (18 c c) diluted with alcohol

(50 c c) It is washed with KOH solution and poured into water (Chautau d, *A Ch* [6] 16, 157).

Properties—Heavy colourless liquid, not inflammable, with an exceedingly irritating vapour Miscible with alcohol, ether, and acetone, sl sol water Is totally decomposed at 100° Very dilute solutions of KOH , NaOH , or NH_4HO attack it slowly in the cold Conc solutions resinify it Mineral acids behave similarly HNO_3 oxidises it to $\text{CH}_3\text{I CH}_2\text{CO}_2\text{H}$ [82°] $\text{Ag}(\text{C}_2\text{H}_5\text{O}_2)$ forms propyl acetate (90°) AgCN and silver sulphocyanide give $\text{CH}_2(\text{CN})\text{CH}_2\text{CO H}$ and $\text{CH}_2(\text{CNS})\text{CH}_2\text{CO H}$ Aniline gives the compound $\text{CH}_3\text{I CH}_2\text{CH}(\text{N}^+\text{C}_6\text{H}_5)_2$.

IODO PROPYL ALCOHOL v Iodhydrin of PROPYLENE GLYCOL

Di iodo-propyl alcohol $\text{C}_3\text{H}_5\text{I}_2$, \pm
 $\text{CH}_3\text{I CHI CH}_2\text{OH}$ *Di iodide of allyl alcohol* [45°] Prepared by dissolving iodine in allyl alcohol (Hubner a. Lellmann, *B* 13, 460, 14, 207) Colourless needles Easily decomposed by light or heat Sol alcohol, ether, and benzene, insol water On heating the chloroform solution it gives a compound crystallising in colourless prisms [160°], which is probably an iodo allyl alcohol

Di iodo isopropyl alcohol $\text{CH}_3\text{I CH}(\text{OH})\text{CH}_2\text{I}$
Glycerin di iodhydrin [α - 18°] S.G. 1.24 Prepared by heating the dichlorhydrin of glycerin with KI and water on a salt bath (Nahmacher, *B* 5, 353, Claus, *A* 163, 24) Faintly yellowish oil Decomposes when distilled

p-IODO PROPYL-BENZENE $\text{C}_6\text{H}_4(\text{C}_2\text{H}_4)\text{I}$ [14] [250°] Formed by heating p diazo propyl benzene with HI (Louis, *B* 16, 110) Volatile with steam Colourless oil Sol ether, m sol alcohol, almost insol water On oxidation with cold CrO_3 in acetic acid it gives p iodo benzoic acid

p Iodo iso propyl benzene $\text{C}_6\text{H}_4(\text{C}_2\text{H}_4)\text{I}$ (234°) Prepared by heating p diazo iso propyl benzene with HI (Louis, *B* 16, 114) Colourless oil On oxidation it gives p iodo benzoic acid

α -IODO PROPYLENE v ALLYL IODIDE

α Iodo propylene $\text{CH}_3\text{CHI CH}_2$, *Allylene hydro iodide* (82°) (S), (93°-103°) (O) S.G. 1.835, 1.803 Formed by decomposing di iodo propane $\text{CH}_3\text{CI}_2\text{CH}_2$ (1 mol) with alcoholic KOH (1 mol), and mixing the distillate with water (Semenoff, *Bl* [2] 5, 446, *Z* 1865, 725, Oppenheim, *Bl* [2] 4, 434, *Z* 1865, 719)

Di iodo propylene $\text{C}_3\text{H}_4\text{I}_2$, \pm $\text{CH}_2\text{CI CHI CH}_2$ *Allylene di iodide* (198°) Obtained by exposing allylene for two months to a solution of I in aqueous KI in sunshine (Oppenheim) Oil. Turns brown in light Yields allylene iodo alcohol KOAc

Tri iodo propylene $\text{CHI CI CH}_2\text{I}$ *Di iodo-allyl iodide Propargyl tri iodide* [41°] Formed by the combination of propargyl iodide with iodine (Henry, *B* 17, 1132) Small colourless needles

Tri iodo-propylene $\text{CH}_2\text{CI CI}_2$ *Iodo-allylene di iodide* [64°] From silver allylene and iodine in ethereal solution (Liebemann, *A* 135, 273) Needles Decomposes at 78° V e sol ether, m sol alcohol. Alcoholic potash gives iodo-allylene

IODO-PROPYLENE-GLYCOL v Iodhydrin of GLYCERIN.

IDO-PROPYL-THIOPHENE C_8H_7S \approx C_8H_7S C_8H_7 . From α propyl thiophene (Rufen, B. 20, 1743) Oil, volatile with steam

Ido-IDO PYROTARTARIC ACID $C_8H_7IO_4$ [185°] From itaconic acid an¹ HI at 150° (Swarts, Z 1866, 722) Nodules Reduced by further treatment with HI to pyrotartaric acid

TETRA IODO PYRROLE C_4I_4NH 'Iodol' S 02, S (90 p.c. alcohol) 6 at 15°, S (ether) 60 Formed by the action of an ethereal solution of iodine on pyrrole potassium (Ciamician a Dennstedt, G 13, 18, B 15, 2582), and by the action of iodine on an alkaline solution of pyrrole (Ciamician a Silber, G 16, 543, B 18, 1766) Long yellowish brown flat prisms or minute yellow needles (from dilute alcohol) Sol ether, acetic acid, and hot alcohol, nearly insol cold alcohol, insol water and aqueous acids Decomposes at about 140°-150° Gives a white ppt with Ag_2O , instantly blackening, green colouration with $HgCl_2$. Has no basic properties It is not decomposed by boiling water, but boiling HCl aq blackens it It does not dissolve in aqueous KOH, but alcoholic potash dissolves it forming a potassium derivative which is decomposed by acids (even CO) but not by water Zinc dust and KOH reduce it to pyrrole (Ciamician a Silber, B 19, 3027) When heated gently with conc H_2SO_4 it gives at first a green, then a dirty violet colouration Its alcoholic solution gives a red colour with nitric acid (Vulpinus) It is employed pharmaceutically as a substitute for iodoform for suppurating sores, fungoid growths, hay fever, &c, having the advantage of possessing no odour, and exerting no poisonous effect upon the system

IDO-QUINOLINE C_8H_5IN [63°] (above 300°) SG 1.93 Prepared by heating quinoline with a KI solution of iodine and HIO_4 (La Coste, B 13, 780) Monoclinic prisms, or long thin needles. Easily volatile with steam

Salts - $B'HCl$ aq small yellow needles - $B'H PtCl_2$ 2aq long orange needles, sl sol water - $B'H C_2O_4$, sparingly soluble yellow needles or plates

Methylo iodide $B MeI$ glistening golden plates, sol hot water v sl sol cold water and alcohol, insol ether

Methylo chloride $B' MeCl$ aq fine yellowish needles or thick yellow prisms, dimorphous (Lehmann, Z K 12, 377), v sol water - ($B' MeCl$), $PtCl_4$ fine orange crystals, sl sol cold water

(Py 3) Iodo quinoline C_8H_5 $\begin{matrix} CH \\ | \\ N \\ | \\ Cl \end{matrix}$ [53°]

Formed by heating (Py 3) chloro quinoline with HI (Friedlander a Weinberg, B 18, 1531) White needles Sl sol water, v sol other solvents - BHI long needles - $B'H_2Cl_2PtCl_4$ aq red needles

Di-iodo-quinoline $C_8H_4I_2N$ [90° uncor] Formed by the action of iodine in CS_2 on quinoline (Claus a Istel, B 15, 824), Dull-green metallic needles Sol alcohol, ether, and acetic acid

DI IODO-QUINONE $C_6H_4I_2O_2$ [159°] Formed by the action of iodic acid on the diacetyl derivative of hydroquinone (Metzler, B 21, 2555) Yellow needles (from alcohol) Is converted by

the action of sulphurous acid into the hydroquinone [142°]

Di iodo quinone $C_6H_2I_2O_2$ [3541] [180°] From di-iodo p amido-phenol (q v), H_2SO_4 , and $K_2Cr_2O_7$ (Seiffert, J pr [2] 28, 438) Formed also by oxidising di iodo phenol sulphonic acid with CrO_3 (Kehrmann, J pr [2] 37, 336) Golden plates (from benzoline) Readily sublims, almost insol cold, v sl sol boiling water With HCl and $SnCl_2$ it gives di iodo hydroquinone [145°] $FeCl_3$ reconverts this into di-iodo quinone

Di iodo quinone chlorimide $C_6H_2I_2$ $\begin{matrix} <O \\ | \\ NCl \end{matrix}$ [123°] From $C_6H_4I_2(NH_2)(OH)$, hydric chloride, and bleaching powder solution (Seiffert, J pr [2] 28, 438) With NMe_3 it gives

$C_6H_2I_2$ $\begin{matrix} <O \\ | \\ N \end{matrix}$ $>C_6H_4NMe_2$

IDO RESORCIN $C_6H_4IO_2$ \approx $C_6H_4I(OH)_2$ [67°] Formed by agitating an ethereal solution containing resorcin (10 pts) and iodine (24 pts) with PbO (110 pts), distilling off the ether and recrystallising from benzene (Stenhouse, C N 26, 279, A 171, 311). Trimetric prisms, sol water

Tri iodo resorcin $C_6H_3I_3(OH)_2$ [145°] (M a N), [154°] (C) Formed, together with a brown substance insoluble in CS_2 , when ICl is added to an aqueous solution of resorcin (Michael a Norton, B 9, 1752) Formed also by adding an aqueous solution of resorcin to a solution of KIO_3 , iodine, and KI (Claassen, B 11, 1443). Prepared by adding bleaching powder to a dilute alkaline solution of resorcin (1 mol) and KI (7 mols), and then acidifying (Degener, J pr [2] 20, 324) Needles (from CS_2), sl sol hot water, v sol alcohol and ether May be sublimed. Boiling HNO_3 gives tri nitro resorcin

Di acetyl derivative $C_6H_3I_3(OAc)_2$ [170°] Needles, v sol alcohol and ether

IDO RESORCIN SULPHONIC ACID $C_6H_3I_3(OH)_2(SO_3H)$ From potassium resorcin sulphionate and iodine (H Fischer, M 2, 340) - KA' 3aq minute crystals

Iodo resorcin disulphonic acid $C_6H_2I_3(OH)(SO_3H)_2$ Formed by digesting potassium resorcin disulphonate (30 g) with iodine (33 g) in dilute alcoholic solution at 100° (F) - KA long needles

IDO SALICYLIC ACID v Iodo o oxy-BENZOID ACID

IDO STEARIC ACID $C_{18}H_{33}IO_2$ From di oxy stearic acid and HI (A Saytzeff, J pr [2] 33, 309) Oil

Iodo stearic acid $C_{18}H_{33}IO_2$ From oleic acid, PI_3 , and a little water The product is mixed with water and extracted with ether (M C a A Saytzeff, J pr [2] 35, 384)

Properties - Oil Resembles the preceding acid

Reactions - 1 Reduced by Zn and HCl to stearic acid - 2 Moist Ag_2O forms a substance that is unsaturated (taking up 55 p.c I from its solution in $HgCl_2$ aq), but is converted by alcoholic KOH into oxy stearic acid - 3 Alcoholic KOH forms solid iso oleic acid [45°], and also ordinary oleic acid

Iodo-stearic acid $C_{18}H_{33}IO_2$ Formed by heating iso oleic acid with tri iodide of phosphorus and water (Coust a Saytzeff, J pr [2] 37, 276, Bk [2] 47, 169) Oil, sol ether.

Reactions—1. Yields an oxy stearic acid [85°] on treating with silver oxide—2 Alcoholic potash regenerates iso-oleic acid [40°-45°], but forms no oleic acid—3 Alkaline KMnO_4 oxidises it to di-oxy-stearic acid [78°]

IODO-STEARIDENIC ACID $\text{C}_{18}\text{H}_{33}\text{IO}_2$ *Iodo-oleic acid* From ricinoleic acid $\text{C}_{18}\text{H}_{33}\text{O}_2$, water, P, and iodine at 100° (Claus, B 9, 1917) Oil Reduced by boiling with zinc and HClAg to stearic acid Combines with bromine

DI-IODO-SUCCINAMIC ACID

$\text{CO}_2\text{HCH}_2\text{CH}_2\text{CONH}_2$

Ethylether A'Et [134°], long needles, slightly soluble in cold water Formed by the action of iodine upon diazo succinamic ether $\text{CO}_2\text{EtCH}_2\text{CH}_2\text{ON}_2$ in ethereal solution (Curtius & Koch, B 19, 2462, J pr [2] 38, 485)

IODO-TARCONINE v NARCOTINE

IODO-THIENYL METHYL KETONE

$\text{C}_6\text{SH}_4\text{COCH}_3$ *Iodo-acetothienone* [129°] Formed by the action of acetyl chloride upon mono- or di iodo thiophene in presence of AlCl_3 (Gattermann & Römer, B 19, 692) Long colourless needles (from alcohol) Very volatile with steam Strong odour By alkaline KMnO_4 it is oxidised to iodo thiophene carboxylic acid [131°]

Phenyl hydrazide $\text{C}_6\text{SH}_4\text{C(NHPh)CH}_3$ [134°], yellow tables, sl sol alcohol

(a) IODO-THIOPHENE $\text{C}_4\text{H}_3\text{IS}$: s

$\text{CHCH} \rangle \text{S}$ (182° cor). Oil Formed by the action of iodine and HgO on thiophene at the ordinary temperature (Meyer & Kreis, B 17, 1558, Egh, B 18, 544) With sodium and EtI it gives ethyl thiophene With Na and n butyl bromide it forms n -butyl thiophene (182°) Gives an iodo-nitro thiophene [74°] (Kreis, B 17, 2078)

Di-iodo-thiophene $\text{C}_4\text{H}_2\text{I}_2\text{S}$ [40½°] White crystals Formed by the action of 2 mols of iodine and HgO on thiophene at the ordinary temperature (Meyer & Kreis, B 17, 1558)

IODO-THIOPHENE CARBOXYLIC ACID

$\text{C}_6\text{SH}_4\text{I(COOH)}$ *Iodo thiophenic acid* [131°] Formed by oxidation of iodo thienyl methyl ketone with alkaline KMnO_4 (Gattermann & Römer, B 19, 693) Colourless silky needles (from water) Sublimes in glistening tables— $\text{NH}_4\text{A'}$ needles, sl sol cold water

(a)-IODO-THIOPHENE-(β)-DI SULPHONIC ACID $\text{C}_4\text{HIS(SO}_3\text{H)}_2$ Formed by sulphonation of (a)-iodo thiophene On reduction with sodium amalgam it yields thiophene (β) di sulphonic acid (Langer, B 18, 559)

IODO-THIOXENE is IODO DI METHYL THIOPHENE (q v)

IODO-THYMOL $\text{C}_{10}\text{H}_7\text{MePr(OH)I}$ [1436] [69°] Prepared by adding iodine (8 g) to a solution of thymol (5 g) in ammonia (6 cc) mixed with alcohol (2 cc) and distilling the only product with steam (Willgerodt & Kornblum, J pr [2] 39, 289) Lustrous white needles, insol water, sol other solvents Oxidised by MnO_2 and H_2SO_4 or by FeCl_3 to thymoquinone Not attacked by aqueous KOH at 800° HNO_3 forms nitro thymol, displacing I by NO_2 H_2SO_4 forms a sulphonic acid.

Ethyl derivative $\text{C}_{12}\text{H}_{15}\text{MePrI(OEt)}$ [52°] Opaque white rimetric tables, insol cold water, sl sol hot water and alcohol

Acetyl derivative $\text{C}_{12}\text{H}_{15}\text{MePrI(OAc)}$

[71°] White needles

Benzoyl derivative $\text{C}_{12}\text{H}_{15}\text{MePrI(OBz)}$

[95°] Large tables

Acryl derivative [155°], yellowish crystalline aggregates

IODO THYMOL SULPHONIC ACID

$\text{C}_{10}\text{HMePr(OH)ISO}_3\text{H}$ [14326] From thymol by successive sulphonation and iodation (Kehrmann, J pr [2] 39, 392) Gives an iodo thymoquinone on oxidation HNO_3 gives di nitro thymol [53°]— KA'2aq crystals melting in water of crystallisation at 80°, decomposed at 120°— BaA' — AgA'

IODO-THYMOQUINONE $\text{C}_{10}\text{HMePrIO}$ [52641] [61°] Formed by oxidising iodo-thymol sulphonic acid with CrO_3 (Kehrmann, J pr [2] 39, 392) Yellowish red prisms Reduces to an iodo hydrothymoquinone [74°]

Oxim $\text{C}_{10}\text{HMePrIO(NOH)}$ [52641] [130°] Formed by heating the quinone with hydroxylamine hydrochloride in diluted (75 p c) alcoholic solution Long yellow prisms and needles, sol alcohol and ether, insol cold water Its acetyl derivative $\text{C}_{10}\text{HMePrIO(NOAc)}$ crystallises in golden needles [70°] Its sodium derivative crystallises in greenish laminae

Iodo thymoquinone $\text{C}_{10}\text{HMePrIO}$ [52341] [66°] From iodo carvacrol sulphonic acid by oxidation with chromic acid mixture (Kehrmann, J pr [2] 40, 188) Garnet red tables, sol alcohol and ether More volatile with steam than its isomeride Smells like quinone Hydroxylamine slowly forms an oxim

o IODO-TOLUENE $\text{C}_6\text{H}_4\text{I}$: s $\text{C}_6\text{H}_4\text{ICH}_3$ [21] (205°) (B & K), (211° V) (K) S.G. 2.1697 (B & K) Formed from o toluidine by the diazo reaction (Beilstein & Kuhlberg, Z 8, 102, A 158, 347, Kekulé, B 7, 1007, Mabery & Robinson, Am 4, 101) Oil

Reactions—1 Oxidised by dilute nitric acid to o iodo benzoic acid [157°]—2 With ClCOEt and Na it forms $\text{C}_6\text{H}_4\text{MeCOEt}$ —3 CrO_3Cl gives $\text{C}_6\text{H}_4\text{ICHCl}$, and a little $\text{C}_6\text{H}_4\text{ICH}_3$ (Stuart & Elliott, C J 53, 805)—4 When heated with H_2SO_4 it forms iodo toluene sulphonic acid, di iodo toluene, and tri iodo toluene (Neumann, A, 241, 62)

m Iodo toluene $\text{C}_6\text{H}_4\text{ICH}_3$ [31] (204°) S.G. 2.1698 From *m* toluidine by the diazo reaction (B & K)

p Iodo-toluene $\text{C}_6\text{H}_4\text{ICH}_3$ [41] [35°] (2115°) From *p* toluidine by the diazo-reaction (Körner, Bull Acad Belg 1867, 157) The same body appears to be formed from mercury ditolyl and iodine, although the melting point is given as 20° (Dreher & Otto, A 154, 171) Laminae May be sublimed Gives *p* iodo benzoic acid when oxidised by chromic acid mixture H_2SO_4 forms iodo-toluene sulphonic acid and di- and tri-iodo-toluene (Neumann, A, 241, 58)

m-Iodo-toluene v BENZYL IODIDE.

o-IODO-TOLUENE SULPHONIC ACID

$\text{C}_6\text{H}_4\text{ISO}_3\text{H}$: s $\text{CH}_3\text{C}_6\text{H}_4\text{SO}_3\text{H}$ From o-iodo-toluene and SO_3 (Mabery & Palmer, Am 6, 170) Syrupy liquid— BaA' , 1½ aq : needles— CaA' , 2½ aq— PhA' , 2 aq

***p*-Iodo-toluene (a)-sulphonic acid**

$\text{CH}_3\text{C}_6\text{H}_4\text{I}\text{SO}_3\text{H}$ Formed, together with the (B) isomeric, by gradually adding *p* iodo toluene to SO_3 , both dissolved in chloroform (Glassner, B 8, 560) — BaA'_4 4aq needles, ν e sol water

***p* Iodo toluene (B)-sulphonic acid**

$\text{CH}_3\text{C}_6\text{H}_4\text{I}\text{SO}_3\text{H}$ Formed as above Deliquescent crystalline mass — KA'_4 4aq very soluble laminæ — NaA'_4 4aq dense aggregates of whetstone shaped very soluble crystals — CaA'_4 3aq very soluble silky needles — $\text{3aA}'_4$ 4q thin laminæ, sl sol water — CuA'_4 6aq light blue needles, ν sol water

Amide $\text{C}_6\text{H}_4\text{MeI}\text{SO}_2\text{NH}$ [179°] Crystals, m. sol hot water, ν so alcohol

Iodo-toluene disulphonic acid

$\text{C}_6\text{H}_3\text{MeI}(\text{SO}_3\text{H})_2$ [143°]

From $\text{C}_6\text{H}_5\text{Me}(\text{NH}_2)(\text{SO}_3\text{H})_2$ by diazo-reaction (E Richter, A 230, 325, Limpriicht, B 18, 2170) Slender white needles, ν sol alcohol and water — BaA''_4 6aq prisms, ν sol water — KA''_4 2aq small prisms

Chloride $\text{C}_6\text{H}_4\text{MeI}(\text{SOCl})_2$ [143°] After one fusion it melts at 126° Long white prisms, sl sol ethe-

Amide [130°-132°] White needles (from water)

***p*-IODO-*o*-TOLUIDINE** $\text{C}_6\text{H}_3\text{MeI}(\text{NH}_2)$ [142°]

[49°] (273°) From iodo *o* nitro toluidine by reduction (Heynemann, Z [2] 6, 402, A 158, 338) Needles Boils, with rapid decomposition, at 273° ν e sol alcohol, ether, and CS_2 — $\text{B}'\text{HNO}_3$ colourless nacreous laminæ S 95 at 16°

Iodo-toluidine $\text{C}_6\text{H}_4\text{MeI}(\text{NH})$ [143°] [189°] From *p* iodo toluene by nitration and reduction (Glassner, B 8, 561) Needles or plates, ν sol alcohol — $\text{B}'\text{HCl}$ needles — $\text{B}'\text{HNO}_3$ plates — $\text{B}'\text{H}_2\text{SO}_4$ needles

Di-iodo-*p*-toluidine $\text{C}_6\text{H}_3\text{MeI}_2(\text{NH}_2)$ [1354°]

[1245°] Formed by the action of ICl on *p* toluidine dissolved in HClAq (Michael a Norton, B 11, 115, Am 1, 263) Ramified groups of slender needles, m sol cold alcohol, sl sol hot water

IODO-TOLUIDINE SULPHONIC ACID

$\text{C}_6\text{H}_3\text{Me}(\text{NH})\text{I}(\text{SO}_3\text{H})$ [1245°] From the diazo derivative of $\text{C}_6\text{H}_3\text{Me}(\text{NO}_2)(\text{NH}_2)(\text{SO}_3\text{H})$ [1245°] and conc HI at 140° (Limpriicht a Foth A 230, 308, B 18, 2185) Satiny needles (containing aq) Sl sol cold water, m sol hot water — BaA''_4 trimetric tables, ν sol water

IODO-TOLUQUINONE $\text{C}_6\text{H}_3\text{MeI}_2$ [6241°]

[117°] Formed by oxidising a solution of iodo cresol sulphonic acid in sulphuric acid with chromic acid (Kehrmann, J pr [2] 37, 340, 39, 392) Long reddish yellow needles, m sol ether Reduced by stannous chloride to iodo hydrotoluquinone [111°] With hydroxylamine it gives the mono-oxim crystallising in short yellow prisms [156°]

Di-iodo-toluquinone $\text{C}_6\text{H}_3\text{MeI}_2\text{O}_2$ [36241°]

[118°] From di iodo *m* cresol sulphonic acid and CrO_3 (Kehrmann, J pr [2] 39, 392) Garnet-red laminæ, ν sol organic solvents May be sublimed

DI-IODO-DITOLYL [341°] $\text{C}_6\text{H}_3\text{MeI}\text{C}_6\text{H}_4\text{MeI}$ [134°] [100°]. From di amido ditolyl by Sandmeyer's modification of the diazo-reaction (Stolle, B. 21, 1096) Yellow needles

***o*-IODO-DI-TOLYL-ETHYLIDENE-DI-AMINE** $\text{CH}_3\text{CH}(\text{NH}_2)\text{CH}_2(\text{Me})_2$ Obtained by

heating iodo acetic aldehyde with *p*-toluidine (Chautard, A Ch [6] 16, 156) Orange yellow prisms or tables, decomposed by heat Does not furnish crystalline salts

IODO TOLYL METHYL KETONE

$\text{C}_6\text{H}_4\text{MeI}\text{COCH}_3$ [125°] [89°] Formed by heating diazo tolyl methyl ketone (from amido-tolyl methyl ketone [102°]) with an excess of HI (Klingel, B 18, 2700) Yellowish prisms ν sol alcohol and ether, sl sol ligroin and benzene, insol water By CrO_3 it is oxidised to iodo iso phthalic acid [204°]

 β IODO-VALERIC ACID $\text{C}_6\text{H}_4\text{IO}_2$ 12

$(\text{CH}_3)_2\text{CI}\text{CH}_2\text{CO}_2\text{H}$ [80°] Solidifies at 59° Separates as crystals when HI is passed into a concentrated solution of β oxy isovaleric acid (Schirokoff, J pr [2] 23, 285) Converted by sodium amalgam to isovaleric acid

Iodo valeric acid $\text{C}_6\text{H}_4\text{MeI}\text{CO}_2\text{H}$ (?) **Hydro-iodide of tiglic acid** [865°] Formed by the combination of tiglic (methyl crotonic) acid with HI which may be effected in the cold (Schmidt a Berendes, A 191, 117) Formed also, together with the following, when angelic acid is heated with HIAq (Schmidt, A 208, 254) Needles, sl sol cold water, decomposed by boiling with water Converted by zinc and dilute H_2SO_4 into $\text{C}_6\text{H}_4\text{MeH}\text{CO}_2\text{H}$ Aqueous AgNO_3 gives AgI , tiglic acid and CO_2

Iodo valeric acid $\text{CH}_3\text{CHI}\text{CHMe}\text{CO}_2\text{H}$ [46°] **Hydro iodide of angelic acid** From angelic acid and conc HIAq in the cold (Fittig, A 216, 162) Prisms, ν sol water Zinc and dilute H_2SO_4 converts it into $\text{C}_6\text{H}_4\text{MeH}\text{CO}_2\text{H}$ Aqueous AgNO_3 gives AgI , tiglic acid, and CO_2 Aqueous Na_2CO_3 at 0° gives CO_2 , ψ butylene $\text{CH}_3\text{CHCHCH}_3$, and HI ν also Angelic acid, vol 1 p 266

IODO-ISOVALERIC-ALDEHYDE $\text{C}_6\text{H}_4\text{IO}_2$ 12

$(\text{CH}_3)_2\text{CH}\text{CHI}\text{CHO}$ (?) SG 1217

Preparation — Isovaleric aldehyde (24 cc) dissolved in alcohol (50 cc) is treated with iodine (20 g), and iodic acid (8 g) The reaction takes place at the ordinary temperature, and is complete in about 15 days The liquid is poured into excess of water, and decolourised by alkali or reduced silver (Chautard A Ch [6] 16, 160).

Properties — Colourless liquid, blackening on exposure to light, and having an extremely irritating and suffocating odour It is completely decomposed at 100°, and is not solidified at -70° ν sol alcohol and ether, less sol benzene, CHCl_3 , and CS Sl sol water

Reactions — Rapidly decomposed by alkalis. The action of ammonia is complex, giving valeric aldehyde ammonia, valeridine $\text{C}_6\text{H}_4\text{N}$, and valentrine $\text{C}_6\text{H}_4\text{N}$ Forms a crystalline compound with NaHSO_4 $\text{Ag}(\text{C}_6\text{H}_4\text{O}_2)$ at 100° yields amyl acetate (138°) AgCN and AgSCN react forming AgI and the corresponding derivatives. Forms a mono- and a di anilide with aniline.

IODO-VANILLIN *o* Methyl derivative of**IODO-DI OXY-BENZEOIC ALDEHYDE****DI-IODO-VINYL-AMINE** $\text{Cl}_2\text{CH}(\text{NH}_2)$ [192°

with decomposition] Formed by the action of cold aqueous NH_3 , CO_2 and alcohol being eliminated upon the di iodo oxy acrylic ether $\text{Cl}_2\text{C}(\text{OH})\text{CO}_2\text{Et}$, obtained by treating diazo-oxy acrylic ether $\text{CN}_2\text{C}(\text{OH})\text{CO}_2\text{Et}$ with an ethereal solution of iodine (Buchner a Curtius, B. 19, 851) Small yellowish prisms. Very

sparingly soluble in cold water and ether, more easily in hot water and hot alcohol. Volatile with steam. It is stable towards acids, but alkalis set free NH_3 in the cold.

Di- α -iodo- α -xylene $\text{C}_8\text{H}_6(\text{CH}_2\text{I})_2$, *o*-Xylyl *ene-iodide* [110°]. Prismatic crystals. Formed by heating phthalyl alcohol (di- α -oxy-xylene) with HI (Lesser, B 17, 1826).

Di- α -iodo- p -xylene $\text{C}_8\text{H}_6(\text{CH}_2\text{I})_2$, [α 170°]. Obtained by boiling [41] $\text{C}_8\text{H}_6(\text{CH}_2\text{OH})_2$ with conc HIAq for a few minutes (Grimaux, Z 1870, 835). Slender needles, sl sol ether, v sol boiling alcohol and chloroform. Turns yellow in air. Not very volatile with steam.

iodoplumbic acid H_2PbI_4 , v **LEAD**, **iodide of, Combinations**, No 3.

IONS. The elements or radicals into which a compound is primarily separated by electrolysis (v **PHYSICAL METHODS**, section *Electrical methods*).

IPECACUANIC ACID $\text{C}_8\text{H}_7\text{O}_5$. An acid existing, according to Willkg (A 76, 342), along with emetine (*q v*) in the root of *Cephaelis Ipecacuanha*. It is extracted by boiling alcohol, ppd. by basic lead acetate, and the pp decomposed by H_2S . Reddish brown, very bitter, amorphous mass, m sol ether, v sol alcohol and water. Colours ferric salts green, the colour being changed to violet by ammonia. Its dilute solution does not ppt $\text{Pb}(\text{OAc})_2$, but ppts basic lead acetate. Its alkaline solution absorbs oxygen from the air. It thus somewhat resembles gallic acid. Podwysotzky (*Ph* [3] 10, 642) by extracting ipecacuanha with ether and light petroleum obtained an acid ('Erythrocephalein'), which formed a purple red alkaline salt, and which crystallised from chloroform in coloured needles.

IPEOMIC ACID $\text{C}_8\text{H}_7\text{O}_5$, [104°]. Produced by the action of nitric acid on jalapin (Mayer, A 85, 143, Poleck a Samelson, C C 1884, 813). Resembles sebacic acid.

IRIDIUM Ir. At w 192.5 Mol w un known [2200°] (Van der Weyde), [1950°] (Viole, C R 89, 702), [2500°] (Pictet, C R 88, 1317) SG 22.42 at 17.5° (Deville a Debray, P M [4] 50, 651) SH 0°-100° = 0.323, 0°-1400° = 0.401 (Viole, C R 89, 702) CE linear 000007 (Fizeau, C R 68, 1125).

Occurrence—As metal, alloyed with Pt, Os, Ru, Rh, and Pd. Specimens of platinum-iridium, osmium-iridium, &c, contain from 25 to 78 p.c. Ir (v Deville a Debray, A Ch [3] 56, 431, Berzelius, P 13, 435, 527, 15, 208). In 1802 Tennant (*T* 1804 411) noticed a metallic residue when he dissolved Pt ore in *aqua regia*, this residue was examined by Descotils (Gehlen's *Journ Chem* 2, 273), and Fourcroy a Vauquelin (Gehlen's *Journ Chem* 3, 362). In 1804 Tennant (*lc*) showed that the residue contained two new metals, to one he gave the name *Iridium* because of the varied colours of its oxides, and to the other the name *Osmium* because of the peculiar smell of its volatile oxide.

Formation—1 By digesting Ir_2O_3 or IrCl_3 with formic acid—2 By action of sunlight on an alcoholic solution of Ir sulphate.

Preparation—When Pt ore is heated with *aqua regia* osmium and platinum-iridium remain as black insoluble scales. Ir is prepared from the residue by various methods. 1 W von

Schneider (*A. Suppl* 5, 261) mixes the insoluble in *aqua regia* with N_2Cl , and heats to low redness in a stream of Cl_2 , Os is thus removed. The residual mixed chlorides are triturated with the smallest possible quantity of water, and then washed on to a filter, the solution is treated with Cl_2 mixed with a little HClAq , and shaken up with excess of KCl , the pp is washed with water, and then with saturated KClAq . The pp is now nearly pure K_2MCl_6 , $\text{M} = \text{Ir, Ru, and Pt}$. The pp is dissolved in much boiling water, and H is passed in for several days (the flask being closed by a caoutchouc cork) until the supernatant olive-green liquid treated with potash becomes colourless and then blue or turbid. Pt and Ru are thus wholly ppd as tartrates, and most of the Ir remains in solution as IrCl_3 . H is removed by a stream of CO_2 —otherwise explosions occur on opening the flask from the action of the ppd Pt and Ru on the H and O in the flask—the corks are removed, and the liquid is again saturated with H, Ir is ppd on the surface of the liquid in large lustrous laminae—2 Matthey (*Pr* 28, 463) prepares Ir free from all metals except Pt as follows. Ordinary Ir, finely divided, is fused with 10 times its weight of Pb, the Pb is dissolved in HNO_3Aq , and the insoluble is digested for a long time with *aqua regia*, the insoluble is fused with KHSO_4 (to remove Rh), and then melted with 10 times its weight of KOH and 3 times its weight of KNO_3 in a gold crucible, the cold mass is treated with cold water, in which K iridate remains insoluble. The insoluble is well washed with water containing a little KOH and NaClO , and then with water, fairly conc cold NaClOAq is added to the residual blue solid, after a time the water is distilled off, the residue is again fused with KNO_3 and KOH, and treatment with dilute NaClOAq and KOHAq is repeated. The blue solid is now dissolved in *aqua regia*, the liquid is evaporated to dryness, and the residue is redissolved and filtered, the dark-coloured liquid is slowly poured into conc NaOHAq containing NaClO , and Cl_2 is passed in, the liquid being kept in a distilling apparatus, the blue oxide of Ir thus obtained is collected, washed, dried, and reduced in a mixture of CO and CO_2 (made by gently warming oxalic acid with sulphuric acid). Metallic Ir is thus obtained, while any Fe present remains as oxide. The mass is heated to redness with KHSO_4 , and repeatedly washed with ClAq to remove Au, and with HFAq to remove SiO_2 , it is finally washed with water and dried. Pt may be removed by dissolving in *aqua regia*, saturating with NH_3Cl , dissolving the double NH_3 , Ir and NH_3 , Pt chlorides in hot water, reducing by SO_2 , when $\text{IrCl}_3 \cdot 3\text{NH}_3\text{Cl}$ is formed, and dissolves with olive green colour, while $\text{PtCl}_2 \cdot 2\text{NH}_3\text{Cl}$ remains insoluble, the reduction should be stopped before the whole of the $\text{IrCl}_3 \cdot 2\text{NH}_3\text{Cl}$ is reduced (the presence of a little of this salt gives a deep red colour to the mixed chlorides). The solution is oxidised by HNO_3 , NH_3Cl is added, and the ppd $\text{IrCl}_3 \cdot 2\text{NH}_3\text{Cl}$ is washed with NH_3ClAq , and strongly heated, when Ir remains as a grey metallic powder (Claus, *J pr* 42, 251). Wöhler a Mucklé (*A* 104, 368) reduce the mixed Pt NH_3 and Ir NH_3 chlorides by digesting with slightly warm KCNaq till the undissolved is light yellow brown, taking care to avoid excess, $\text{IrCl}_3 \cdot 3\text{NH}_3\text{Cl}$ is formed and

dissolves, while $\text{PtCl}_2 \cdot 2\text{NH}_4\text{Cl}$ remains unreduced and insoluble. For other methods of preparing Ir v Deville a Debray (*A Ch* [8] 56, 481), Claus (*C C* 1862 129), Wöhler (*P* 81, 161, 104, 363; 107, 857), Fremy (*A Ch* [8] 44, 885), Martius (*A* 107, 360), Bunsen (*A* 146, 274).

Properties -- As prepared by heating $\text{IrCl}_3 \cdot 2\text{NH}_4\text{Cl}$, Ir forms a grey spongy mass, Ir as a grey powder is produced by heating $\text{IrCl}_3 \cdot 2\text{KCl}$ with excess of Na_2CO_3 , washing and reducing in H, by heating to whiteness and compressing, a compact lustrous mass is obtained. When fused in a very powerful O H flame, Ir forms a white, lustrous mass resembling polished steel, brittle when cold, somewhat malleable at red heat (D a D). Harder than iron. SG of porous Ir varies from 16 to 19. Ir which has been very strongly heated is insoluble in all acids, Ir black is sol in *aqua regia*. Heated in Cl, Ir black forms IrCl_3 .

The atomic wt of Ir has been determined by heating $\text{IrCl}_3 \cdot 2\text{HCl}$ in H (Berzelius, *P* 13, 435, 527, 15, 208), by reducing $\text{IrCl}_3 \cdot 2\text{NH}_4\text{Cl}$ in H and weighing the residual Ir, and by reducing the double Ir-K chloride in H and dissolving out the KCl (Seubert, *B* 11, 1767). As no compound of Ir has been gasified, the valency of the atom of Ir in gaseous molecules is not determined. From the crystalline form of osmium G Rose (*P* 77, 143) concluded that Os and Ir are isomorphous, and that both belong to the hexagonal system.

In its chemical reactions, Ir is closely related to Os and Pt, and less closely but very distinctly related to Ru, Rh, and Pd, all these elements are metallic, but the instability of their salts, the solubility of the hydrated oxides MO_2 in alkalis, and the formation of acids H_2MCl_4 and H_2MBr_4 , show the non-metallic tendencies of the Pt metals (v. NOBLE METALS).

Ir is used alloyed with Pt for making instruments, &c., which remain unchanged in air, & q for making the normal metre preserved as the standard of reference. Stylographic pens are sometimes tipped with Ir. When Ir powder is heated to whiteness with $\frac{1}{4}$ its weight of P the whole melts, on cooling, the mass may be readily worked, by strongly heating with lime, the P is entirely removed.

Reactions and Combinations -- 1 Heated in oxygen, compact Ir is not oxidised, Ir black is oxidised to Ir_2O_3 , which decomposes again at c 1000° (Deville a Debray, *C R* 1878 441, cf. Dissociation, vol II p 397). -- 2 Ir black is oxidised by fusion with *nitre*, *potash*, *potassium carbonate*, or *potassium hydrogen sulphate*. -- 3 Heated in *chlorine* IrCl_3 is formed, mixed with NaCl or KCl, and heated in Cl, soluble $\text{IrCl}_3 \cdot 2\text{Na}(\text{K})\text{Cl}$ is formed. -- 4 Heated in an *alcohol flame*, Ir becomes covered with a blackish layer which disappears on heating in air, leaving a portion of the Ir combined with C (t. IRIDIUM CARBID). -- 5 Ir probably combines with *phosphorus* when the elements are heated together, when the product is heated in air, Ir and a phosphate of Ir are formed.

Iridium alloys of Faraday a Stodart (*A Ch* 21, 73), Deville a Debray (*C R* 81, 839), Fixeau (*C R* 78, 1205), Morin (*C R* 78, 1502), Wöhler (*A* 146, 875).

Osmium-iridium alloy Occurs in South

America, California, Australia, Ural Mountains, &c. Hexagonal prisms, lustrous, steel grey. Slightly malleable, SG 19.3 to 21.1. Analyses show composition approximating to formulae Ir_2Os , IrOs and IrOs_2 , with Ir more or less replaced by Pt, Rh, and Ru, but it is probably non-homogeneous (D a D). Heated in air, OsCl_4 is given off, heated with KNO_3 and KOH K osmate and iridate are formed. Scarcely acted on by *aqua regia*. Alloys with Fe (F).

Platinum-iridium alloy Occurs native in octahedra. 1 pt Ir heated in O H flame with 10 pts Pt gives a ductile alloy, Pt alloyed with 15-20 p.c. Ir is insoluble in *aqua regia*. For accounts of the Pt-Ir alloy used for making the normal standard metre v. D a D and also M. Alloys of Ir with Pt and Rh, and with Pt, Rh, and Sn were prepared by D a D. Alloys of Ir with Cu, Au, Pb, Hg, Ru, and Sn have been prepared.

Iridium, ammonio salts of, or Irid ammonium salts (Ammoniacal iridium bases) IrCl_3 dissolves in $(\text{NH}_4)_2\text{CO}_3$ aq and on neutralising with HCl aq, the compound $\text{IrCl}_3 \cdot 2\text{NH}_3$ is produced, treated with H_2SO_4 aq, this compound yields $\text{IrSO}_4 \cdot 2\text{NH}_3$. $\text{IrCl}_3 \cdot 2\text{NH}_3$ dissolves in boiling NH_4Aq , and the compound $\text{IrCl}_3 \cdot 4\text{NH}_3$ forms on cooling, this compound gives a sulphate and nitrate, SO_4 and 2NO_3 replacing Cl_2 . From $\text{IrCl}_3 \cdot 3\text{NH}_3$ Cl is obtained $\text{IrCl}_3 \cdot 5\text{NH}_3$, and this again yields a basic hydroxide $\text{Ir}(\text{OH})_3 \cdot 5\text{NH}_3$, a carbonate $\text{Ir}_2(\text{CO}_3)_3 \cdot 10\text{NH}_3$, and other salts. Finally by treating $\text{IrCl}_3 \cdot 2\text{NH}_3$ with conc. HNO_3 aq the compound $\text{IrCl}_3 \cdot (\text{NO}_3)_2 \cdot 4\text{NH}_3$ is obtained, AgNO_3 aq does not ppt Cl from this salt, treated with HCl aq the salt $\text{IrCl}_3 \cdot \text{Cl}_2 \cdot 4\text{NH}_3$ is produced from which AgNO_3 aq ppts only half the Cl. The foregoing compounds may be formulated as double salts of IrCl_3 , IrCl_4 , IrCl_5 , &c., with NH_3 , but many of their reactions are better explained if we regard them as salts of condensed ammoniums containing Ir. The following classification is that usually adopted --

1 IRIDOSAMMONIUM COMPOUNDS $\text{N}_2\text{H}_5\text{IrX}_2$, and $\text{N}_2\text{H}_5\text{IrX}_2$, or $\text{NH}(\text{NH}_4)\text{IrX}_2$, and $\text{N}_2\text{H}_5(\text{NH}_4)_2\text{IrX}_3$, &c., obtained from IrCl_3 .

Iridosammonium chloride $\text{N}_2\text{H}_5\text{IrCl}$ (simplest formula $\text{IrCl}_3 \cdot 2\text{NH}_3 = \text{di-ammonio-iridium dichloride}$) A yellow, curdy pp, obtained by dissolving IrCl_3 in excess of $(\text{NH}_4)_2\text{CO}_3$ aq and neutralising the greenish yellow liquid with HCl aq. Insol hot and cold water. Heated gives NH_4Cl , HCl and Ir (Skoblikoff, *A* 84, 275). Heated with H_2SO_4 aq, easily soluble orange coloured **iridosammonium sulphate**, $\text{N}_2\text{H}_5\text{IrSO}_4$, is produced.

Iridoso diammonium chloride $\text{N}_2\text{H}_5\text{Ir}_2\text{Cl}_3$ or $\text{N}_2\text{H}_5(\text{NH}_4)_2\text{Ir}_2\text{Cl}_3$ (simplest formula $\text{IrCl}_3 \cdot 4\text{NH}_3 = \text{tetrammonio-iridium dichloride}$) $\text{IrCl}_3 \cdot 2\text{NH}_3$ is boiled with NH_4Aq until almost all dissolved, the solution is filtered and allowed to cool. Nearly white pp, insol in cold water, heated, or placed in boiling water, goes to $\text{IrCl}_3 \cdot 2\text{NH}_3$ (Skoblikoff, *l.c.*) The **sulphate**, $\text{N}_2\text{H}_5\text{Ir}_2\text{SO}_4$, is obtained by the action of H_2SO_4 aq on the chloride, and the nitrate, $\text{N}_2\text{H}_5\text{Ir}_2(\text{NO}_3)_4$, by addition of Ba_2NO_3 aq to a solution of the sulphate in warm water.

2 IRIDAMMONIUM COMPOUNDS $\text{N}_2\text{H}_5\text{Ir}_2\text{X}_4$, or $\text{N}_2\text{H}_5(\text{NH}_4)_2\text{Ir}_2\text{X}_4$, obtained from IrCl_3 .

Irido pentammonium chloride $\text{N}_2\text{H}_5\text{Ir}_3\text{Cl}_4$ (simplest formula $\text{IrCl}_3 \cdot 10\text{NH}_3 = \text{decammonio-}$

iridium trichloride $\text{IrCl}_3 \cdot 3\text{NH}_4\text{Cl}$ is dissolved in water, an equal volume of conc. NH_4Aq is added, a flask is quite filled with the liquid, and allowed to stand for several weeks in a warm place until the olive green colour of the liquid has changed to deep rose red, after gentle warming, to remove excess of NH_3 , the liquid is saturated with HClAq and evaporated to dryness, the greenish yellow salt which separates during evaporation is washed with cold water, and then crystallised from water acidified with HCl . Flesh-coloured, crystalline, powder, v. sol water (Claus, *J pr* 63, 99, A 98, 317).

Irido pentammonium hydroxide

$\text{N}_5\text{H}_4\text{Ir}_2(\text{OH})_2$. Known only in solution which is obtained by digesting the chloride with moist Ag_2O . Solution is alkaline, neutralised by acids gives carbonate, nitrate, and sulphate.

3 IRIDI AMMONIUM COMPOUNDS $\text{N}_5\text{H}_4\text{IrCl}_2 \text{X}$, or $\text{N}_5\text{H}_4(\text{NH}_4)_2\text{Cl}_2\text{IrX}$, obtained from series 1 by oxidation.

Iridi-diammonium chloride? better *chloriridi-diammonium chloride* $\text{N}_5\text{H}_4\text{IrCl}_2\text{Cl}_2$ (simplest formula $\text{IrCl}_4 \cdot 4\text{NH}_3 = \text{tetrammonioiridium tetrachloride}$). Obtained by slowly warming iridosammonium chloride ($\text{N}_5\text{H}_4\text{IrCl}_2$) with conc. HNO_3Aq , dissolving the salt which separates in hot water, adding excess of HClAq , and crystallising from hot water. Violet octahedra, AgNO_3Aq ppts only $\frac{1}{2}$ the Cl from this salt. The salt which separates after treating $\text{N}_5\text{H}_4\text{IrCl}_2$ with HNO_3 is the *nitrate*, $\text{N}_5\text{H}_4\text{IrCl}_2(\text{NO}_3)_2$, AgNO_3Aq does not ppt Cl from this salt. When this salt is evaporated with dilute $\text{H}_2\text{SO}_4\text{Aq}$, the *sulphate*, $\text{N}_5\text{H}_4\text{IrCl}_2\text{SO}_4$, is obtained (Skoblikoff, *lc*). Palmaer (*B* 22, 15) describes $\text{N}_5\text{H}_4\text{IrCl}_2\text{Cl}_2$, $\text{N}_5\text{H}_4\text{IrCl}_2\text{SO}_4 \cdot 2\text{H}_2\text{O}$, $\text{N}_5\text{H}_4\text{IrCl}_2(\text{NO}_3)_2$, $\text{N}_5\text{H}_4\text{IrCl}_2\text{Cl}$, and $(\text{N}_5\text{H}_4\text{IrCl}_2)_2\text{SO}_4 \cdot 2\text{H}_2\text{O}$.

References — For other memoirs besides those referred to, v. Weltzien, A 97, 19, Gibbs a Genth, *J* 1858 214, Birnbaum, *B* 12, 1544.

Iridium boride of. When Ir is heated with B to a very high temperature a fusible mass is obtained, no definite boride has been isolated (Wöhler a Deville, *A Ch* [3] 52, 71).

Iridium bromides of (Birnbaum, A 133, 161). Only one bromide has been isolated, $\text{IrBr} \cdot 4\text{H}_2\text{O}$, there are indications of the existence in solution of a tetrabromide, IrBr_4 . Ir does not combine directly with Br, nor is it soluble in a mixture of HNO_3 and HBrAq .

IRIDIUM TRIBROMIDE $\text{IrBr}_3 \cdot 4\text{H}_2\text{O}$ (*Iridobromide* *iridium sesquibromide*). When the blue hydrate IrO_2H_2 (obtained by heating a solution of IrCl_3 with alkali) is dissolved in HBrAq and evaporated *in vacuo*, olive green six-sided crystals separate, and then steel blue needles. The olive-green crystals are $\text{IrBr}_3 \cdot 4\text{H}_2\text{O}$, and the steel-blue crystals are $\text{IrBr}_3 \cdot 3\text{HBr} \cdot 3\text{H}_2\text{O}$. The bromide loses $3\text{H}_2\text{O}$ at 100° – 120° , it is sol water, insol alcohol or ether, aqueous solution is green, but becomes blue probably with formation of IrBr .

Irido-bromhydric acid

$\text{IrBr}_3 \cdot 3\text{HBr} \cdot 3\text{H}_2\text{O} (= \text{H}_3\text{IrBr}_3 \cdot 3\text{H}_2\text{O})$. Crystallises in steel-blue crystals which melt at 100° , giving off $8\text{H}_2\text{O}$, deliquescent, easily sol water, alcohol, ether, oxidised by HNO_3 , probably to IrBr_4 .

Ammonium-iridium tribromide, or *Ammonium iridobromide* or *bromiride*

$2(\text{IrBr}_3 \cdot 3\text{NH}_4\text{Br}) \cdot 3\text{H}_2\text{O}$, is obtained by reducing $\text{IrBr}_3 \cdot 2\text{NH}_4\text{Br}$ (formed by adding $\text{IrCl}_3 \cdot 2\text{NH}_4\text{Cl}$ to hot NaBrAq and cooling) by SO_2 and neutralising by $(\text{NH}_4)_2\text{CO}_3$. The following other double salts are described by Birnbaum (A 133, 161): $\text{IrBr}_3 \cdot 3\text{H}_2\text{GBr}$, $\text{IrBr}_3 \cdot 3\text{AgBr}$, $\text{IrBr}_3 \cdot 3\text{KBr} \cdot 3\text{H}_2\text{O}$, $\text{IrBr}_3 \cdot 3\text{NaBr} \cdot 12\text{H}_2\text{O}$.

IRIDIUM TETRABROMIDE (*Iridibromide*). When IrO_2H_2 is dissolved in HBrAq , or when $\text{IrBr}_3 \cdot 2\text{KBr}$ is decomposed by $\text{H}_2\text{SiF}_6\text{Aq}$, a solution is obtained which loses Br on evaporation, and on addition of HNO_3Aq , after evaporation, gives a deliquescent, blue, crystalline mass, easily sol in water and alcohol. The blue solution probably contains $\text{IrBr}_3 \cdot 2\text{HBr}$ (Birnbaum). This solution does not yield double compounds with other metallic bromides.

Ammonium iridium tetrabromide, or *Ammonium iridibromide* or *bromiride* $\text{IrBr}_3 \cdot 3\text{NH}_4\text{Br}$ ($(\text{NH}_4)_3\text{IrBr}_4$). Dark blue octahedra, by adding $\text{IrCl}_3 \cdot 2\text{NH}_4\text{Cl}$ to hot NaBrAq , and cooling.

Potassium bromiride $\text{IrBr}_3 \cdot 2\text{KBr}$ (K_2IrBr_5). When solution of IrCl_3 , or $\text{IrCl}_3 \cdot 2\text{NaCl}$, or $\text{IrCl}_3 \cdot 2\text{NH}_4\text{Cl}$ is warmed with conc. KBrAq , a greenish blue liquid is obtained from which very dark blue crystals of K_2IrBr_5 separate, the salt is recrystallised from hot water.

Sodium bromiride also exists

Iridium carbide of (? IrC). When a coherent mass of Ir is held in the flame of a spirit lamp, black masses appear on the surface, these burn when heated in air and leave 80.2 pct Ir (Berzelius, *P* 15, 213).

Iridium chlorides of. Three chlorides are known IrCl_3 , IrCl_4 , and IrCl_5 , as none has been gashed the formulae are not necessarily molecular (v. *Iridium, haloid compounds* of). When Ir black is heated in Cl_2 , IrCl_3 is formed, when Cl_2 is passed over a heated mixture of Ir black and KCl , $\text{IrCl}_3 \cdot 2\text{KCl}$ is produced.

IRIDIUM DICHLORIDE (*Iridosochloride*) IrCl_2 . A blackish green, insoluble, solid. By passing Cl over Ir black heated to low redness (Berzelius, *P* 13, 470). The change is not complete, as the temperatures of formation and decomposition are nearly the same (Claus, A 59, 219). By heating Ir sulphite in Cl_2 (Fellenberg, *P* 50, 66). By carefully heating IrCl_3 (Skoblikoff, A 84, 275). Seubert (*B* 11, 1761) describes an acid $\text{IrCl}_3 \cdot \text{H}_2\text{SO}_4 \cdot 4\text{NH}_4\text{Cl}$ from which salts of NH_4 and Na are obtained.

IRIDIUM TRICHLORIDE (*Iridochloride* *iridium sesquichloride*) IrCl_3 . Olive green solid, insol in acids or alkalis (Claus, A 107, 129). Obtained by heating Ir black in Cl_2 , also by strongly heating an alkali double salt of IrCl_3 with conc. H_2SO_4 , and pouring into cold water (Berzelius, *P* 13, 470). A soluble hydrate $\text{IrCl}_3 \cdot 4\text{H}_2\text{O}$ is obtained by dissolving IrO_2H_2 in HClAq , reducing by H_2S , and evaporating.

Double salts *Ammonium iridium trichloride* (*Ammonium chloriride*) $2(\text{IrCl}_3 \cdot 3\text{NH}_4\text{Cl}) \cdot 3\text{H}_2\text{O}$, formed by slowly evaporating mixed solutions of $\text{IrCl}_3 \cdot 3\text{NaCl}$ and NH_4Cl , or by reducing $\text{IrCl}_3 \cdot 2\text{NH}_4\text{ClAq}$ by H_2S , adding conc. NH_4ClAq , filtering if necessary, and slowly evaporating (Claus, *J pr* 42, 851; Seubert, *B* 11, 1761), olive green, rhombic plates.

Potassium chloriridate $\text{IrCl}_3 \cdot 3\text{H}_2\text{O}$ By reducing $\text{IrCl}_3 \cdot 2\text{KCl}$ in H_2S , and evaporating with addition of KCl . Olive-green efflorescent prisms. The following double salts are also known — $2\text{IrCl}_3 \cdot (12\text{NH}_3, \text{Co})\text{Cl}_6$, $\text{IrCl}_3 \cdot 3\text{H}_2\text{O}$, $\text{IrCl}_3 \cdot 3\text{AgCl}$, $\text{IrCl}_3 \cdot 3\text{NaCl} \cdot 12\text{H}_2\text{O}$ (Claus, *l. c.*, Karmrodts a Uhrlaub, *A* 81, 120)

IRIDIUM TETRACHLORIDE (Iridochloride) IrCl_4 . The solution obtained by decomposing $\text{IrCl}_3 \cdot 2\text{NH}_4\text{Cl}$ suspended in water by Cl contains IrCl_4 (Vauquelin, *A Ch* 89, 150, 225), a solution of IrCl_4 is also obtained by dissolving $\text{IrO}_2 \cdot \text{H}_2\text{O}$ in HCl aq, digesting $\text{IrCl}_3 \cdot \text{aq}$ with *aqua regia*, or decomposing $\text{IrCl}_3 \cdot 2\text{KCl}$ with $\text{H}_2\text{SiF}_6 \cdot \text{aq}$ (Berzelius, *P* 13, 470). By evaporating these solutions to dryness at temperatures not above 40° , IrCl_4 is obtained as a black mass, translucent with dark-red colour at the edges, very deliquescent, decomposed by heat to IrCl_3 and Cl_2 , and then Ir combines with alkali chlorides to form double salts **Ammonium iridium tetrachloride (Ammonium chloriridate)** $\text{IrCl}_4 \cdot 2\text{NH}_4\text{Cl}$, obtained by mixing conc solutions of NH_4Cl and IrCl_3 , or $\text{IrCl}_3 \cdot 2\text{NaCl}$. Crystallises from hot water in dark red brown octahedra, isomorphous with $\text{PtCl}_4 \cdot 2\text{NH}_4\text{Cl}$. Soluble in cold water. Reduced by SO_2 to the more soluble $\text{IrCl}_3 \cdot 3\text{NH}_4\text{Cl}$.

Potassium chloriridate $\text{IrCl}_3 \cdot 2\text{KCl}$ Formed by mixing conc solutions of its constituent salts, also by dissolving $\text{IrO}_2 \cdot \text{H}_2\text{O}$ in HCl aq (solution probably contains H_2IrCl_6) and adding KCl , also by passing Cl_2 at a gentle heat over an intimate mixture of Ir black and KCl , dissolving in hot water, filtering from Ir , evaporating to dryness with addition of *aqua regia*, dissolving out KCl by small quantities of cold water, dissolving in hot water, adding a little *aqua regia*, and evaporating to the crystallising point. Black octahedra, very soluble in cold water, much more soluble in hot water, insoluble in alcohol. Heated, goes to $\text{IrCl}_3 \cdot 3\text{KCl}$ and at higher temperature to Ir and KCl .

The following **chloriridates** have also been obtained — $\text{IrCl}_3 \cdot 2(\text{NH}_4, \text{CH}_3)\text{Cl}$ (Vincent, *C R* 100, 112), $\text{IrCl}_3 \cdot 2\text{NaCl} \cdot 6\text{H}_2\text{O}$ (Vauquelin, *l. c.*, Berzelius, *l. c.*)

Iridium cyanides of, also **Iridicyanides**, *v* vol II p 332

Iridium haloid compounds of The only haloid compound certainly formed by direct union of the elements is IrCl_3 . When an intimate mixture of Ir black and KCl is heated in Cl_2 , or of Ir black and KI is heated in I_2 , the double salt K_2IrCl_6 or K_2IrI_6 is obtained. The following formulæ present the composition of the haloid compounds and the double salts which they form with alkali haloid compounds —

double alkali-iridium haloid salts are probably better regarded as salts of the acids H_2IrX_6 , H_2IrX_4 , and H_2IrX_2 . The chloriridates (salts of H_2IrCl_6) are readily reduced to chloriridites (salts of H_2IrCl_4).

Iridium hydroxides of, *v* **Iridium oxides** and **hydrated oxides** of

Iridium iodides of (Oppler, *Ueber Iodverbindungen des Iridiums* (Göttingen, 1857), *J* 1857 263) Two iodides, IrI_3 and IrI_2 , have been isolated. There are indications of the existence of IrI_2 . Ir and I do not directly combine.

IRIDIUM DI IODIDE (Iridoso iodide) The brown solid obtained by leading SO_2 into IrI_3 suspended in water is perhaps IrI_2 . When a solution of $\text{IrCl}_3 \cdot 2\text{NH}_4\text{Cl}$ in boiling KIAq is allowed to cool, a black crystalline powder separates, this is **ammonium iridium di iodide**, $\text{IrI}_2 \cdot 2\text{NH}_4\text{I}$.

IRIDIUM TRI IODIDE (Irido iodide Iridium sesqui iodide) IrI_3 . A black crystalline pp., by adding NH_4Cl to $\text{IrI}_3 \cdot 2\text{KIAq}$. Very slightly soluble in cold water, more soluble in hot water.

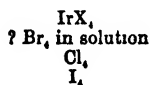
Double salts — Ammonium iridium tri iodide (Ammonium iodiridate) $2(\text{IrI}_3 \cdot 3\text{NH}_4\text{I}) \cdot \text{H}_2\text{O}$. Crystalline needles, by dissolving $\text{IrCl}_3 \cdot 2\text{NH}_4\text{Cl}$ in boiling KIAq , cooling, filtering from $\text{IrI}_2 \cdot 2\text{NH}_4\text{I}$, concentrating, and recrystallising from hot water.

Potassium iodiridate, $\text{IrI}_3 \cdot 3\text{KI}$. Green lustrous crystalline powder, by reducing IrCl_3 by H_2S with addition of KIAq . **Silver iodiridate**, $\text{AgI} \cdot 3\text{KI}$, has also been obtained.

IRIDIUM TETRA IODIDE (Iridi iodide) IrCl_4 . A black powder, by boiling $\text{IrCl}_3 \cdot \text{aq}$ with KI in presence of a little HCl .

Double salts — Ammonium iridium tetra iodide (Ammonium iodiridate) $\text{IrI}_4 \cdot 2\text{NH}_4\text{I}$. Separates after some weeks from solution of $\text{IrCl}_3 \cdot 2\text{NH}_4\text{Cl}$ in cold conc KIAq , dark brown lustrous crystals, easily decomposed by heat, aqueous solution gently heated deposits IrI_3 and $\text{IrI}_2 \cdot 2\text{NH}_4\text{I}$. **Potassium iodiridate**, K_2IrI_6 . Separates, after IrI_3 , from solution of IrCl_3 in KIAq . Also formed by dissolving IrI_3 in KIAq , and allowing to crystallise, also, in small quantity, by action of I vapour on an intimate mixture of Ir black and KI at 60° – 70° . **Sodium iodiridate**, $\text{IrI}_4 \cdot 2\text{NaI}$, has also been obtained.

Iridium oxides and hydrated oxides of The only oxides of Ir which have been certainly isolated are Ir_2O_3 and IrO . The former forms a hydrate with $3\text{H}_2\text{O}$, and another hydrate probably with $5\text{H}_2\text{O}$. IrO forms the hydrate $\text{IrO}_2 \cdot 2\text{H}_2\text{O}$. Hydrated Ir_2O_3 dissolves in alkalis probably with formation of **iridates**, $\text{IrO}_2 \cdot 2\text{H}_2\text{O}$.



Double salts.



None of the Ir haloid compounds has been gasified. The formulæ are given from the analogies between these compounds and those of other Pt metals. The only binary compounds of Pt metals which have been gasified are OsO_4 and RuO_4 . The acid H_2IrBr_6 has been obtained. The

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dissolves in some acids, but no salts have thus been prepared. **Alkaline iridates** appear to exist. There are indications of the existence of an oxide with less O than Ir_2O_3 , probably IrO . When Ir black is heated in O , Ir_2O_3 is formed, the oxide decomposes again at c 1000° . IrO_2 is the most

E

stable oxide, $\text{Ir}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ rapidly absorbs O from the air, and passes into $\text{IrO}_2 \cdot x\text{H}_2\text{O}$.

IRIDIOUS OXIDE (*Iridium monoxide*) IrO When IrCl_3 is boiled with KOH aq a black powder separates, this is probably IrO (Berzelius, P 13, 479) Claus (A 59, 249) thinks that this oxide is obtained by heating one of the double salts of IrSO with K_2CO_3 in a stream of CO_2 . On adding KCl aq to $\text{IrCl}_3 \cdot 3\text{NaCl}$ aq, and filtering, a solution of IrCl_2 is obtained, when K_2CO_3 aq is added to this liquid (or to $\text{IrCl}_3 \cdot 3\text{KCl}$ aq) a greyish green pp is formed, which is probably a hydrate of IrO , the pp is sol in excess of K_2CO_3 aq (Berzelius, *lc*). A few salts derived from IrO are known, e.g. $\text{IrSO}_2 \cdot 4\text{H}_2\text{O}$, they are not, however, formed directly from the oxide, but by reducing $\text{IrO}_2 \cdot 2\text{H}_2\text{O}$ or chloriridates by SO_2 .

IRIDIC OXIDE AND HYDRATE (*Iridium dioxide*) IrO_2 and $\text{IrO}_2 \cdot 2\text{H}_2\text{O}$ By adding excess of alkali to IrCl_3 aq and heating, a heavy indigo blue pp $\text{IrO}_2 \cdot 2\text{H}_2\text{O}$ is produced. The same compound is obtained by using IrCl_3 aq in place of IrCl_3 , and allowing the pp to stand in the air, when it absorbs O. The pp is soluble in HCl aq with formation of IrCl_4 , it is insoluble in dilute H_2SO_4 aq or HNO_3 aq. When the hydrate is heated in a stream of CO_2 , IrO_2 remains as a black powder quite insoluble in acids (Claus, A 59, 249). No salts corresponding with IrO_2 have been prepared. By adding CaO aq to a solution of $\text{IrO}_2 \cdot \text{H}_2\text{O}$ in HCl aq a blue pp is obtained, which is a compound of IrO_2 with CaO . When Ir black is fused for some time with KNO_3 , and the blackish-green mass is treated with water, part dissolves, forming a deep indigo blue solution, and part remains as a black crystalline powder. The quantity of K in this powder is variable, but the ratio of IrO is always 1.3 (Claus, A 59, 249). The powder is probably an iridate of K, $\text{IrO}_2 \cdot x\text{K}_2\text{O}$.

IRIDO-IRIDIC OXIDE AND HYDRATES (*Iridium sesquioxide*) Ir_2O_3 and $\text{Ir}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ The oxide Ir_2O_3 is obtained by heating $\text{IrCl}_3 \cdot 2\text{KCl}$ with 2 parts K_2CO_3 , or Na_2CO_3 , in a stream of CO_2 , washing the fused mass with boiling water, and then with water containing NH_4Cl , heating to remove NH_4Cl , treating with acid to remove alkali, and again washing with water (Claus, A 59, 249). A hard blue black powder, decomposed by heating to c. 1000° into Ir and O (Deville & Debray, C R 1878 441, cf. Dissociation, vol. II p 397). Reduced to Ir by H at the ordinary temperature. The hydrate $\text{Ir}_2\text{O}_3 \cdot 8\text{H}_2\text{O}$ ($= \text{Ir}_2\text{O}_3 \cdot \text{H}_2\text{O}$) is obtained by adding to $\text{IrCl}_3 \cdot 2\text{NaCl}$ aq so much KOH aq that all remains dissolved, and then heating or ppg by alcohol. Black pp insol in acids except conc HCl aq, which dissolves it very slightly. By adding a little KOH aq to solution of an alkaline chloriridate, and quite filling a flask with the liquid, a yellow green pp forms, which is probably $\text{Ir}_2\text{O}_3 \cdot 5\text{H}_2\text{O}$, it oxidises very easily to the blue $\text{IrO}_2 \cdot 2\text{H}_2\text{O}$, easily soluble in the slightest excess of alkali. De Boisbaudran (C R 96, 1236, 1406, 1551) describes a violet coloured pp, which is probably $\text{Ir}_2\text{O}_3 \cdot x\text{H}_2\text{O}$, obtained by adding alkali to $\text{Ir}_2(\text{SO}_4)_3 \cdot 3\text{K}_2\text{SO}_4$ aq (this salt is formed by heating Ir compounds with KHSO_4 , and treating the product with K_2SO_4 aq). A few salts, and some double and basic salts, corresponding with Ir_2O_3 are known, they are pro-

duced indirectly. The sulphites and double sulphites are produced by reducing $\text{IrO}_2 \cdot 2\text{H}_2\text{O}$ or chloriridates by SO_2 . By adding CaO aq to IrCl_3 aq a yellow pp of $\text{Ir}_2\text{O}_3 \cdot 8\text{CaO}$ is produced (Claus, A 59, 249). Compounds of Ir_2O_3 with NO_2 and with nitrites are described by Gibbs (B 4, 280).

IRIDIUM, oxychloride of An oxychloride of Ir is said by Berzelius (P 13, 484) to be formed as a yellowish-brown pp by adding a small quantity of HgNO_3 aq to $\text{IrCl}_3 \cdot 2\text{KCl}$ aq, HgCl_2 goes into solution, addition of more HgNO_3 causes ppg of HgCl .

IRIDIUM, phosphide of. When Ir is very strongly heated with $\frac{1}{2}$ its weight of P, the whole melts. By heating the product with CaO the P is removed. Ir and P probably combine when heated together, but no phosphide of Ir has been isolated.

IRIDIUM, salts of Very few salts of Ir are known other than the haloid compounds. Iridous sulphite, $\text{IrSO}_2 \cdot 4\text{H}_2\text{O}$, and some double sulphites, irido-iridic sulphite, $\text{Ir}_2(\text{SO}_4)_3 \cdot 6\text{H}_2\text{O}$, and several double and basic sulphites, and a few other salts (chiefly double and basic), corresponding with Ir_2O_3 , have been isolated. None of these salts is obtained from the corresponding oxide, the sulphites, which are the best known salts, are formed by reducing $\text{IrO}_2 \cdot 2\text{H}_2\text{O}$ or chloriridates, by SO_2 aq.

IRIDIUM, sulphides of Three sulphides of Ir are known, IrS , Ir_2S_3 , and IrS_2 . Ir combines with S when heated with it.

IRIDOUS SULPHIDE IrS (*Iridium monosulphide*) A blue black solid, by heating IrS_2 or IrS_3 (Berzelius, P 13, 487, Böttger, J pr 3, 227). The pp obtained by passing H_2S into a solution of an iridous compound is probably IrS . Does not decompose by heating in absence of air, roasted in air gives SO_2 and a basic sulphate. The ppd sulphide dissolves in KHSAq .

IRIDIC SULPHIDE IrS_2 (*Iridium disulphide*) A black powder, obtained by strongly heating Ir black with S and Na_2CO_3 , and washing with water (Fellenberg, P 50, 66). Heated in air gives SO_2 and Ir . Berzelius (*lc*) obtained a dark brownish-yellow pp by passing H_2S into IrCl_3 aq, heated, this gave IrS .

IRIDO-IRIDIC SULPHIDE Ir_2S_3 (*Iridium sesquisulphide*) A brown black pp by passing H_2S into solution of an irido-iridic compound. Heated gives SO_2 and S, and leaves IrS (Berzelius, *lc*).
M M P M

IRIDOLINE $\text{C}_{10}\text{H}_8\text{N}$ ($252^\circ - 257^\circ$ S G 15 1072). A base occurring in coal tar oil (G Williams, Tr E 21, 377, C J 16, 375).

IRIDOSMIUM v *Iridium*, Alloys of, p 47.

IRIS CAMPHOR $\text{C}_{15}\text{H}_{12}\text{O}_2$. Occurs in the root of *Iris florentina* (Dumas, A 15, 158). Pearly plates, volatile with steam. Insol water. Flückiger (Ar Ph [3] 8, 481), by distilling iris root with steam, obtained crystals of myristic acid saturated with a fragrant oil.

IRISH PEARL MOSS v CARAGHEEN MOSS.

IRISIN $\text{C}_6\text{H}_8\text{O}_4$ aq $[\alpha]_D = -51^\circ$. A carbohydrate extracted from the rhizomes of the water lily (*Iris pseudacorus*) with water and then ppg with lead acetate. Probably identical with graminin (Wallach, A 284, 364, B 21, 396).

Properties — More strongly laevorotatory than inulin — 1. Does not reduce Fehling's solu-

tion Is very easily converted into sugar (chiefly levulose) by weak acids Gives no colouration with iodine V sol strong HI solution from which solution NaOH separates iodoform Also sol HCl, and NaOH Irisin is nearly 4 times as soluble as inulin in water at 22°. Under the microscope the globules of irisin resemble those of inulin, but do not exhibit double refraction

IRON Fe At w 55.9 Mol weight unknown [1550°] (Pouillet), [1587°] (Daniell, Tr 1880), [1600°] (Pictet, P M 1879 446) S G determinations vary from 6.95 (Roberts a Wrightson, A Ch [5] 30, 274) to c 8.2 Almost pure Fe has S G c 7.85 at 16° according to Caron (C R 70, 1263), 8.139 according to Chandler-Roberts (C N 31, 137) S H 111641 at 0°, 112359 at 50°, 113795 at 100° (Byström, *Oefversk Vetensk Förhandl* [Stockholm, 1860] 17, 307, v also Weinhold, P 149, 214) C E 0000.233 at 18° to 100° (Kopp, A 81, 1, v also Fizeau, A Ch [4] 2, 143, C R 68, 1125) T C (Ag=100) 11.9 (Wiedemann a Franz, P 89, 497) E C (Hg at 0°=1) 9.68 at 0°, 6.19 at 100° (Lorenz W 13, 422, 582) Crystallises in regular system (Fuchs, A 84, 257) Emission spectrum is very complex (Angström, P 94, 141, Cornu, *Spectre Normal* [Paris, 1881], Liveing a Dewar, Pr 29, 402, 32, 402) Some of the thermal data for Fe are presented in the following table (Th 3, 293) —

X [Fe, X, Aq]

Cl = 99,950

Br = 78,070

I = 47,650

[Fe, Cl, Aq] = 127,720, [Fe, O, 3H²O] = 191,150,

[Fe²O²H², 3H SO⁴Aq] = 33,840,

[Fe²H²SO⁴Aq] = 24,840

For further details regarding physical properties v Rammelsberg's *Hand der Kryst physikal Chemie*, 1, 193 For the physical properties of different kinds of pig iron, wrought iron, and steel, v DICTIONARY OF APPLIED CHEMISTRY

Occurrence—The metal itself is found in small quantities in rocks of volcanic origin and in lavas, it also occurs in meteorites Compounds of iron are very widely distributed, and occur in immense quantities The spectroscopie shows the existence of iron (or iron compounds) in the sun and other stars The chief ores of iron are *hematite*, FeO, *brown iron ore*, Fe₂O₃·H₂O, *yellow ochre*, Fe₂O₃·2H₂O, *magnetic iron ore*, Fe₃O₄, *spathe ore*, FeCO₃, *pyrites*, FeS₂ (the 'formulae express approximately the compositions of the ores)

Preparation—Commercial iron, whether pig iron, malleable iron, or steel, always contains more or less C and Si, and generally small quantities of P and S, sometimes also traces of Mn, Ti, Ni, Co, Cu, Sb, and As 1 Wöhler (A 94, 126, 95, 192) prepares Fe₂O₃ by heating FeSO₄ crystals with 2-3 parts NaCl in a crucible and washing with water, and reduces the Fe₂O₃ by heating in H (cf G de Claubry, C C 1859 214, Luca, C R 1851 332, 1352, 202). Moissan (C R 89, 176) says the reduction must be carried out at c 700° —2 By reducing FeCl₃ in a stream of H (Peligot, C R 19, 670), or in neutral solution at b point by Zn (Capitaine, C R 9, 757), or by Zn vapour at a high temperature (Poumarède, C R 29, 518) —8 Troost melts

pig iron in a lime crucible by the O-H flame (Bl [2] 9, 250) —4 A mixture of dry Na₂SO₄ and FeSO₄ is heated in a Pt crucible so long as SO₂ comes off, the residue is washed with cold water, and the crystalline Fe₂O₃ is reduced in H in a Pt crucible, and the Fe is melted in the O-H flame (Ma'ithessen a Prus Szczepanowski, C N 20, 501) —5 By electrolysis FeSO₄ saturated with NH₄Cl, Fe appears at the negative pole (a large iron plate), it contains H and other gases, which may be removed by heating (Varrentrap, D P J 187, 152, Lenz, J pr 108, 438)

Properties—A lustrous, greyish white metal Crystallises in regular cubes or octahedra Annealed wrought iron wire is extremely tenacious. Such mechanical treatment as hammering, bending, or torsion changes many of the physical properties of iron The physical properties which have been determined are for the most part those of iron containing small quantities of foreign substances Iron is magnetic (cf Faraday, P 70, 24, Gore, P M [4] 40, 170) Pure, or almost pure, Fe is softer and more malleable, but less tenacious, than ordinary malleable iron The iron obtained by reducing Fe₂O₃ in H at temperatures below c 600° is pyrophoric, the powder obtained at c 700° is non pyrophoric (Moissan, C R 89, 176) By passing a weak electric current through FeSO₄·Aq mixed with MgSO₄, Lenz (C C 1870 188) obtained a greyish, non crystalline, very brittle mass, easily pulverised by the finger, this iron contained c 200 times its volume of gases, chiefly H (v *Iron, hydrides of*), mixed with CO, CO₂, N, and H₂O vapour, by heating *in vacuo* the gases were removed, and the iron then resembled Pt in appearance Iron is unchanged in dry O, but in moist O or in ordinary air it is slowly oxidised to FeO, xH₂O, heated in air or O a mixture of FeO and Fe₂O₃ is produced Iron combines directly with the halogens, also with S, C, B, Si, P, As, it forms alloys with many metals Iron dissolves in the common mineral acids with formation of salts Finely divided Fe decomposes water at 100°

The atomic weight of Fe has been determined (1) by analysing and determining V.D. of FeCl₃ and FeCl₂, (2) by determining S.H. of Fe, (3) by reducing Fe₂O₃ in H (Berzelius, P 8, 165, Svanberg a Norlin, A 50, 432, Erdmann a Marchand, J pr 33, 5, Rivet, A Ch [3] 30, 192), (4) by transforming Fe into Fe₂O₃ (B, A 50, 432, S a N, lc, Maumené, A Ch [3] 30, 380), (5) by determining the Cl in FeCl₃ and in FeCl₂ (Dumas, A Ch [3] 55, 157) The atom of Fe is trivalent in the gaseous molecule FeCl₃, and divalent in the gaseous molecule FeCl₂ (v *Iron, chlorides of*)

Iron is distinctly a metallic element, it replaces the H of most acids, forming two series of salts, the simplest formulae for which are FeX₂ and FeX₃, respectively, X=Cl, NO₃, SO₄, PO₄, &c The ferrous salts, FeX₂, are easily oxidised to ferric salts, FeX₃, very many salts, both normal and basic, of both series have been isolated, numerous double salts are also known. Fe₂O₃ forms compounds with several oxides more basic than itself, e.g. with K₂O, BaO, CaO, MgO, these compounds may be

regarded as *ferrites* derived from the hydroxide $\text{Fe}_2\text{O}_3\cdot\text{H}_2\text{O}$ (= $\text{Fe}_2\text{O}_3\cdot\text{H}_2\text{O}$). There are also some salts known as *ferrates*, e.g. K_2FeO_4 and BaFeO_4 , which may be looked on as derived from the hypothetical *ferric acid* H_2FeO_4 , but neither this acid nor its anhydride, FeO_3 , has been isolated (v *FERRATES and FERRITES*, vol. II pp 546-7). Fe_2S_3 , like Fe_2O_3 , forms some double compounds with more basic sulphides, these double compounds may be regarded as *thioferrites* (v *Iron, sulphides of*). Fe is closely related to Co and Ni, these three metals show some marked analogies with the Pt metals, Fe is also related to Cr and Mn, and it shows some resemblance to Al and the other earth metals (v *IRON GROUP OF ELEMENTS*, also *NOBLE METALS*, and *CHROMIUM GROUP OF ELEMENTS*, and *EARTHS, METALS OF THE*).

Reactions and Combinations—1 Heated in air or oxygen Fe is oxidised to Fe_2O_3 and Fe_3O_4 . 2 Unchanged in dry air at ordinary temperature, but rusts in moist air to $\text{Fe}_2\text{O}_3\cdot x\text{H}_2\text{O}$. 3 Finely divided Fe decomposes water at 100° (v Ramann, B 14, 1433), and rapidly decomposes steam. Compact Fe decomposes steam at red-heat, Fe_2O_3 is produced. Iron slowly rusts in contact with water and ordinary air, the formation of $\text{Fe}_2\text{O}_3\cdot x\text{H}_2\text{O}$ proceeds slowly at first, and then more rapidly, probably because the Fe and Fe_2O_3 exert an electrolytic action on the water, presence of ammoniacal salts increases the rate of rusting, alkalis and alkaline carbonates hinder the rusting. According to Calvert (C N 23, 98) Fe does not rust in water if CO_2 is absent. For an account of the retarding or hastening action of various salts on the rusting of Fe v Wagner, D P J 218, 70. Various processes have been used for preventing iron rusting, covering the iron with Zn is frequently done, Barff & Bower heat the iron to c 650° in water vapour, whereby a hard protecting film of Fe_2O_3 is formed on the surface of the iron. 4 Fe decomposes solution of hydrogen peroxide, forming $\text{Fe}_2\text{O}_3\cdot\text{H}_2\text{O}$, the Fe becomes covered with bubbles of O (Weltzien, A 138, 129). 5 Iron dissolves in dilute sulphuric or hydrochloric acid with evolution of H . Cold conc sulphuric acid is almost without action on Fe, when heated, SO_2 is evolved, and $\text{Fe}_2(\text{SO}_4)_3$ formed. Dilute nitric acid dissolves Fe, forming $\text{Fe}(\text{NO}_3)_2$ and NH_4NO_3 , with evolution of N oxides. Conc nitric acid is almost without action on Fe, v *infra*, Passivity of iron. 6 Heated repeatedly with caustic soda solution Fe forms crystals of $\text{Fe}_2\text{O}_3\cdot\text{H}_2\text{O}$ (Brunck & Graebe, B 13, 725). 7 When Fe is heated to redness in a stream of ammonia a nitride of Fe is formed, Fe_3N_2 , according to Stahl Schmidt (P 125, 37). 8 Fe heated in a closed tube with solution of sulphur dioxide to 200° forms solution of FeSO_4 and crystals of FeS_2 (Geitner, A 129, 350). 9 Fe is oxidised to Fe_2O_3 , or K_2FeO_4 (v vol. II p 547) by fusion with potassium nitrate. 10 Fe combines with the halogens, with arsenic, boron, carbon, hydrogen, nitrogen, phosphorus, and sulphur (v *Iron, chlorides of, borides of, carbides of, &c.*), it also forms alloys with several metals (v *Iron, alloys of*).

Passivity of iron Iron is not acted on by conc nitric acid, iron which has been immersed in conc nitric acid does not dissolve in dilute

nitric acid, nor does it ppt Cu from $\text{CuSO}_4\cdot\text{Aq}$; such Fe is said to be *passive*. Co and Ni, and perhaps Bi, also show passivity. According to Varenne (A Ch [5] 19, 251, 20, 240) immersion of iron in acid of 100 p.c. HNO_3 is not accompanied by evolution of any gas, if the acid has SG 1.882 (c 63 p.c. HNO_3) NO is evolved copiously for 3 to 20 seconds and then ceases, in both cases the Fe becomes passive. Acid of less SG than 1.299 (c 47 p.c. HNO_3) does not produce passivity. If part of a rod of Fe is dipped into conc acid, and then the whole is carefully immersed in a dilute acid, the whole rod is passive. These facts are explained by Varenne by supposing that a gaseous film is deposited on the surface of the iron, and protects the iron from the action of the acid, the gas dissolves in the more dilute acid, in the case of the partly immersed rod the gas bubbles are removed from one part and then adhere to the other part of the rod. This view was also upheld by Mousson (P 39, 330), but was controverted by Faraday and Schönbein (P 39, 342), and Beetz (P 67, 286, 365). Ramann (B 14, 1430) considers the passivity to be due to a layer of Fe_2O_3 , which is soluble in dilute, but insoluble in conc, nitric acid. R thinks that immersion in the acid produces $\text{Fe}(\text{NO}_3)_2$, and that this then reacts with the Fe to produce Fe_2O_3 and NH_4NO_3 , iron wire, according to R, becomes passive when partly heated, also by making it the positive electrode in an O containing liquid. Various oxidising agents acting on Fe render it passive, e.g. HClO_4 , CrO_3 , H_2O_2 (Keir). The passivity of Fe is removed by strongly rubbing the iron, or heating it in reducing gases, also by bringing it into contact with Zn. According to Saint Edme (C R 106, 1079) commercial sheet Ni is passive in ordinary $\text{HNO}_3\cdot\text{Aq}$, Fe in contact with Ni becomes passive in the acid, passive Ni remains passive when heated to bright redness in H , whereas Fe loses its passivity.

Detection and Estimation—Fe compounds are detected by giving a reddish green colour to the hot borax bead in the oxidising flame, which becomes bottle green to nearly colourless when cold, also by the ppn. of brown red $\text{Fe}_2\text{O}_3\cdot x\text{H}_2\text{O}$ by alkalis from ferric compounds, $\text{K}_2\text{FeCy}_4\cdot\text{Aq}$ gives a deep blue pp with ferrous salts, and no pp but a brownish colour with ferric salts. Fe is estimated, gravimetrically, by ppn as $\text{Fe}_2\text{O}_3\cdot x\text{H}_2\text{O}$ and weighing as Fe_2O_3 , volumetrically, by titration with $\text{KMnO}_4\cdot\text{Aq}$ or $\text{K}_2\text{Cr}_2\text{O}_7\cdot\text{Aq}$. For details and for other methods v *ANALYSIS OF ANALYSIS*.

Metallurgy of iron Pure iron is too soft and has too little tenacity for use in making machinery, &c. Ordinary iron is divided into three kinds—pig or cast iron containing from 2 to 5.75 p.c. C, besides small quantities of Si, P, S, and traces of metals other than Fe (chiefly As, Ti, V, Cr, Mn, Cu), malleable or wrought iron containing less than 5 p.c. C, and steel containing about 1 p.c. C. Pig iron is obtained by very strongly heating Fe ores with lime and coal in large furnaces, and blowing in air at the bottom of the furnaces, the C of the coal is burnt chiefly to CO , and this reacting with oxides of Fe produces Fe, which then combines with, or it may be only dissolves, carbon, the C seems to be produced by a reaction between the partly reduced Fe_2O_3

and CO (v Bell, *C J* [2] 7, 203) The reduction of Fe_2O_3 is probably aided by cyanides present in the furnace The greater part of the foreign matter of the iron ores is removed in the form of a fusible slag consisting of silicate of Fe, Ca, Mg, Al, Mn, &c Wrought iron is obtained by *puddling* cast iron, in this process the cast iron is strongly heated along with lumps of Fe_2O_3 in a reverberatory furnace, the Fe_2O_3 is partially deoxidised, and the O is used in burning the C, Si, S, and P, the C being thus almost entirely removed as CO or CO_2 Steel is obtained by completely removing the C from wrought iron by blowing in air, and then adding a sufficient quantity of Fe containing a known quantity of C Steel is also sometimes made by causing wrought iron to take up C by strongly heating the iron with charcoal Cast iron is more fusible, harder, and more brittle than wrought iron, which is more ductile and malleable than cast iron Steel combines the malleability and ductility of wrought iron with the fusibility of cast iron, it is also very hard and elastic Cast iron is divided into two kinds *grey cast iron*, granular, with low malleability, SG c 7.1, melting at c 1600° , this variety contains C partly in combination with Fe and partly uncombined as graphite, *white cast iron*, harder but not so strong as grey, SG c 7.5, melts at a lower temperature than grey, this variety contains only combined C By melting grey, and cooling rapidly, white, cast iron is obtained, by melting white, and cooling slowly, grey, cast iron is produced For details of the metallurgy of iron v DICTIONARY OF APPLIED CHEMISTRY

Iron, alloys of Iron alloys with many metals, the softer metals—Ag, Sn, Zn &c—become harder and more tenacious when alloyed with a few parts of iron per thousand, the alloys may generally be prepared by fusing the various metals with iron wire, a little Fe_2O_3 being added to remove C from the iron

Alloys with *aluminum* have been described, approximately AlFe (Wohler, A 115, 102) approximately AlFe, and AlFe, (Calvert a Johnson, *J* 1855, 326) Sonnenschein (*J pr* 66, 168) described an alloy with *lead* having a composition approximating to Pb_2Fe An *amalgam* of iron is formed by mixing finely divided Fe with Na amalgam, adding a little water, and squeezing out excess of Hg, the crystals correspond in composition to Hg_2Fe_2 , they slowly undergo decomposition, rapidly when much compressed (Raman, *B* 14, 1433) Alloys with the following metals have been described—Sb, Cu (Musket, *P M* [3] 6, 81), Mn (M, *lc*), Mo (Heine, *J pr* 9, 176, Stromeyer, *P* 28, 551, Wiggins, *P* 28, 565, Steinberg, *J pr* 18, 379), Ni, Pd (Faraday a Stodart, *Tr* 1822 254), Pt (Schönbein, *P* 42, 17, Boussingault, *A Ch* [2] 53, 441), K (Calvert, *P M* 1855), Rh (F a S, *lc*), Sn (Lassaigne, *J Chim méd* 6, 609), Ti (Riley, *C J* 16, 387), W (Bernoulli, *P* 21, 573), Zn (Percy, *Metallurgy*, 2, 153)

Iron, arsenides of The minerals *arsenical pyrites* and *mispickel* are compounds of Fe with As and with As and S respectively, their compositions approximate to the formulae FeAs_2 and FeAsS Brittle masses are obtained by fusing together As and Fe out of contact with air, but they have not been much examined

Iron, borides of When Fe is heated in a stream of BCl_3 vapour, a hard, white substance is obtained, which is decomposed by boiling water forming H_2BO_3 (?) and Fe_2O_3 , and evolving H (Fremy, Wurtz's *Dict de chim* 1, 1417) by heating ferric borate in H, Lassaigne (*J Chim mét* 3, 535) obtained a similar compound, no analyses are given

Iron, bromides of Two exist, FeBr_2 and FeBr_3 , both are produced by the direct union of their elements Neither has been gasified, the formulae are given because of the analogies between these salts and FeCl_2 and FeCl_3 (v *Iron, chlorides of*)

FERROUS BROMIDE (*Protobromide of iron*) FeBr_2 A yellow solid, obtained by heating excess of Fe in Br vapour, soluble in water, solution deposits crystals $\text{FeBr}_2 \cdot 6\text{H}_2\text{O}$, decomposed by heating in air to FeBr_3 and Fe_2O_3 (Scheufelen, *A* 231, 156) Thomsen gives $[\text{FeBr}_2, \text{Aq}] = 78,070$ (*Th* 3, 294)

FERRIC BROMIDE (*Sesquibromide of iron*) FeBr_3 Dark red crystals, obtained by heating Fe in excess of Br vapour, may be sublimed, with partial decomposition, out of contact with air, deliquescent Aqueous solution very easily partially reduced by heating to FeBr_2 (L de Koninck, *Zeit anorgan Chem*, 1889 149)

Iron, carbides of Whether pig iron is a carbide of iron, or a mixture of carbides and free C, cannot yet be regarded as finally settled The presence of a few per cents of C in iron very much modifies the properties of the whole, by making it more fusible, if from 2 to 1.5 p c C is present, the mass is only fusible with difficulty, but is very hard and tough, it has the properties of wrought iron or steel, according to the quantity of C Iron takes up C directly, and by so doing has its properties changed On the temperatures at which carburization of Fe occurs with amorphous C, diamond, and graphite, v Hempel, *B* 18, 998

The following carbides of Fe have been described, but the investigation of none of these bodies is complete—(1) Fe_3C , a brittle, fusible mass, by melting together an intimate mixture of equal parts of Fe and C (Faraday a Stodart, *G A* 66, 183), (2) FeC_2 , a black pyrophoric powder, by heating H_2FeCy_3 , or $(\text{NH}_4)_2\text{FeCy}_6$, out of contact with air (Berzelius), or by fusing K_2FeCy_6 , and washing away the KCN produced (Rammelsberg), (3) FeC , was obtained by Karsten by the action of dilute acids on grey pig-iron (*S* 68, 182), (4) FeC , by the action of Br or I on pig iron (Berthier, *J* 1862 127)

Iron, chlorides of Two chlorides of iron, FeCl_2 and FeCl_3 , have been isolated, and a third, Fe_2Cl_4 , perhaps exists When Fe is heated in a stream of Cl, both FeCl_2 and FeCl_3 are produced

FERROUS CHLORIDE FeCl_2 (*Iron dichloride* or *protochloride*) Mol w 126.64 (v *infra*) SG 2.528 (Fulhol, *A Ch* [3] 21, 415), 2.988 at 17.9° (Clarke's *Specific Gravity Tables* [new ed.] 24) VD 62.79 at 1300° – 1400° , 61.55 at 1400° – 1500° (Nilson a Pettersson, *C J* 53, 828) VD at yellow heat 94.4° (V Meyer, *B* 17, 1835), calc for FeCl_2 , 63.32, for Fe_2Cl_4 , 26.64 (v *infra*). HF $[\text{FeCl}_2] = 82,050$, $[\text{Fe}_2\text{Cl}_4, \text{Aq}] = 99,950$; $[\text{FeCl}_2, 4\text{H}_2\text{O}] = 15,150$ (*Th* 3, 293)

Formation.—1. By heating excess of Fe filings

or wire in Cl.—2 By partially reducing FeCl_2 in H (Wöhler, *A* 4, 255)—3 By dissolving Fe in HClAq out of contact with air, evaporating, and drying in a stream of H—4 By heating Fe filings with NH_4Cl out of contact with air

Preparation—Fe wire is heated in a stream of dry HCl, and the product sublimed in the same gas (Wöhler & Liebig, *P* 21, 258)

Properties and Reactions—White, lustrous, six-sided, deliquescent, tablets Heated very strongly in N forms FeCl_2 and Fe (V Meyer, *B* 17, 1335) Heated in O, forms Fe_2O_3 with evolution of Cl Oxidises readily in air to a mixture of FeCl_2 and Fe_2O_3 Heated to redness in steam forms Fe_2O_3 , HCl, and H Heated to redness in a stream of NH_3 , Fe_3N_4 is produced (Fremy, *C R* 52, 321) Heated with PH_3 forms HCl and Fe phosphide A mixture of FeCl_2 and O is reduced by Zn vapour with separation of Fe in crystals (Poumarède, *C R* 29, 520) S c 50 at 80° , less sol in HClAq

Molecular weight of gaseous ferrous chloride By volatilising in HCl at a yellow heat, V Meyer got values for V D which were about the mean between those calculated for FeCl_2 and Fe_2Cl_2 (*B* 17, 1335), calculated for $\text{FeCl}_2 = 63.32$, for $\text{Fe}_2\text{Cl}_2 = 126.64$, observed 92.32 and 96.51 Meyer concluded that molecules of Fe_2Cl_2 probably exist at lower temperatures and molecules of FeCl_2 at a very high temperature Nilson & Pettersson (*C J* 53, 827) obtained values for V D at $c 1400^\circ$ agreeing with that calculated for FeCl_2 , calculated = 63.32, observed at $1300^\circ - 1400^\circ = 62.79$, at $1400^\circ - 1500^\circ = 61.55$ There can be no doubt then that molecules of the composition FeCl_2 exist at $c 1300^\circ - 1500^\circ$

Combinations—1 With water to form the hydrates $\text{FeCl}_2 \cdot 2\text{H}_2\text{O}$ and $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$ The former is obtained by saturating warm conc HClAq with FeCl_2 or by passing HCl into saturated FeCl_2Aq , the latter is produced by dissolving Fe in HClAq and concentrating out of contact with air $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$ forms blue green, monoclinic, deliquescent, transparent crystals, S G 1.93—2 With nitric oxide, solution of FeCl_2 , especially an alkaline solution, absorbs NO forming a green-brown liquid which contains $\text{FeCl}_2 \cdot \text{NO}$, NO is removed by warming—3 With ammonia to form $\text{FeCl}_2 \cdot 6\text{NH}_3$ and $3\text{FeCl}_2 \cdot 2\text{NH}_3$ (Rogstadius, *J* pr 86, 810) The former is a white powder, produced by passing NH_3 over FeCl_2 at the ordinary temperature, the latter is formed by slowly heating $\text{FeCl}_2 \cdot 6\text{NH}_3$ to melting By strongly heating $3\text{FeCl}_2 \cdot 2\text{NH}_3$ in NH_3 the nitride Fe_3N_4 is formed (Fremy, *C R* 52, 321)—4 With ammonium chloride to form $\text{FeCl}_2 \cdot 2\text{NH}_4\text{Cl}$, obtained by crystallising mixed solutions of FeCl_2 and NH_4Cl , or, according to Hisinger & Berzelius (*G A* 27, 273) (with evolution of NH_3 and H), by boiling Fe filings with NH_4ClAq (cf. Winkler, *R. P.* 59, 171, Vogel, *J* pr 2, 192) A compound $\text{FeCl}_2 \cdot 2\text{KCl} \cdot 2\text{H}_2\text{O}$ is also known (Schabus, *W. A. B* 1850 475) Compounds with CdCl_2 and HgCl_2 , viz $\text{FeCl}_2 \cdot 2\text{CdCl}_2 \cdot 12\text{H}_2\text{O}$ and $\text{FeCl}_2 \cdot \text{HgCl}_2 \cdot 4\text{H}_2\text{O}$ are also described (v Hauer, *W. A. B* 17, 331)

FERRIC CHLORIDE (*Iron sesquichloride* or *perchloride*) FeCl_3 , Mol w 162.01, and probably also 824.03 (v *infra*) ($280^\circ - 285^\circ$) (Friedel & Crafts, *C R* 107, 801) S G 2.804 at 10.8° (Clarke's *Tables of Specific Gravity* [new ed.],

24) V D varies from $c 165$ at 320° to $c 80$ at $1000^\circ - 1800^\circ$ (v *infra*) Crystallises in hexagonal forms $a = 1.1295$, Nordenskiöld (v. Rammelsberg's *Hand der Krystal physikal Chemie*, 1, 260) H F [FeCl_3] = 96.040, [$\text{FeCl}_3 \cdot \text{Cl}$] = 13.990, [$\text{FeCl}_3 \cdot \text{Aq}$] = 127.720, [$\text{FeCl}_3 \cdot \text{Aq} \cdot \text{Cl}$] = 27.770 (*Th* 8, 293)

Occurrence—In the craters of volcanoes

Formation—1 By heating Fe in a stream of Cl—2 By heating FeCl_2 in Cl—3 By strongly heating FeCl_2 in a wide mouthed vessel, Fe_2O_3 remains and FeCl_3 sublimes—4 By passing HCl over strongly heated amorphous Fe_2O_3 (Deville & Troost, *C R* 45, 821)—5 By heating a mixture of calcined FeSO_4 with CaCl_2 or NaCl The product obtained by boiling down a solution of Fe_2O_3 in HCl always contains oxychloride

Preparation—Fine iron wire is strongly heated in a hard glass tube in a stream of dry Cl, and the product is sublimed in the same gas

To prepare a solution of FeCl_3 , hydrated Fe_2O_3 is dissolved in hot HClAq , the solution is decanted, Cl is passed in until K_2FeCl_6 ceases to give any trace of blue colour, and excess of Cl is removed by warming, or 2 parts warm HClAq are saturated with Fe (filings or wire), to the clear decanted liquid 1 part HClAq is added, the liquid is warmed and HNO_3Aq is added little by little until all FeCl_2 is oxidised to FeCl_3 , excess of HNO_3 is removed by evaporation with a little HClAq Geissler (*Ar Ph* [2] 58, 159) recommends to evaporate the ferrous solution to S G 1.3 at 17° and then to add HNO_3Aq to the cold solution, when oxidation proceeds rapidly

Properties—By slowly cooling the vapour, FeCl_3 condenses in hexagonal crystals which appear very dark red by transmitted, and greenish by reflected, light (Deville & Troost, *C R* 45, 821) As ordinarily prepared FeCl_3 forms a brownish black crystalline mass Volatilises at somewhat over 100° Melts at $306^\circ - 307^\circ$ in a sealed tube (Carnelley & Williams, *C J* 37, 126) Deliquescent, conc solution in water is dark brown and somewhat viscid, it becomes limpid and yellow on dilution, a very dilute solution is colourless when cold (v *Reactions*, No 4)

Franz (*J* pr [2] 5, 283) gives the following table showing S G and percentage composition of FeCl_3Aq at 17.5° —

S G	p.c. FeCl_3	S G	p.c. FeCl_3	S G	p.c. FeCl_3
1.0146	2	1.1746	23	1.3870	42
1.0292	4	1.1950	24	1.4118	44
1.0439	6	1.2155	26	1.4377	46
1.0578	8	1.2365	28	1.4617	48
1.0734	10	1.2568	30	1.4876	50
1.0894	12	1.2778	32	1.5153	52
1.1054	14	1.2988	34	1.5439	54
1.1215	16	1.3199	36	1.5729	56
1.1378	18	1.3411	38	1.6023	58
1.1542	20	1.3622	40	1.6317	60

Schult (*Forh skand Natur*, 1868 452) obtained the following results at 14.6° —

S G FeCl_3Aq	p.c. FeCl_3	S G FeCl_3Aq	p.c. FeCl_3
1.0882	4.65	1.339	33.25
1.0918	10.45	1.3624	36.95
1.1517	16.80	1.4361	41.0
1.2107	22.64	1.554	49.61
1.2318	24.60		

FeCl_3 is soluble in alcohol or ether, the solution

easily decomposes in sunlight with formation of FeCl_2 .

Molecular weight of gaseous ferric chloride. — Deville a Troost (*C R* 45, 821) found the V D of ferric chloride at 440° to be 162.7, this corresponds to the formula Fe_2Cl_3 . Grönwald a V Meyer (*B* 21, 687) made a series of determinations of V D, in an atmosphere of N, with the following results 151.75 at 448° (mean of 4), 138.5 at 518° (mean of 3), 121.3 at 606° (mean of 6), 78.66 at 750° (mean of 2), 71.2 at 1036° , 77.6 at 1077° , 4.8 at 1300° . These numbers seem to indicate a gradual resolution of molecules of Fe_2Cl_3 into FeCl_3 , none of the results, however, agrees exactly with the value calculated for FeCl_3 (81.905), but this was to be expected, as G a M showed that even at 518° ferric chloride is partially resolved into FeCl_2 and Cl in an atmosphere of N. According to Friedel a Cratz (*C R* 107, 301) ferric chloride is not dissociated at 440° in an atmosphere of Cl. F a C made a series of determinations of V D in Cl, using a modification of Dumas' method, the results are appended — 165.1 at 321.6° , 180.4 at 325.2° , 174.2 at 356.9° , 171.5 at 357° , 168.7 and 163.5 at 412.2° . These values are all somewhat greater than 162.1, the value required by the formula FeCl_3 . The most probable conclusion from all the results seems to be that at high temperatures, c 700° and upwards, the vapour of ferric chloride consists of molecules of FeCl_3 , but that as temperature falls these molecules are mixed with more complex molecules, some of which at any rate have the composition Fe_2Cl_3 (cf Young, *N* 39, 195).

Reactions. — 1 When FeCl_3 is heated in oxygen, Fe_2O_3 and Cl are produced (cf Schulze, *J pr* [2] 21, 407). — 2 Heated in hydrogen, FeCl_2 is formed (Wohler, *A Suppl* 4, 255). — 3 Heated in water vapour, Fe_2O_3 and HCl are formed. — 4 FeCl_3 dissolves in water, the conc solution is a syrupy dark brown liquid, becoming yellow on dilution, and colourless when very dilute and cold. By heating FeCl_3 aq, colloidal soluble Fe_2O_3 , $x\text{H}_2\text{O}$ and HCl are produced, fairly conc FeCl_3 aq forms oxychlorides, Fe_2O_3 , $x\text{FeCl}_3$, at c 100° , and at higher temperatures Fe_2O_3 is formed. The amount of decomposition and the composition of the products depend on the concentration of the solution and the temperature, very dilute solutions, c 30,000 water to 1 FeCl_3 , are decomposed with formation of colloidal soluble ferric hydrate even without warming (cf Ivor, *hydrated oxides or hydroxides of p 59*). Krecke (*J pr* [2] 3, 286) gives the following table (cf Tichborne, *C N* 24, 123 199, 209, 230, 25, 133, Muller, *J* 1873 40, Fous sereau, *C R* 103, 42) —

FeCl_3 aq containing $\frac{1}{10}$ p.c. FeCl_3 or less is decomposed by light at 5° – 6° (Krecke, *lc*). When FeCl_3 aq is heated in a sealed tube to 250° – 300° it decomposes to Fe_2O_3 and HCl (Senarmont, *C R* 32, 762). From a very dilute cold colourless solution of FeCl_3 , K_2FeCl_6 aq ppts a pure bl. e pp., and NaCl produces no change. If the solution is heated it becomes yellow, and contains soluble ferric hydrate and HCl. From this solution K_2FeCl_6 aq ppts greenish blue solid, and on addition of NaCl a soluble ferric hydrate separates, which, after long contact with NaCl aq, becomes insol in water (Debray, *C R* 68, 913). Evaporation of FeCl_3 aq is accompanied by evolution of some HCl, the residue contains oxychloride or hydrated oxide FeCl_3 is not volatilised from violently boiling FeCl_3 aq containing HCl (Fresenius, *Fr* 6, 92). G Wiedemann connects the decomposition of FeCl_3 aq, and also of other ferric salts, with the different magnetic behaviour of colloidal soluble Fe_2O_3 , $x\text{H}_2\text{O}$, and that hydrate which remains in combination with the acid (*W* 5, 45). — 5 FeCl_3 dissolves in alcohol or ether. An ethereal solution mixed with alcohol and allowed to stand in sunlight loses its yellow colour, and FeCl_3 and chlorinated derivatives of alcohol and ether are formed. — 6 An aqueous solution of FeCl_3 is readily reduced to FeCl_2 , e.g. by As, Sb, Bi, Fe, Pb, or Zn, also by finely divided Pt, more slowly by Pd and Au (cf Béchamp, *C R* 52, 757, Saint Pierre, *C R* 54, 1077). Also reduced by stannous chloride, sulphurous acid, or sulphuretted hydrogen, hydriodic acid reduces dilute FeCl_3 aq slowly (v Mohr, *A* 105, 53). Many organic compounds also reduce FeCl_3 aq, especially in sunlight, e.g. alcohol and ether, or tartaric acid (v Schorax, *B* 3, 11, Poitevin, *C R* 52, 94). — 7 A very little FeCl_3 is said to be formed by passing hydrogen for 48 hours through FeCl_3 aq (Brunner, *J* 1864 125). — 8 FeCl_3 dissolves freshly ppd ferric hydrate forming oxychlorides (q.v.). — 9 When FeCl_3 aq is dropped into solution of potassium nitrite, in an atmosphere of CO, brisk evolution of NO proceeds, and a pp of soluble ferric metahydroxide, Fe_2O_3 , H_2O , separates (Pesci, *G* 18, 183). — 10 FeCl_3 aq reacts with potassium iodide to form FeCl_2 , I, and KCl, probably FeI_3 is at first produced, and is then resolved into FeI_2 and I, and then the FeI_2 reacting with unchanged FeCl_3 forms FeCl_2 and I (v Carnegie, *C N* 60, 87). — 11 When electrolysed, conc FeCl_3 aq gives FeCl_2 at the negative, and Cl with a little O at the positive, electrode.

Combinations. — 1 With water to form various hydrates (1) FeCl_3 , $6\text{H}_2\text{O}$, yellow solid, formed by dissolving 100 parts FeCl_3 in 63.5 parts H_2O (S.G. of solution 1.67 at 25°), or by slowly eva-

P.c. FeCl_3 in solution	Formation of Graham's colloidal hydrate	Formation of colloidal hydrate of P de Saint-Gilles	Formation of oxychlorides	Formation of Fe_2O_3
82	100–130	—	over 100	140
16	100–120	—	"	120
8	100–110	—	"	110
4	90–100	—	90	
2	87	—	87	
1	83	100–130		
.5	75	"		
25	64	"		
125	54	"		
.0625	36	"		

FeCl_3 re formed on cooling

FeCl_3 not re formed on cooling

porating FeCl_2 aq on the water bath, and allowing to crystallise, or by exposing FeCl_2 to the air (Mohr, *A* 29, 178), or by passing HCl over commercial ferric chloride, filtering the liquid which forms through glass wool, and evaporating over KOH *in vacuo* (Engel, *C R* 104, 1708), melts at 35.5° , and solidifies again at 28° (Ordway, *Z* 1866 23) (2) $2\text{FeCl}_2 \cdot 5\text{H}_2\text{O}$, reddish tablets, melting at 81° , formed by melting the hexahydrate, warming to 100° , for some hours, and cooling slowly (Engel, *l c*), or by evaporating FeCl_2 aq to a syrup (SG c 15), with addition of a little HCl aq, and crystallising (Fritzsche, *J pr* 18, 479, Goble, *J Ph* [3] 5, 301, 25, 259) According to Wittstein (*R P* [2] 36, 30) and Ordway (*Z* 1866 23), the hydrate thus obtained is $\text{FeCl}_2 \cdot 3\text{H}_2\text{O}$. The hydrate $2\text{FeCl}_2 \cdot 5\text{H}_2\text{O}$ is said to be formed by placing $\text{FeCl}_2 \cdot 6\text{H}_2\text{O}$ over H_2SO_4 , the hexahydrate at first liquefies and then loses water—2 With *hydrogen chloride and water*, to form $\text{FeCl}_2 \cdot \text{HCl} \cdot 2\text{H}_2\text{O}$, thin, transparent, amber yellow laminae, formed by passing dry HCl gas over $2\text{FeCl}_2 \cdot 5\text{H}_2\text{O}$, saturating the liquid thus produced with HCl at 25° , and cooling to 0° (Engel, *C R* 104, 1708, Sabatier, *Bl* 1881 197)—3 With *ammonia*, to form $\text{FeCl}_2 \cdot \text{NH}_3$. When heated, this compound partly sublimes, and is partly decomposed, giving FeCl_2 —4 With *phosphoric chloride*, to form a brown fusible compound, $\text{FeCl}_2 \cdot \text{PCl}_3$ (Weber, *J pr* 76, 410)—5 With *hydrocyanic acid*, to form a brown liquid, which then crystallises to $\text{FeCl}_2 \cdot 2\text{HCN}$ (Klein, *A* 74, 87)—6 With *cyano gen chloride*, but the compound has not been obtained pure (v Klein, *l c*)—7 With *nitrosyl chloride*, to form $\text{FeCl}_2 \cdot \text{NOCl}$, a dark coloured, very deliquescent mass, obtained by placing dry FeCl_2 in the fumes of *aqua regia* placed over CaCl₂ (Weber, *J pr* 89, 152)—8 With *alkali chlorides*, to form $\text{FeCl}_2 \cdot 2\text{MCl} \cdot \text{H}_2\text{O}$, $\text{M} = \text{K}, \text{Na}, \text{Rb}, \text{NH}_4$. The NH_4 compound forms garnet red crystals, by evaporating mixed solutions of the constituent salts. The crystals are rhombic according to Fritzsche (*J pr* 18, 484), dimorphous according to Gentz (*J pr* 71, 164). The double compounds $\text{FeCl}_2 \cdot 2\text{MCl} \cdot \text{H}_2\text{O}$ are all decomposed by water. The K salt is described by Fritzsche (*J pr* 18, 483), the Na salt by Deville (*C R* 43, 970), the Rb salt, said to be $\text{FeCl}_2 \cdot 3\text{RbCl}$, by Godeffroy (*Ar Ph* [3] 9, 343) According to Kremers (*J pr* 55, 191), a compound of the form $\text{FeCl}_2 \cdot 2\text{MCl} \cdot \text{H}_2\text{O}$, in which M is partly K (12.1 p c), partly Na (16 p c), and partly NH_4 (6.2 p c), is found in certain volcanic craters—9 With *thallium chloride*, to form $\text{FeCl}_2 \cdot 3\text{TlCl}$ (Wöhler, *A* 144, 250)—10 With *magnesium and beryllium chlorides*, to form $\text{FeCl}_2 \cdot \text{MCl}_2 \cdot \text{H}_2\text{O}$, formed by adding MgCl_2 or BeCl_2 to a conc hot solution of FeCl_2 in HCl aq (Neumann, *A* 244, 328)

FEROUS FERRIC CHLORIDE Fe_2Cl_3 ($\text{FeCl}_2 \cdot 2\text{FeCl}_3$) When $\text{Fe}_2\text{O}_3 \cdot \text{H}_2\text{O}$ is treated with a little HCl aq, FeCl_2 goes into solution, and Fe_2O_3 remains, but $\text{Fe}_2\text{O}_3 \cdot \text{H}_2\text{O}$ dissolves wholly in considerable excess of conc HCl aq. When the solution is cooled and placed *in vacuo* over H_2SO_4 , yellow crystals of $\text{Fe}_2\text{Cl}_3 \cdot 18\text{H}_2\text{O}$ are said to separate (Lefort, *C R* 69, 179). The crystals are deliquescent, at 50° they lose water, heated to 90° HCl is given off and O absorbed, by warming an aqueous solution of the crystals, FeCl_2 and FeCl_3 are formed.

Iron, cyanides of. No cyanides of iron have been isolated, but *ferric* and *ferro cyanides* are numerous, v vol ii pp 332, 334, 338

Iron, ferricyanide and ferrocyanide of, v vol ii pp 334 a 338

Iron, fluorides of. Two fluorides of Fe are known, FeF_2 and FeF_3 . The former is produced by dissolving Fe in HFAq, the latter by dissolving $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ in HFAq

FEROUS FLUORIDE FeF_2 . Small green crystals of this composition separate from a solution of Fe in HFAq (Scheurer Kestner, *A Ch* [3] 68, 472). By careful heating, FeF_2 is obtained as a white solid. FeF_2 is unchanged by heat, the hydrate is partly decomposed to HF and Fe_2O_3 . Slightly sol in water, more easily on addition of HF. The double salt $\text{FeF}_2 \cdot 2\text{KF}$ is a green pp, obtained by adding KF aq to FeSO_4 aq, the salts $\text{FeF}_2 \cdot \text{KF} \cdot 2\text{H}_2\text{O}$, $\text{FeF}_2 \cdot 2\text{NH}_4\text{F}$, and $\text{FeF}_2 \cdot \text{NH}_4\text{F} \cdot 2\text{H}_2\text{O}$ are described by Wagner (*B* 19, 896). The *Silicofluoride* $\text{FeF}_2 \cdot \text{SiF}_4$ ($= \text{FeSiF}_6$) is said to be formed as blue green crystals by dissolving Fe in H_2SiF_6 aq and crystallising (Wurtz, *Dictionnaire*, 1 1408)

FERRIC FLUORIDE FeF_3 . Crystals of the hydrate $2\text{FeF}_3 \cdot 9\text{H}_2\text{O}$ are obtained by dissolving $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ in HFAq and evaporating, or by oxidising FeF_2 in HFAq by HNO_3 . The crystals are yellowish according to Berzelius, colourless according to Scheurer Kestner (*A Ch* [3] 68, 472). At 100° $3\text{H}_2\text{O}$ is removed, at higher temperatures H_2O and HF escape. By heating the salt dried at 100° in a Pt crucible colourless crystals of FeF_3 are formed on the surface of the mass, isomorphous with AlF_3 (Deville, *C R* 43, 970). Slowly dissolved by water, addition of NH_4Aq ppt the *oxyfluoride* $3\text{Fe}_2\text{O}_3 \cdot 2\text{FeF}_3 \cdot 4\text{H}_2\text{O}$. FeF_3 forms double salts with K, Na, and NH_4 , these are obtained by adding FeF_3 aq to solutions of the alkali fluorides, the salts described are $\text{FeF}_3 \cdot 3\text{KF}$ (Berzelius, Wagner, *B* 19, 896), $2\text{FeF}_3 \cdot 4\text{KF} \cdot \text{H}_2\text{O}$ (Nicklès, *Z* 7, 480), $\text{FeF}_3 \cdot 3\text{NaF}$ (B, W, *l c*), $\text{FeF}_3 \cdot 2\text{NaF}$ (Nicklès, *J* 1869 268), $\text{FeF}_3 \cdot 2\text{NH}_4\text{F}$ (N, *l c*), to this salt Mangano gives the formula $\text{FeF}_3 \cdot 3\text{NH}_4\text{F}$ (*A Ch* [3] 60, 306)

Iron, haloid compounds of. Iron readily combines with the halogens. With each halogen, two compounds are formed, ferrous FeX and ferric FeX_3 , ferric iodide, however, has not been isolated with certainty, the compound Fe_2Cl_6 , corresponding to Fe_2O_3 , also perhaps exists. The only compounds that have been gasified are ferrous and ferric chloride. The former has the molecular formula FeCl_2 at 1300° – 1500° , but there are indications of the possible existence of molecules of Fe_2Cl_4 at lower temperatures, the latter appears to exist as a gas both as Fe_2Cl_6 and FeCl_2 . The haloid compounds of Fe are soluble in water, solutions of FeBr_3 and FeCl_3 are decomposed by much water, the compounds all form several double compounds with the chlorides &c of the more positive metals

Iron, hydrides of. No definite hydride of Fe has been certainly isolated, but there are several indications of the existence of a compound or compounds of Fe and H. Fe heated to redness absorbs about 46 vols H (Deville, *A* Troost, *C R* 57, 965, 59, 102). Electrolytically deposited Fe contains H along with N, CO, and CO_2 (Lenz, *P. Ergänzbd* 5, 252, cf Meidinger,

D P J 163, 288, Chandler Roberts, *C N* 31, 137, Iron was takes up H when immersed in dilute $\text{H}_2\text{SO}_4\text{Aq}$ (Johnson, *Pr* 23, 168, Reynolds, *C N* 29, 118) According to Caillaet (*C R* 80, 319) Fe npd electrolytically from neutral FeCl_2Aq containing NH_4Cl gives off 235-238 vols H when heated *in vacuo* Observations on the supposed existence of a gaseous hydride of Fe were made by Thomson, Dupasquier (*C R* 14, 511), and Reinsch (*J* 1858 '90) The non existence of such a gas was proved by Fresenius a Schlossberger (*A* 51, 415) and by Erlemeyer (*N J P* 9, 97) Thoma has investigated the conditions under which H is absorbed by Fe (*Z P C* 3, 91) He finds that when Fe is made the negative electrode in a voltameter, or when it is placed in dilute $\text{H}_2\text{SO}_4\text{Aq}$, it absorbs much H, but that Fe placed in an atmosphere of H does not absorb any of that gas, a certain portion of the H absorbed is given off again on exposure to air or immersion in water, the rest of the H is more firmly retained by the Fe According to Wanklyn a Carus (*A* 120, 69) when Fe_2 reacts with ZnEt , in presence of ether, various gases including C_2H_4 , CH_4 , C_2H_6 , and H are evolved, and a metal like powder remains, which, after washing with ether, is decomposed by H_2O giving H and a mixture of H and FeO They regard this powder as a hydride of Fe, perhaps FeH_2 , but no analyses are given

Iron, hydrated oxides or hydroxides of Compounds with water of each of the three oxides of iron are known The compositions of the various hydrated oxides are differently represented by different chemists they appear to vary with small changes of conditions, so that it is impossible to say whether all the bodies described are true hydrates or not The readiness with which most of the hydrated oxides undergo change of composition points to their being hydrates, i.e. compounds of water, rather than hydroxides, i.e. compounds of OH Thomson (*Th* 3, 293) gives the thermal data $-(\text{Fe}, \text{O}^2, 3\text{H O}) = 191,140$, $-(2\text{FeO H}^2, \text{O}, \text{H O}^2) = 54,590$, $[(\text{Fe}, \text{O}, \text{H O}) = 68,280$, $[(\text{FeO H}, \text{H SO}^2\text{Aq}) = 24,920$, $[(\text{Fe}^2\text{O}^2\text{H}^2, 3\text{H SO}^2\text{Aq}) = 33,840$

Only one hydrate of ferrous oxide, FeO H O , appears to exist, but it is doubtful whether any hydrate of this oxide has been obtained free from hydrates of Fe_2O_3 Descriptions are given of two hydrates of ferrous ferric oxide, viz $\text{Fe}_2\text{O}_3 \cdot \text{H}_2\text{O}$ and $\text{Fe}_2\text{O}_3 \cdot 4\text{H}_2\text{O}$, but here again there is great doubt as to the composition of the substances described as definite hydrates Many hydrates of ferric oxide have been obtained, the following are known as minerals ($\text{M} = \text{Fe}_2\text{O}_3$) $\text{M H}_2\text{O}$, $\text{M 2H}_2\text{O}$, $\text{M 3H}_2\text{O}$, $\text{2M H}_2\text{O}$, $\text{2M 3H}_2\text{O}$, $\text{3M 5H}_2\text{O}$, the following are said to have been obtained artificially $\text{M H}_2\text{O}$, $\text{M 2H}_2\text{O}$, $\text{M 3H}_2\text{O}$, $\text{2M H}_2\text{O}$, $\text{2M 3H}_2\text{O}$, many of these appear to exist in different modifications each distinguished by special properties, broadly they fall into two classes, those which are insoluble in water, and those which dissolve in water

HYDRATE OF FERROUS OXIDE $\text{FeO H}_2\text{O}$ The white pp obtained by adding air-free KOH Aq to an air free solution of a ferrous salt, and washing out of contact with air, is probably $\text{FeO H}_2\text{O}$ To obtain the pp nearly free from ferric oxide Schmidt (*A* 86, 101) pptts in a closed flask,

allows pp to settle, draws off water by a syphon, carefully pours warm water which has been boiled for some time on to the surface of the cold water in the flask, withdraws water by a syphon, &c., he then quickly transfers the moist pp by a syphon to a retort containing ether, and having attached to it a long glass tube dripping under Hg, the ether covers the pp and drives the air out of the retort, on warming, the water vapour and condensed water escape through the Hg When the ether is removed and the retort is cold, dry air free H (not CO, as the hydrate combines with this gas) is passed through the apparatus, and portions of the solid are transferred in an atmosphere of H to small wide mouthed stoppered tubes Dried in this way, the hydrate is a pale greenish, friable solid, which rapidly oxidises and glows in the air

A de Schulten (*C R* 109, 266) states that $\text{FeO H}_2\text{O}$ may be obtained in green six sided flat prisms, which almost instantly become red in air, by mixing 5 grams FeCl_2 dissolved in 100 c.c. air free water, and 200 c.c. NaOH Aq , containing 20 grams NaOH , in a flask filled with coal gas, heating, and allowing to stand for 24 hours

According to Bineau (*C R* 41, 509) ferrous hydrate is soluble in c. 150,000 parts water Ferrous hydrate is very easily oxidised, it is therefore an energetic reducer, e.g. it reduces salts of Au, Ag, and Pt, and also HIO_4Aq It dissolves in acids to form ferrous salts FeX_2 ,

$\text{X} = \text{NO}_3, \text{SO}_4, \text{PO}_4, \text{ \&c (cf Ferrous oxide,}$

under Iron, oxides of)

HYDRATES OF FERROUS FERRIC OXIDE

$\text{Fe}_2\text{O}_3 \cdot \text{H}_2\text{O}$ and $\text{Fe}_2\text{O}_3 \cdot 4\text{H}_2\text{O}$, a composition of both doubtful Hydrates of Fe_2O_3 are produced by ppg a mixture of equivalent quantities of a ferrous and ferric salt by alkali, and also by digesting ppg $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ with Fe The black powder obtained by Wöhler (*A* 28, 92), by ppg a mixture of equivalent parts of ferrous and ferric sulphates by slight excess of NH_4Aq , boiling, washing by decantation, and drying under 100° is said by Lefort (*C R* 69, 179) to be $\text{Fe}_2\text{O}_3 \cdot 4\text{H}_2\text{O}$ By ppg a mixture of equivalent quantities of a ferrous and ferric salt by hot conc KOH Aq , Lefort (*lc*) obtained a hydrate more stable than that got by using NH_4Aq , to this hydrate he gives the formula $\text{Fe}_2\text{O}_3 \cdot \text{H}_2\text{O}$ The black hydrate obtained by digesting $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ with Fe under water does not seem to have been analysed (v. Preuss, *A* 26, '96) Ferrous ferric hydrates are described as brown black, magnetic, brittle, powders, when heated out of contact with air they give Fe_2O_3 , heated in air, Fe_2O_3 is produced Ferrous ferric hydrates dissolve in acids to produce mixtures of ferrous and ferric salts, in some cases definite salts of the form Fe_2X_3 , $\text{X} = \text{SO}_4$, &c., are formed, according to Lefort (*C R* 69, 179) (cf Ferrous-ferric oxide, under Iron, oxides of)

HYDRATES OF FERRIC OXIDE A great many supposed hydrates of Fe_2O_3 have been described, but there is much doubt as to the composition and properties of definite hydrates of the form $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ The following occur as minerals, the compositions of which more or less accurately agree with the formulae—limonite $\text{Fe}_2\text{O}_3 \cdot 2\text{H}_2\text{O}$, $\text{Fe}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$, and $2\text{Fe}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$;

göthite $\text{Fe}_2\text{O}_3 \cdot \text{H}_2\text{O}$, *turgite* $2\text{Fe}_2\text{O}_3 \cdot \text{H}_2\text{O}$ and $3\text{Fe}_2\text{O}_3 \cdot 5\text{H}_2\text{O}$. According to Ramsay (*C J* 32, 895), and Bresscius (*J pr* [2] 3, 272), the pp obtained by adding alkali to a ferric salt, either hot or cold, washing, and drying at 100° , is $\text{Fe}_2\text{O}_3 \cdot \text{H}_2\text{O}$, by ppg FeCl_3Aq with NH_4Aq , washing first with water and then with alcohol and ether, and drying at 100° the hydrate $\text{Fe}_2\text{O}_3 \cdot 2\text{H}_2\text{O}$ is obtained, according to Bresscius (*lc*) Wittstein (*C C* 1853 367) says that the hydrate obtained by ppg a ferric salt by NH_4Aq , washing with cold water, and drying at a gentle heat, or at 100° for a short time, is $\text{Fe}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$. The experiments of Carnelley & Walker (*C J* 53, 89) on the dehydration of ppd $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ through a range of temperature showed that the pp after drying at 15° for 18 days contained more water than is required by $\text{Fe}_2\text{O}_3 \cdot 5\text{H}_2\text{O}$, that when heated to 55° the residue had the composition $\text{Fe}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$, but that water was gradually and continuously given off from 15° upwards until Fe_2O_3 was formed at 500° , the composition, however, remained constant from 385° to 415° , and corresponded with the formula $10\text{Fe}_2\text{O}_3 \cdot \text{H}_2\text{O}$. Van Bemmelen (*R T C* 7, 106) has examined the composition of the colloidal pp by adding NH_4Aq to FeCl_3Aq , his experiments, which were very extended, showed the great readiness with which the ratio of Fe_2O_3 to H_2O varies, *e.g.* exposed to air for a year the composition became constant with the ratio $\text{Fe}_2\text{O}_3 \cdot 4$ or $4\text{H}_2\text{O}$, after 4 months in dry air the ratio was $\text{Fe}_2\text{O}_3 \cdot 1.6\text{H}_2\text{O}$, heated to 100° in dry air the ratio was $\text{Fe}_2\text{O}_3 \cdot 96\text{H}_2\text{O}$, the same sample kept for 6 years in a closed flask and then heated to 15° until constant gave $\text{Fe}_2\text{O}_3 \cdot \text{H}_2\text{O}$, at 100° $\text{Fe}_2\text{O}_3 \cdot 45\text{H}_2\text{O}$, between 100° and 300° the final state was $\text{Fe}_2\text{O}_3 \cdot 25\text{H}_2\text{O}$, and so on. Besides the papers referred to above on the composition of ppd. $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$, the following are of importance.—Lefort (*J pr* 54, 305), P de Saint Gilles (*J pr* 66, 137), Schaffner (*A* 51, 177), Muck (*Z* [2] 4, 41), Davies (*C J* [2] 4, 69), Tommasi (*B* 12, 1929, 2334).

The *monohydrate* $\text{Fe}_2\text{O}_3 \cdot \text{H}_2\text{O}$ is said to be obtained (1) by adding NaOH Aq and KClO Aq to boiling FeSO_4Aq , washing, and drying at 100° (Muck, *Z* [2] 4, 41), (2) by keeping ppd $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ in contact with boiling water for 6-8 hours (P de Saint-Gilles, *A Ch* [3] 46, 47), (3) by repeatedly melting NaOH in a cast iron vessel and washing (Brunck & Gräbe, *B* 13, 725), (4) by keeping ppd $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ under water for many years at the ordinary temperature (Schiff, *C C* 1860 1768). This hydrate also occurs native as *göthite* $\text{Fe}_2\text{O}_3 \cdot \text{H}_2\text{O}$ is described as a dark-red powder, slowly dissolved by dilute warm HCl Aq , $\text{H}_2\text{SO}_4\text{Aq}$, or HNO_3 , insol in cold conc acids. The monohydrate obtained by Brunck & Gräbe was in the form of small tabular, brownish-violet, hexagonal crystals, SG 2.91. Göthite forms rhombic crystals, SG 3.8 to 4.2. The monohydrate obtained by Péan de Saint-Gilles, by keeping $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ in contact with boiling water for 6-8 hours, is described as a brick-red powder, scarcely acted on by conc boiling HNO_3Aq , slowly dissolved by conc hot HCl Aq , it dissolves in dilute HNO_3Aq , HCl Aq , or acetic acid, forming a red liquid which appears clear by transmitted, but

turbid by reflected light, and is ppd by a very small quantity of an alkali salt, on addition of conc HNO_3Aq or HCl Aq , this solution gives a red pp which dissolves on adding water. This modification of $\text{Fe}_2\text{O}_3 \cdot \text{H}_2\text{O}$ does not form Prussian blue with K_2FeCy_6 and acetic acid.

The *dihydrate* $\text{Fe}_2\text{O}_3 \cdot 2\text{H}_2\text{O}$ is said to be obtained by ppg cold FeSO_4Aq by NaOH Aq and KClO Aq or $\text{H}_2\text{O Aq}$, washing, and drying at 100° (Weltzien, *A* 138, 126, Muck, *lc*). This hydrate also occurs native as *limonite*.

The *hydrates* $2\text{Fe}_2\text{O}_3 \cdot \text{H}_2\text{O}$ and $2\text{Fe}_2\text{O}_3 \cdot 8\text{H}_2\text{O}$ are said to be formed by the action of water on ppd $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$, the first by boiling for 100 to 1,000 hours, Davies (*C J* [2] 4, 66), the second by long continued action of water at the ordinary temperature (Wittstein, *Ar Ph* 74, 158), or, crystalline, by freezing $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ suspended in water (Lamberger, *C C* 1853 783). By heating any of the hydrates in sealed tubes Fe_2O_3 is eventually produced (Sénarmont, *C R* 32, 762).

Muck (*Z* 1868 41) thinks that the ferric hydrates obtained by oxidising ferrous hydrate or carbonate in air differ essentially in properties from the ferric hydrates obtained directly from ferric salts. Tommasi (*Bl* [2] 38, 152, T a Pellizzari, *Bl* [2] 37, 196) arranges the ferric hydrates in two classes the red hydrates obtained by ppg ferric salts by alkali, and the yellow hydrates obtained by oxidising hydrates of FeO or Fe_2O_3 , or by oxidising FeCO_3 .

Ferric hydrates give up O to readily oxidised compounds such as SO Aq , SnCl_2Aq , in contact with decaying organic bodies the hydrates part with O, but again take it up if exposed to air, they absorb gases *e.g.* NH_3 and CO_2 (v Reinhardt, *Fr* 7, 187). They dissolve in acids to form ferric salts FeX , $\text{X} = \text{NO}_3$, $\frac{\text{SO}_4}{2}$, $\frac{\text{PO}_4}{3}$ &c

These hydrates when freshly ppd also dissolve in FeCl_3Aq forming oxychlorides (*q v*) (*v Ferric oxide*, under *Iron*, *oxides of*, p 62).

SOLUBLE FERRIC HYDRATES. A modification of $\text{Fe}_2\text{O}_3 \cdot \text{H}_2\text{O}$ soluble in water was prepared by P de Saint Gilles in 1855 (*A Ch* [3] 46, 47). Ordinary ppd $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ is dissolved in acetic acid, the solution is heated to 100° for a long time in a closed vessel, the blood red colour of the liquid changes to brick red, the liquid appears opalescent in reflected light, and the taste is no longer metallic but merely that of acetic acid, on now adding conc HCl Aq or HNO_3Aq , or the merest trace of $\text{H}_2\text{SO}_4\text{Aq}$ or an alkali salt, the whole of the Fe is thrown down as a brown red curdy pp, which, when dried on a porous tile, appears as a brown, lustrous, varnish like solid, having the composition $\text{Fe}_2\text{O}_3 \cdot \text{H}_2\text{O}$. This modification of ferric hydrate, sometimes called *ferric metahydrate*, is insol in conc acids, but dissolves easily in water, forming a deep-yellow, opalescent, tasteless liquid, unchanged by K_2FeCy_6 , or KCNS . Traces of H_2SO_4 or alkali salts, and also conc HCl Aq or HNO_3Aq , ppt the hydrate from its aqueous solution. The same hydrate has been obtained by Soheuer & Kestner by prolonged heating an aqueous solution of basic ferric nitrate in a sealed tube at 190° (*C R* 48, 1160), also by Debray by heating dilute FeCl_3Aq to 100° (*C R* 68, 913, *cf. Ferric chloride*, *Reactions*,

No 4, p 55) Another soluble modification of ferric hydrate, sometimes called *dialysed* or *colloidal ferric hydrate*, is obtained by dialysing a solution of FeCl_3 containing $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ so long as the dialysate contains HCl (Graham, *Tr* 1861 183) The solution is prepared by digesting FeCl_3Aq with $\text{ppd Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$, or by adding $(\text{NH}_4)_2\text{CO}_3\text{Aq}$ to FeCl_3Aq so long as the pp dissolves on stirring, the solution should contain 4-5 p.c. of solid matter, and have 5-6 equivalents of ferric hydrate in it After dialysis a deep red liquid is left on the dialyser, this liquid gelatinises by exposure to the air for some days, or by traces of H_2SO_4 , alkalis, alkaline salts, but not by HCl , HNO_3 , or acetic acid, the jelly like mass thus obtained slowly becomes insol in water, and is then sol in dilute acids, it appears to be changed to the ordinary $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ (Graham *Le*, cf Shuttlesworth, *Ph* [3] 8, 545, Becquerel, *A* 126, 208)

Krecke (*J pr* [2] 3, 286) has examined the formation of soluble ferric hydrates from FeCl_3Aq Formation of the colloidal hydrate occurs in FeCl_3Aq containing $\frac{1}{10}$ to $\frac{1}{2}$ p.c. FeCl_3 at ordinary temperatures, solutions containing 8 to 32 p.c. must be heated to above 100° in closed tubes, on cooling the more conc solutions (if not heated too long and too highly) re formation of FeCl_3 occurs, but in solutions of less than 1 p.c. the base and acid remain uncombined on cooling If a solution containing less than 1 p.c. is boiled for some time formation of ferric metahydrate begins, and after a time the hydrate is ppd as an orange yellow powder A table showing the various products obtained by heating solutions of FeCl_3Aq of different concentrations to different temperatures is given under *Ferric chloride, Reactions*, No 4 (p 55)

Iron, iodides of Only one iodide of iron, FeI_3 , has been isolated with certainty

FERRIC IODIDE (Proto iodide of iron) FeI_3 , H F [FeI_3Aq] = 47.650 (*Th* 3, 294) Fe and I combine by rubbing together with or without water, Fe and I heated together form a crystal line grey mass which melts at 177° (Carius a Wanklyn, *A* 120, 69) According to De Luca FeI_3 is white, but goes green on addition of water (*C R* 55, 615) Green deliquescent crystals of $\text{FeI}_3 \cdot 5\text{H}_2\text{O}$, SG 2873, are obtained by digesting 1 part Fe filings with 3 4 parts I in presence of water, adding Fe filings, evaporating out of contact with air, and filtering FeI_3 , especially FeI_3Aq , rapidly absorbs O with formation of oxyiodide (*q v*) Addition of sugar syrup to FeI_3 renders the solution more stable $\text{H}_2\text{O}_2\text{Aq}$ decomposes FeI_3 with formation of $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ and separation of all the I FeI_3Aq dissolves I, addition of K_2CO_3 to a solution containing 3 FeI_3 , 21 forms KI and $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$, the solution may contain FeI_3

FERRIC IODIDE (FeI_3) This iodide has not been isolated, it is probably contained in a solution of FeI_3 to which I is added in the ratio FeI_3 , I. FeI_3Aq containing I is separated by heating to FeI_3 and free I Fleury (*J Ph* [5] 16, 529) says that the action of I on Fe in presence of water soon ceases if the temperature is kept down to 15° , on filtering, only Fe is obtained, if, however, the mixture is boiled the red colour of the liquid disappears and a considerable quantity of Fe_2O_3 is found in the re-

sidue A large excess of I is required to dissolve all the Fe, and a large excess of Fe is required to convert all the I into FeI_3 F supposes that FeI_3 is at first formed and is then decomposed by the heat to Fe_2O_3 and HI, and the HI then acts on the free Fe forming FeI_3 , Mohr observed that a very dilute solution of FeCl_3 (1 in 12,300) gives a blue colour on addition of starch and KI only after a considerable time (*A* 105, 53) Nicklés found that a solution of $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ in HIAq in presence of ether did not at once give a blue pp with K_3FeCy , (*A Ch* [4] 5, 161, cf Ordway, *Am. S* [2] 26, 197) A solution made by digesting 126 parts I with sufficient iron, filtering, and adding 63 parts I, gives an apple green pp on addition of 201 parts citric acid previously neutralised by alkali carbonate (Creuse, *Ph* [3] 3, 953) According to Carius a Wanklyn (*A* 120, 70) if Fe is heated with excess of I until a little I is vapourised, and the mass is allowed to cool in a covered crucible, I is suddenly evolved, and the residue consists of FeI_3

Iron, nitrides of A nitride of Fe, Fe_3N_2 , is found as a silver like lustrous deposit in the lavas of Lina (Silvestri, *P* 157, 165), heated in H it yields Fe and NH_3 Finely divided Fe, prepared by heating ferrous oxalate or reducing Fe_2O_3 by H, takes up about 2 p.c. N when heated in N (Geuther a Briegleb, *A* 133, 228, Rogstadus, *J pr* 86, 307, Remsen, *Am* 3, 134) By heating Fe in NH_3 the Fe increases in weight 12-13 p.c. and becomes brittle (Berthollet, *G* 4 30, 378) By electrolysis a mixture of NH_4Cl and a ferrous salt a lustrous deposit is obtained, supposed by Kramer to be a nitride of Fe with 15 p.c. N (*Ar Ph* [2] 106, 284), but said by Meidinger (*D P J* 163, 283) to be an FeNH_3 amalgam A nitride (or nitrides) of iron seems to be produced by heating FeCl_3 or FeCl_2 in NH_3 The product is a brittle, grey white, solid, heated alone it gives N and Fe, heated in H it gives NH_3 and Fe, it is slowly and partially decomposed by boiling water, sol. in acids with formation of H, ferrous and NH_3 salts, not acted on by Cl, Br, or I in presence of H O at ordinary temperatures, but when heated ferric and NH_3 salts are formed, and H, and probably also N, are evolved The formula Fe_3N_2 agrees with most of the analyses, the formula Fe_2N_3 was given by Stahlschmidt (*P* 125, 37, cf Rogstadus, *J pr* 86, 307, Fremy, *C R* 52, 321)

Iron, nitroprusside of, v vol. II p 340

Iron, nitrosulphides or nitrososulphides of. In 1858 Roussin obtained a salt by the reaction of NH_4HS and KNO_3 on FeCl_3Aq , which contained Fe, but in which Fe could not be detected by the ordinary tests (*A Ch* [3] 52, 285) Procznsky (*A* 125, 302) obtained the same compound by adding NH_4HS to a solution of a ferrous salt saturated with NO A similar salt was obtained by Pavel by using K_2S in place of NH_4HS (*B* 12, 1407) By treating these salts with alkali others were obtained more or less closely related to the original salts The compounds thus obtained were analysed and examined by Roussin (*l c*), Procznsky (*l c*), Rosenberg (*B* 3, 812, 12, 1715), Demel (*B* 12, 461), and Pavel (*B* 12, 1407, 1949, 15, 2600) Roussin's analyses of the compound formed by using

NH_4HS led to the formula $\text{Fe}_3\text{S}_2(\text{NO})_4$, to the same compound Proczinsky gave the formula $\text{Fe}_3\text{S}_2(\text{NO})_2 \cdot 2\text{H}_2\text{O}$, and Rosenberg assigned the composition $\text{Fe}_3\text{S}_2(\text{NO})_{10} \cdot 4\text{H}_2\text{O}$. In his second paper R. showed that the compound prepared by using K_2S contained K, this was confirmed by Pavel. All the observers admitted the presence of the NO group in these compounds.

Pavel (B 15, 2600) has given an account of the compounds of which Roussin's salt is the type, and has indicated what appear to be their relations to other compounds of iron. As the compounds contain the group NO in addition to S they are better called nitroso than nitrosulphides. Pavel regards the nitroso sulphides of Fe as salts of acids, the acidic radicals of which are composed of Fe, NO, and S, the salts may be formulated generally as derived from the acids $\text{H}_n\text{Fe}_x(\text{NO})_y\text{S}_z$. P. compares the *ferronitrososulphides* with the *ferronitrosocyanides* or *nitroprussides*, which are salts of $\text{HFe}(\text{NO})\text{Cy}$, (*v Nitroprussides*, vol. II p. 340). According to P. there are two series of ferronitrososulphides derived from the two acids $\text{HFe}(\text{NO})_2\text{S}_2$ (or $\text{H}_2\text{Fe}_2(\text{NO})_4\text{S}_4$) and $\text{HFe}(\text{NO})_2\text{S}$ (or $\text{H}_2\text{Fe}_2(\text{NO})_4\text{S}_2$). The former acid has been isolated, the latter is not as yet known in the free state. The two series may perhaps be called *ferrinitroso* and *ferronitrososulphides*. In the following account of the salts Pavel's formulæ are adopted —

Series I *Ferrinitrososulphides* $\text{MFe}(\text{NO})\text{S}_2$

Potassium ferrinitrososulphide $\text{KFe}(\text{NO})\text{S}_2$. A solution of KHS, made from 44 grams KOH in 400 c.c. water, is added to a boiling solution of 35 grams pure NaNO_2 in 400 c.c. water, the mixture is heated just to boiling, and a solution of 159 grams ferrous sulphate in 1200 c.c. water, to which a drop of dilute H_2SO_4 has been added, is allowed to flow into it with constant stirring, the liquid is kept warm on a water bath, and is shaken from time to time until a greenish red pp. (of Fe_2O_3 , FeO , and S) forms on the sides of the flask, when the liquid is rapidly filtered, a little dilute KOH aq. is added to it when cold, and it is allowed to stand for 48 hours. The salt which separates is dissolved in water at 70° , a little KOH aq. is added as the liquid cools, and the crystals thus obtained are recrystallised in the same way, after standing 48 hours; the crystals are then dried over H_2SO_4 *in vacuo* protected from the light. About 30 grams of the pure salt are thus obtained (Pavel, B 15, 2601). If more KHS is used than directed above double sulphides of Fe and K are obtained and no nitrososulphide.

$\text{KFe}(\text{NO})\text{S}_2$ crystallises in large, nearly black, lustrous, rhombic prisms, it is fairly stable in the air, decomposition begins at $c 115^\circ$ with evolution of NO, at $c 130^\circ\text{S}$, $(\text{NH}_4)_2\text{SO}_4$, NH_4NO_3 , &c. sublimes, heated in air H_2O , NO, N, SO_2 are evolved, and the residue consists of FeS , Fe_2O_3 , Fe_3O_4 , and K_2SO_4 , strong mineral acids evolve H_2S . Various metallic salts form metallic sulphides and NO, while Fe salts remain in solution, heated with alkali, crystalline Fe_2O_3 is ppd, NO evolved, and ferronitrososulphide $\text{KFe}(\text{NO})\text{S}$ is formed (Pavel, l.c.). $\text{KFe}(\text{NO})\text{S}_2$ is soluble in about 2 parts hot water, slightly sol. in cold water, very sol. with decomposition (Pavel, B.

12, 1410) in alcohol or ether, the salt is ppd by NH_4Aq or KOH aq . The salts of this series do not react with K_2FeCy_6 .

The following salts of this series have been isolated NH_4X , BaX , CsX (the most stable salt of the series), CaX_2 , FeX_2 , LiX , MgX_2 , RbX , TiX ($\text{X} = \text{Fe}(\text{NO})\text{S}_2$). The Na salt is formed similarly to the K salt, also by the reaction of Na_2CS_2 with NaNO_2 , the compound $\text{Fe}_3\text{S}_2(\text{NO})_4\text{CS}_2$ said by Low (C. C. 1865 548) to be formed in this reaction does not exist according to Pavel (B 15, 2603). For other reactions of the salts of the series *v* Pavel, B 15, 2604.

Ferrinitrososulphydic acid $\text{HFe}(\text{NO})_2\text{S}_2$ seems to be obtained by adding a slight excess of dilute H_2SO_4 aq. to a dilute solution of the Na salt, it is insol. in water, alcohol, and ether, sol. in CHCl_3 or CS. The acid has not been obtained pure (P, l.c.).

Series II *Ferronitrososulphides*
 $\text{MFe}(\text{NO})\text{S}$. These salts are formed by the action of dilute alkali solutions on the salts of Series I, in these reactions NO is evolved and FeO ppd, but NH_3 is not evolved except from the salt $\text{NH}_4\text{Fe}(\text{NO})\text{S}_2$. The salts of Series II are very unstable, they easily pass into Series I, the Cs salt is the most unstable of all, these salts are insol. in ether (except the Fe salt), CS_2 , and CHCl_3 , they decompose in the air with separation of S and Fe_2O_3 , and evolution of a little H_2S , the solutions then have an alkaline reaction, and contain salts of Series I along with alkaline nitrite and thiosulphate. If CO is passed into a solution of K_2FeCy_6 , and the K salt of Series II is then added, NO is evolved, S and Prussian blue are ppd., and the filtrate contains K_2FeCy_6 , and some KNO.

Potassium ferronitrososulphide
 $\text{KFe}(\text{NO})_2\text{S} \cdot 2\text{H}_2\text{O}$ is obtained by warming $\text{KFe}(\text{NO})_2\text{S}_2$ on the water bath with dilute KOH aq., washing, and drying *in vacuo* over H_2SO_4 and CaO in the dark, it is very unstable, and can be obtained approximately pure only with difficulty (P, B 15, 2606). By warming an alcoholic solution of $\text{KFe}(\text{NO})_2\text{S}$ with a considerable excess of EtI, quickly distilling, washing the residue first with water and then with aqueous alcohol, and crystallising from benzene, *ethyl ferronitrososulphide* $(\text{C}_2\text{H}_5)_2\text{Fe}(\text{NO})_2\text{S}$ is obtained in black, lustrous, monoclinic crystals (P, l.c.). For a discussion of the probable constitution of these nitrososulphides and a comparison of them with the nitroprussides *v* Pavel, B 15, 2613.

Iron, oxides. Three oxides of iron have been isolated, FeO , Fe_2O_3 , and Fe_3O_4 . The molecular weight of none is known with certainty, as none has been gasified. FeO and Fe_2O_3 are basic oxides, they produce salts by reactions with acids, of the forms FeX_2 and FeX_3 , $\text{X} = \text{Cl}$, NO_2 , $\frac{\text{SO}_4}{2}$, $\frac{\text{PO}_4}{3}$, &c., Fe_2O_3 also combines with some

more positive metallic oxides to form compounds $\text{Fe}_2\text{O}_3 \cdot x\text{M}^{II}\text{O}$, which are usually called *ferrites* (*v* vol. II p. 547). Fe_2O_3 reacts with acids to form both ferrous and ferric salts, FeX_2 and FeX_3 , according to Lefort a few salts corresponding with Fe_2O_3 are known, e.g. Fe_2Cl_3 , $\text{Fe}_2(\text{SO}_4)_3 \cdot 2\text{SO}_4 \cdot 15\text{H}_2\text{O}$ (*v Ferric ferric oxide*, p. 61). Besides the ferrites a series of *ferrates* is known (*v* vol. II p. 546), these salts are of

the form M^xFeO_x , they may be regarded as salts of the hypothetical ferrie acid H_xFeO_x , the anhydride of this acid would be Fe_2O_3 , this oxide of iron has not been isolated, but the corresponding oxide of Mn, MnO_2 , is known. All the oxides of Fe form hydrates (*v Iron, hydrated oxides of*, p 57) FeO very readily and rapidly absorbs O and changes to Fe_2O_3 , at a very high temperature Fe_2O_3 parts with O, and is reduced to Fe_3O_4 .

FEROUS OXIDE (*Protoxide of iron*) FeO This oxide cannot be obtained free from Fe_2O_3 , by pptn from ferrous salts and dehydrating the ppt.

Preparation—1 Pure Fe_2O_3 is heated to c 300° in a stream of pure H (Wackenroder a Stromeyer, *Ar Ph* 36, 27)—2 By heating ferrous oxalate out of contact with air (Liebig, *A* 95, 116, Moissan, *A Ch* [5] 21, 199)—3 By heating FeO for 20 minutes to c 500° in H or CO, or in CO at 440° for 6 hours (Moissan, *lc*) 4 By heating Fe in CO_2 to c 1000° (Moissan, *lc*)—5 By adding ferrous oxalate to boiling KOHAq (Böttger, *J pr* 76, 239)

Properties and Reactions—An ivory black powder. According to Moissan (*lc*) FeO obtained by heating ferrous oxalate, or reducing Fe_2O_3 in CO at 500° , is pyrophoric, burns to Fe_2O_3 , and decomposes water at ordinary temperatures, while the FeO obtained by heating Fe in CO_2 at c 1000° is non pyrophoric, burns to Fe_2O_3 , and does not decompose water at 100° . Siewert (*J* 1864 266) says that pyrophoric FeO , produced by reduction of Fe_2O_3 in H, becomes non pyrophoric by remaining in an atmosphere of H for 12 hours after cooling. FeO is readily oxidised to Fe_2O_3 (*cf* Moissan, *supra*). It dissolves in acids to form ferrous salts FeX_2 , $X = Cl, NO_3, \frac{SO_4}{2}$, &c

Combinations—The compound $FeO \cdot Al_2O_3$ occurs native as *hercynite*, and $FeO \cdot Cr_2O_3$ as *chrome ironstone*, magnetic oxide of iron may be regarded as $FeO \cdot Fe_2O_3$. The hydrate $FeO \cdot H_2O$ has been isolated (*v Hydrate of ferrous oxide*, p 57)

FEROUS FERRIC OXIDE (*Black or magnetic oxide of iron*) Fe_3O_4 . This oxide occurs native in large quantities as *magnetite*. According to Spring (*Bl* [2] 50, 215) it is produced on the surface of iron which has been subjected to great pressure, the formation of a film of this oxide on iron rails is the reason why the surfaces of the rails of railroads do not rust (Spring, *lc*) When iron is strongly heated in air a film (*hammer scale*) is produced consisting of Fe_2O_3 , mixed with, or combined with, Fe_3O_4 , the outer layer of this film contains a large quantity of Fe_2O_3 , the inner layer, which is blackish grey and magnetic, is approximately $6FeO \cdot Fe_2O_3$ (Mosander, *P* 6, 35, *cf* Berthier, *A Ch* [2] 27, 19, Beaujeu a Mène, *C R* 61, 1135, Maumené, *Bl* 16, 25, Völcker, *W A B* 66, 193, Glasson, *A* 62, 89) *SG* 5 453 (Playfair a Joule, *C S Mem* 3, 81), 5 3 at 165° (Herapath, *P M* 64, 321) *SG* of *magnetite* 5 1 to 5 18 (Kopp, Rammelsberg) According to Moissan (*A Ch* [5] 21, 223) two varieties of Fe_3O_4 exist, one having *SG* 4 86 obtained by reducing Fe_2O_3 at 500° , the other having *SG* 5 18 obtained by reducing Fe_2O_3 at a higher

temperature Berthelot (*A Ch* [5] 23, 118) describes two varieties, one with *SG* 4 86 obtained by heating $FeCO_3$ to 350° , the other with *SG* 5 5 9 obtained by highly heating the first in N. *Magnetite* crystallises in regular octahedra, isomorphous with Mn_2O_3 (Rammelsberg) *C E*. 000029 (Kopp) *S H* $24^\circ-99^\circ$ 16779 (Regnault, *A Ch* [3] 1, 129) *H F* [Fe^+O^+]=26,900, [FeO, Fe^+O^+]=8,800 (Berthelot, *A Ch* [5] 23, 118)

Preparation—1 By heating $FeCO_3$ to c 350° in a current of CO , (Berthelot, *A Ch* [5] 23 118)—2 By heating $FeCl_2 \cdot 2NH_4Cl$ in air (Hauer, *W A B* 13, 456)—3 By heating FeO in a gentle stream of HCl (Deville, *C R* 53, 199)—4 By melting ferric phosphate with 3.4 pts. Na_2SO_4 (Debray, *C R* 52, 985)—5 By melting $CaCl_2$ with ferrous sulphate in a closed crucible (Kuhlmann, *C R* 52, 1283)—6 By the action of FeF_3 on boric acid at a high temperature in absence of air (Deville a Caron, *A* 108, 56)—7 By heating $FeCl_2$ with excess of Na_2CO_3 to low redness and washing with water (Liebig a Wohler, *P* 21, 582)—8 By heating Fe in steam 9 By long continued heating Fe_2O_3 to white heat (Sidot, *C R* 69, 201)—10 By reducing Fe_2O_3 in CO at 350° for about an hour (Moissan, *A Ch* [5] 21, 223, *cf* Birnie, *R T C* 2, 273, Ackermann a Sarnstrom, *B* 16, 783)—11 By dehydrating $Fe_2O_3 \cdot xH_2O$ at as low a temperature as possible (*v Hydrates of ferrous ferric oxide*, p 57)

Properties and Reactions—A black, magnetic powder, the crystals obtained by methods 4-7 (*supra*) are black octahedra. According to Moissan (*A Ch* [5] 21, 223) and Berthelot (*A Ch* [5] 23, 118), Fe_3O_4 obtained by heating $FeCO_3$ in CO , at 350° , or by reducing Fe_2O_3 by CO at 350° , or by heating reduced Fe in H saturated with H_2O , or in CO_2 at 440° , has a lower *SG* than Fe_2O_3 , produced by reductions at higher temperatures, or by strongly heating the first variety in N, the first variety is said to be easily soluble in conc HNO_3 Aq, and to give Fe_2O_3 when heated on Pt foil, the second variety is described as almost unacted on by boiling conc HNO_3 Aq and as unchanged to Fe_2O_3 by heat. Fe_3O_4 is reduced to Fe by heating with C, or by strongly heating in H. With a little HCl Aq it gives $FeCl_2$ Aq and Fe_2O_3 , with more HCl Aq it gives a solution showing the reactions of both ferrous and ferric chloride. According to Lefort (*C R* 69, 179) Fe_2Cl_6 is obtained by cooling a solution of $Fe_2O_3 \cdot xH_2O$ in excess of conc HCl Aq and evaporating *in vacuo* over H_2SO_4 . (*v Ferrous ferric chloride*, p 56) Lefort (*lc*) also describes a ferrous ferric sulphate— $Fe_2O_3 \cdot 6SO_3 \cdot 15H_2O$ —obtained by evaporating over H_2SO_4 solutions of Fe_2O_3 in H_2SO_4 , he says that arsenates and phosphates of Fe_2O_3 also exist, the solutions of these salts are decomposed to a mixture of ferrous and ferric salts by warming to $60^\circ-70^\circ$ (for hydrates of Fe_2O_3 , *v Hydrates of ferrous-ferric oxide*, p 57)

FERRIC OXIDE (*Peroxiide, sesquioxide, or brown oxide, of iron*) Fe_2O_3 . Occurs native in large quantities widely distributed as *red hematite*, *specular iron ore*, and *martite* *SG* native 5 2 to 5 28 (Rammelsberg, Kopp), *calciné*, 5 135 (Playfair a Joule, *C S Mem* 3, 80), *ppd*, 5 959 at 165° (Herapath, *P M* 64, 321) *S.H.*

15°-98° 16695 (Regnault, *A Ch* [8] 1, 129) CE 00004 (Kopp) Crystallises in hexagonal forms, $a = 1.186537$ (Rammelsberg)

Preparation—1. By ppg a ferro salt by NH_4Aq , thoroughly washing, and strongly heating the pp—2 By strongly heating ferrous oxalate in air, Vogel (*J pr* 68, 187) says this method gives a very pure product—3 By heating ferrous carbonate in air—4 By strongly heating ferrous sulphate in air, the product seems always to contain a little basic sulphate unless the temperature is kept very high for a long time

The oxide is obtained in crystals by the following methods—5 By heating a mixture of equal parts ferrous sulphate and NaCl , and washing with water, the temperature must not be too high, else some Fe_2O_3 is produced—6 By heating Fe_2O_3 in a slow stream of HCl gas (Deville, *C R* 52, 1364), in a rapid current of HCl , FeCl_3 is produced—7 By melting Fe_2O_3 with CaCl_2 (Kuhlmann, *C R* 52, 1283)—8 By the action of FeCl_3 vapour on strongly-heated lime (Daubré, *C R* 49, 143)—9 By melting Fe_2O_3 with borax and treating the fused mass with warm dilute HClAq (Hauer, *W. A B* 13, 456)

Properties—A reddish-brown, very hard, non magnetic powder, extremely hygroscopic, slightly volatile at c 3000° (Elsner, *J pr* 99, 257) According to Malaguti a magnetic variety of Fe_2O_3 exists (*A Ch* [3] 69, 214, cf Lallemand, *A Ch* [3] 69, 233, Lawrence Smith, *B* 8, 183)

Reactions—1 Heated to a very high temperature Fe_2O_3 is partially deoxidised to Fe_3O_4 (Sidot, *C R* 69, 201)—2 Reduced by heating in hydrogen, according to Moissan (*C R* 74, 1296) Fe_2O_3 (prepared from ferrous oxalate) is reduced to Fe_3O_4 by heating to 350° for an hour in H_2 , reduced to FeO by heating to 500° for 20 minutes in H_2 , and to Fe when heated in H_2 at 700° (cf Siewert, *J* 1864 265) Wright a Luff (*C J* 83, 1) found that the temperature at which reduction of Fe_2O_3 by H begins varies from 195° to 260° according to the physical state of the Fe_2O_3 used—3 Reduced by heating in carbon monoxide, to Fe_3O_4 at 350°, to FeO at 500°, and to Fe at 700°-800° (*M, lc*, cf Gruner, *C R* 73, 28) Temperature at which reduction begins varies from 90° to 220° according to physical state of Fe_2O_3 used (*W a L, lc*)—4 Reduced by heating with carbon to Fe_3O_4 , and then to Fe which combines with part of the C (*v Iron, carbides of*, p 53) Reduction begins at 430°-450° (*W a L, lc*) According to Parry (*C N* 27, 313) reduction of Fe_2O_3 by C in a vacuum begins at above 400°, at 1200° c $\frac{2}{3}$ of the oxide is reduced.—5 Reduced to FeO by heating in a mixture of equal volumes of carbon monoxide and dioxide (Debray, *C R* 45, 1018)—6 A mixture of carbon dioxide and cyanogen (6 vols 1 vol) partially reduces Fe_2O_3 to Fe at c 600°-700° (Bell, *J* 1874 266)—7 Heated in ammonia Fe_2N_3 is produced (*v Iron, nitrides of*, p 59) 8 Chlorine forms FeCl_3 slowly at full red heat (Weber, *P* 112, 619)—9 Heated with sulphur SO_2 and FeS are produced (Brescius, *D P J* 192, 125)—10 Heated strongly in a rapid stream of hydrogen chloride FeCl_3 and H_2O are formed (Deville, *C R* 52, 1264)—11 Sulphurated hydrogen does not act on dry Fe_2O_3 ; with

moist Fe_2O_3 ($\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$) FeS , S , and H_2O are formed (Wright, *C J* 43, 156, Brescius, *D P J* 192, 125)—12 Phosphor *c chloride* produces FeCl_3 , which then combines with part of the PCl_3 , forming FeCl_2 , PCl_3 (Weber, *J pr* 76, 410)—13 Fe_2O_3 is slowly dissolved by acids, the more slowly the denser the specimen of Fe_2O_3 , most easily dissolved by 16 times its weight of a boiling mixture of 5 pts H_2SO_4 and 3 pts H_2O (Mitscherlich, *J pr* 8^o 103) After strongly heating with alkalis, alkaline carbonates, or acid sulphates, Fe_2O_3 is readily dissolved by acids—14 Fe_2O_3 readily parts with O when heated with oxidisable organic compounds, on exposure to air O is again taken up (Robin, *C R* 49, 500, Grager, *A* 111, 124) Moist Fe_2O_3 ($\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$) in presence of ordinary air serves as an oxidiser of various vegetable matters, e.g wood (Kuhlmann, *D P J* 155, 81, cf P Thénard, *C R* 49, 289)

Combinations—1 Fe_2O_3 is extremely hygroscopic, it forms several hydrates by combining with water For conditions of formation, composition, and properties *v Hydrates of ferric oxide*, p 57—2 With some more ositive metallic oxides to form ferrites (*q v* in vol II p 547)

HYPOTHETICAL FERRIC ANHYDRIDE (Fe_2O_3) This oxide has not been isolated, but a number of salts are known, which are best regarded as derived from the hypothetical acid H_2FeO_4 , of which Fe_2O_3 would be the anhydride These salts are described under the heading FERRATES (vol II p 546) In addition to the descriptions there given it is to be noted that when air is blown into hot conc NaOHAq containing c 84 p.c NaOH , a perceptible quantity of Fe is dissolved to form a colourless liquid, from which $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ slowly separates (Zirmité, *Chem Zeitung*, 12, 355), possibly the solution contains a ferrate of Na , or, according to the suggestion of Z, a perferate NaFeO_4 (?)

Iron, oxybromides of Oxybromides of Fe are said to be formed by evaporating FeBr_3Aq , and by the action of $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ on FeBr_3Aq or HBrAq (Ordway, *J pr* 76, 19, Bechamp, *A Ch* [3] 57, 296, Scheufelen, *A* 231, 177)

Iron, oxychlorides of The oxychlorides of Fe are numerous and of complex composition, they belong to the form $x\text{FeCl}_3 \cdot y\text{Fe}_2\text{O}_3 \cdot z\text{H}_2\text{O}$ They are divided into two classes, those which are soluble in water, and those which are insoluble

Soluble oxychlorides Prepared by digesting freshly pptd $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ in FeCl_3Aq , also by digesting excess of $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ in HClAq By the former method Ordway obtained $2\text{FeCl}_3 \cdot 28\text{Fe}_2\text{O}_3$ (*J pr* 76, 19) By the second method Béchamp (*A Ch* [3] 51, 296) obtained liquids containing FeCl_3 and Fe_2O_3 in ratio 2 5, 1 3, 1 4, and 1 5 All these solutions evaporated at c 40° give residues soluble in water, solutions containing more Fe_2O_3 than the foregoing (up to $10\text{Fe}_2\text{O}_3$) give residues on evaporation which are insol in water Solutions of soluble oxychlorides are not decomposed by dilution or heating, but addition of various acids and salts causes pptn. of $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$, mixed with oxychlorides, which are soluble in water From solutions of soluble oxychlorides NH_4Aq ppts $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ free from chloride, whereas the pp obtained from the insol oxychlorides contains chloride

Insoluble oxychlorides Prepared by oxidation of FeCl_2Aq , by exposure to air or by HNO_3 in presence of a little HCl (Béchamp, *l.c.*) 10 saturated FeCl_2Aq , HCl is added in less proportion than 3HCl to FeCl_2 , the solution is heated to 100° , and HNO_3Aq is gradually added, a violent reaction occurs, the nearly black liquid becomes yellow, and oxychlorides separate which are insol in water. The composition of these oxychlorides varies according to the temperature and the proportion of HCl used. Insol oxychlorides are also formed by adding alkali to FeCl_2Aq in quantity insufficient for complete pptn. These oxychlorides vary in colour from yellow to brown, they are slowly soluble in HClAq , when heated they lose H_2O and HCl .

Iron, oxyfluoride of By adding NH_4Aq to a solution of FeF_3 , a yellow salt of the composition $2\text{FeF}_3 \cdot 3\text{Fe}_2\text{O}_3 \cdot 4\text{H}_2\text{O}$ is obtained (Deville, *Ch* [3] 49, 85).

Iron, oxysulphide of According to Rammeisberg (*P* 121, 339) the product obtained by heating Fe_2O_3 in H_2S gas to near redness is $8\text{Fe}_2\text{S}_3 \cdot \text{Fe}_2\text{O}_3$.

Iron, reduced *Ferrum reductum*. Very finely divided Fe is obtained by reducing Fe_2O_3 in a stream of pure H . Reduced iron is very readily oxidised, commercial specimens generally contain 40 p.c. or more of oxide.

Iron, phosphides of Meyer in 1780, and Bergmann somewhat later, obtained a phosphide of Fe , supposing it to be a new metal they called it *Hydrosiderium* (*Berl Ges d naturforsch Freunde* [1781] 2, 334, Bergmann's *Opusc* 3, 109). Several Fe phosphides have been described. According to Lucas (*P* 132, 225) only three exist, Fe_3P , FeP , and Fe_2P . They are non magnetic and almost infusible, unchanged by cold HClAq , slowly acted on by boiling HClAq giving $\frac{1}{2}$ of their P as phosphoric acid and $\frac{1}{4}$ as PH_3 , oxidised at ordinary temperature by HNO_3 , and *aqua regia*.

Fe_2P . By heating FeS in PH_3 gas (H Rose, *P* 6, 212, 24, 333), also by heating reduced Fe or FeCl_2 in PH_3 (Freese, *l.c.*) A blue grey powder, $\text{SG } 5.04$, loses P by heating in H or CO , probably forming FeP .

FeP . By heating FeS in PH_3 for a long time (Freese, *l.c.*), also by passing a mixture of H and P vapour over strongly heated Fe_2O_3 , and by the action of PH_3 on FeCl_2 (H Rose, *l.c.*) A black crystalline mass, not acted on by HNO_3 , or HClAq , nor by I .

Fe_3P . 7 parts dry FePO_4 with 1 part lamp black are covered with a layer of NaCl and heated to whiteness, and the fused mass is treated with HClAq , FeP remains as a grey crystalline solid, $\text{SG } 5.74$, unchanged by heating in H , CO , or CO_2 (Freese, *l.c.*)

The following phosphides have been described in addition to those mentioned, according to Freese they are not definite compounds— Fe_2P (Hvoslef, *A* 100, 99), Fe_2P_2 (Schenk, *O J* [2] 11, 826), Fe_2P_2 (Struve, *J pr* 79, 321), Fe_2P (Sidot, *O R* 74, 1425).

Iron, salts of Two series of Fe salts, obtained by replacing H of acids by Fe , are known, *ferrous* salts FeX , and *ferric* salts FeX_3 ($\text{X} = \text{Cl}$, &c., NO_2 , &c., $\frac{\text{SO}_3}{2}$, &c., $\frac{\text{PO}_4}{3}$, &c.). The ferrous

salts are the more stable, as a class ferrous salts are readily oxidised to FeX_3 . Both series contain many well marked and very definite compounds. Lefort (*C R* 69, 179) asserts the existence of a few salts, e.g. chloride, sulphate, phosphate, corresponding to Fe_2O_3 (*vs Ferrosio-ferric oxide*, p 61). Ferrous salts are usually obtained by dissolving Fe in various acids, or in the cases of the insol salts by double decomposition from FeSO_4 , they are for the most part sol in water, a few basic ferrous salts are known, but the greater number are normal salts. Many double ferrous salts, especially with the alkali sulphates, have been isolated, many ferrous salts are isomorphous with the corresponding salts of Co , Ni , Mn , Zn , and Mg . A double ferrous aluminium sulphate, $\text{FeSO}_4 \cdot \text{Al}_2\text{SO}_4 \cdot 24\text{H}_2\text{O}$, is said to exist and to be isomorphous with the alums. Ferric salts are generally prepared by dissolving $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ in the different acids, many basic, and numerous double, ferric salts are known, the double salt $(\text{NH}_4)_2\text{SO}_4 \cdot \text{Fe}(\text{SO}_4)_2 \cdot 24\text{H}_2\text{O}$ belongs to the class of alums. The ferric salts are generally analogous to the persalts of Al , Cr , Co , Ni , and Mn (*cf IRON GROUP OF ELEMENTS*). The following list comprises the more important salts of Fe derived from oxyacids, for details *cf* CARBONATES, NITRATES, SULPHATES, &c. *antimonates*, *arsenates* and *ites*, *borates*, *bromates*, *carbonates*, *chlorates*, *chromates*, *hypophosphites*, *iodates*, *molybdates*, *mobates*, *nitrates*, *perchlorates*, *periodates*, *phosphates* and *ites*, *selenates* and *ites*, *silicates*, *sulphates* and *ites*, *tantalates*, *tellurates* and *ites*, *thiosulphates*, *titانات*, *tungstates*, *vanadates*.

Iron, selenides of Selenides of Fe seem to be obtained (1) by passing vapour over strongly heated Fe , (2) by heating the product obtained in (1) with Se (Berzelius), (3) by method (2) covering the mass with borax, Little (*A* 112, 211) obtained Fe_2Se_3 , thus, $\text{SG } 6.38$, (4) by ppg Fe salts by H_2Se (Reeb, *J Ph* [4] 9, 173). By heating Fe filings with Se to redness Diversa a Shimidzu (*C J* 47, 441) obtained FeSe , resembling FeS in appearance, with acids yields H_2Se .

Iron, selenocyanides of None has been isolated, Crookes (*C J* 4, 12) mentions various reactions which do not yield a definite salt.

Iron, silicide of It is doubtful whether any definite compound has been isolated. Silicides are apparently formed (1) by heating Fe with Si (Deville a Caron, *C R* 45, 163), (2) by the action of molten Fe on silicates. Hahn (*A* 129, 57) obtained a body approximately Fe_2Si by fusing 60 parts Na_2SiF_6 , 20 parts Na , 60 parts Zn , and 22 parts steel, under NaCl . Other bodies, approximately FeSi and Fe_2Si_3 , are described by Deville a Caron (*l.c.*) and Hahn (*l.c.*, *cf* Bous singault, *A Ch* [3] 16, 15).

Iron, silicofluoride of, FeSiF_6 , *vs Ferrous fluoride*, p 56.

Iron, sulphocyanides of, *vs* vol II p 350.

Iron, sulphides of Four are known, FeS , Fe_2S_3 , Fe_3S_4 , and FeS_2 , a subsulphide, Fe_2S , also probably exists. According to Gautier a Hallopeau (*C R* 108, 806) a yellow grey lustrous body, Fe_2S , is formed by heating Fe in CS_2 for several hours at 1300° – 1400° .

Iron subsulphide Fe_2S . Said by Arfwedson (*P* 1, 72) to be produced, as a greyish-black powder, by heating dry FeSO_4 in H .

Ferrous sulphide FeS (*Iron monosulphide* or *protosulphide*) Occurs sometimes in meteorites, also in combination with NiS

Formation—1 By heating FeS_2 with Fe until the mass is thoroughly molten—2 By reducing FeS_2 in H—3 By the reduction of Fe_2O_3 or ferric salts by decomposing organic matter in presence of sulphates (Chevreul, *C R* 43, 218) 4 By gently heating a mixture 1 part S with 1½ parts Fe filings moistened with water, if a considerable quantity of such a mixture is made into a paste with water and covered with earth, the mass after a time becomes red hot and evolves much steam which throws up the earth with some violence—5 By strongly heating Fe_2O_3 or a ferric salt in H_2S (Sidot, *C R* 66, 1257, cf Carnot, *Bl* [2] 32, 162)—6 By heating a mixture of Fe_2O_3 and S in H (Rose, *P* 110, 120)—7 By heating to redness a mixture of Fe_2O_3 and excess of $\text{Na}_2\text{S}_2\text{O}_3$ (Gibbs, *Am S* [2] 37, 346)

Preparation—1 A mixture of 8 parts Fe filings and 2 parts S is strongly heated in a covered crucible until the whole mass is thoroughly melted, if the temperature is not high enough some Fe remains and some Fe_2S_3 is formed—2 A white hot rod of iron is plunged into molten S, the FeS formed flows off, the operation is repeated as long as any S remains (Gahn, Rammelsberg, *B B* 1862 681)—3 By ppg a ferrous salt by an alkali sulphide, preferably NH_4 sulphide

Properties— FeS prepared in the dry way is yellow-brown, lustrous, metal-like solid, crystallising in hexagonal prisms, *SG* 4.69 Non magnetic, not changed by heating out of contact with air or in H FeS prepared in the wet way is an amorphous, greenish black powder, which on gently heating in air is partly changed to FeSO_4 , it is slightly soluble in water By ppg FeS in a very dilute solution, and then dialysing away the other products of the reaction, Wimsinger (*Bl* [2] 49, 452) obtained a very dilute solution of colloidal FeS , the solution was greenish brown, oxidised and coagulated very readily

Reactions— FeS prepared in dry way 1 Heated in hydrogen, or out of contact with air, is unchanged—2 Heated in steam evolves H and H_2S , and leaves a black magnetic mass containing Fe_2O_3 (Regnault, *A Ch* [3] 62, 379)—3 Heated strongly in air forms Fe_2O_3 and SO_2 —4 With dilute hydrochloric or sulphuric acid evolves H_2S , and forms FeCl_2 or FeSO_4 —5 Oxidised by nitric acid—6 Heated with chlorine from S_2Cl_2 and FeCl_3

FeS prepared in the wet way 7 Oxidises in air at ordinary temperature, forming $\text{Fe}_2\text{O}_3 \cdot x\text{H}_2\text{O}$, S, and a little basic Fe sulphate (Wagner, *D P J* 192, 131)—8 Slightly soluble in water, re-ppd by NH_4HS —9 Soluble in sulphurous acid solution, also in potassium cyanide solution (Goueron, *C R* 75, 1276)—10 Slightly soluble in solutions of alkaline sulphides—11 Very easily decomposed by acids, with evolution of H_2S

Ferroso ferric sulphide Fe_2S_3 (*Magnetic sulphide of iron*) Occurs native as *magnetic pyrites* in hexagonal crystals, *SG* 4.51 to 4.64, the composition may be expressed as $x\text{FeS} \cdot \text{Fe}_2\text{S}_3$, x is very seldom = 1, generally = 5 and 6 The first product of the action of H_2S on strongly

heated Fe_2O_3 is Fe_2S_3 , but this decomposes to FeS (Sidot, *C R* 66, 1257)

Ferric sulphide Fe_2S_3 (*Sesquisulphide of iron*) By heating a mixture of powdered FeS and S to redness, or by passing H_2S over Fe_2O_3 at 100° (Berzelius, *P* 7, 393) A greenish-yellow mass, magnetic according to Proust (*Scher J* 10, 54), non magnetic according to Berzelius (*lc*) When heated, gives Fe_2S_3 , when moist soon changes to a mixture of S and Fe_2O_3 , decomposed by dilute acids forming ferrous salts, H_2S , and $\text{H}_2\text{S} \cdot x\text{S}$

Combines with Fe_2O_3 , when heated with it. The product of the reaction of H S on Fe_2O_3 at somewhat above 100° is $\text{FeO} \cdot x\text{Fe}_2\text{S}_3$, according to Rammelsberg (*P* 121, 337) Combines with CuS to form $\text{CuS} \cdot \text{Fe}_2\text{S}_3$, also with K_2S , Na_2S , Ag_2S These compounds may be regarded as *sulphoferrites*, they belong to the form $\text{M}_2\text{Fe}_2\text{S}_3$, and are produced by fusing 1 part Fe filings with 5-6 parts S and 5-6 parts alkaline carbonate, the Ag salt is obtained by adding AgNO_3 to the K salt suspended in water (v Schneider, *P* 136, 460, Preis, *J pr* 107, 10) Fe_2S_3 is said to form a hydrate, $\text{Fe}_2\text{S}_3 \cdot 3\text{H}_2\text{O}$, a greenish black pp obtained by adding $\text{NH}_4\text{HS} \cdot \text{Aq}$ to a ferric solution containing an oxidiser such as Cl or a hypochlorite (Phipson, *C N* 30, 139)

Iron disulphide FeS_2 (*Iron pyrites*)

Occurrence—In large quantities, in regular crystals as *yellow pyrites*, and in rhombic crystals as *white pyrites*

Formation—1 By slowly heating a mixture of Fe_2O_3 , S, and NH_4Cl (Wöhler, *P* 37, 238)—2 By the action of CS_2 on Fe_2O_3 (Schlagdenhaufen, *J Ph* [3] 34, 175)—3 By heating Fe, or Fe_2O_3 , with $\text{SO}_2 \cdot \text{Aq}$ in a sealed tube to 200° (Geitner, *A* 129, 350)—4 By passing H_2S over an oxide or chloride of Fe at a temperature above 100° but below redness—5 By the action of organic matter on water containing Fe and sulphates in solution—6 Glatzel (*B* 33, 37) obtained crystals of FeS , by strongly heating FeCl_3 with P_2S_5

Preparation—1 An intimate mixture of 2 parts FeS and 1 part S, or of 1 part Fe with 1½ parts S, is heated in a retort to a temperature somewhat under red heat, and the product is treated with dilute $\text{HCl} \cdot \text{Aq}$ (Berzelius), below 100° the chief product is FeS_2 , and at red heat Fe_2S_3 is formed—2 By mixing an alkaline per sulphide solution with $\text{FeCl}_2 \cdot \text{Aq}$ at 180° or with $\text{FeSO}_4 \cdot \text{Aq}$ at 165° (Benarmont, *A Ch* [3] 30, 129)

Properties—A bulky dark yellow powder, or crystallised in small brass yellow cubes and octahedra Not magnetic The rhombic form of FeS_2 oxidises in moist air forming $\text{FeSO}_4 \cdot \text{S}$, and H_2SO_4

Reactions—1 White iron pyrites, finely-divided yellow pyrites, or FeS_2 prepared in the wet way, oxidise in air, forming chiefly FeSO_4 , and at higher temperatures SO_2 and Fe_2O_3 —2 Calcined with carbon, gives CS_2 —8 Not acted on by dilute acids, but decomposed by conc hydrochloric acid, giving H_2S and S—4 Oxidised by heating with conc nitric acid

Iron, telluride of Produced by reducing ferrous telluride in H (Berzelius)

M M P M

IRON ALUMS

K_2 (or $(\text{NH}_4)_2$) $[\text{SO}_4 \cdot \text{Fe}_2(\text{SO}_4)_2 \cdot 24\text{H}_2\text{O}]$, v ALUMS, vol I. p 148, and SULPHATES. M M P. M.

IRON GROUP OF ELEMENTS The four elements, IRON, NICKEL, COBALT, and MANGANESE, are more or less closely connected. Fe, Ni, and Co occur in meteorites, some of which also contain minute quantities of Mn. The chief ores of Fe and Mn are oxides, carbonates, and sulphides, of Ni and Co sulphides and arsenides. Fe has been known and used for ages, Ni, Co, and Mn have been known from about the middle of the eighteenth century. The name *iron* is probably a form of the Sanscrit word *ayas* (= metal), the names *nickel* and *cobalt* are derived from terms used by miners in the Middle Ages to express false or spurious metals, or minerals from which no useful metals could be obtained, *manganese* is said to be a corruption from *mag-*

nesia nigra, a name long given to the mineral pyrolusite. Compounds of Fe occur in large quantities, widely distributed, compounds of Mn are not so common, nor so widely distributed, and compounds of Ni and Co occur only sparingly. Fe, Ni, Co, and Mn are obtained by reducing the oxides with charcoal at high temperatures. The metals are all hard, lustrous, fairly malleable and ductile, they crystallise in the regular system, they are more or less magnetic, they are unacted on by dry air, but oxidise slowly in moist air, they decompose steam, and react with acids to form salts. The following table presents some of the characteristic properties of the four metals —

	MANGANESE	IRON	NICKEL	COBALT
Atomic weights	55	55.9	58.6	58.8
One compound of Mn ($MnCl_2$), and two of Fe ($FeCl_2$ and $FeCl_3$) have been gasified, no Ni or Co compounds have been gasified, specific heats of the four metals have been directly determined. Molecular weights unknown				
Melting points (approx)	1800°–1900°	1600°	1500°	1500°
Spec. gravs (approx)	8	7.8	8.9	8.6
Atom. weight.	6.9	7.2	6.6	6.8
Spec. grav (approx)				
Occurrence and Preparation.	Occurs chiefly as MnO_2 . Obtained by reducing the oxides by C at a high temperature, Also by reducing the chloride or fluoride by Na.	Chiefly as Fe_2O_3 , Fe_3O_4 , and $FeCO_3$. Obtained by reducing Fe_2O_3 by C, or CO, at a high temperature, also by reducing Fe_2O_3 or Fe_3O_4 in H, and by electrolysis of $FeSO_4$ in Aq.	Chiefly as $NiAs$ and $NiAsS$. Obtained by reducing Ni_2O_3 by C at a high temperature, or by heating in H.	Chiefly as $CoAs_2$ and $CoAsS$. Obtained by reducing Co_2O_3 by heating with C or in H.
Physical properties	Greyish-white, brittle, very hard, lustrous, magnetic. Melts above M.P. of Fe.	Lustrous, greyish-white, very tenacious, crystallises in regular cubes and octahedra, magnetic. Ordinary iron, containing from 2 to 5 p.c. C, is very hard, malleable, and fairly ductile. Fe obtained by reduction of Fe_2O_3 in H below 600° is pyrophoric. By electrolysis of $FeSO_4$ in Aq a soft, amorphous, brittle mass is obtained.	Silver white, very tenacious, hard, ductile, malleable, magnetic up to c. 250°, crystallises in regular cubes and octahedra.	Steel grey, lustrous, hard, very ductile at red heat and upwards, somewhat malleable, magnetic, crystallises in regular cubes and octahedra.
Chemical properties	Oxidises easily in air, Mn obtained by reducing chloride by Na is said not to tarnish in ordinary air. Oxidised by heating in air. Decomposes steam, said to decompose	Unchanged in dry air, in moist air forms $Fe_2O_3 \cdot xH_2O$, heated in air or O, burns to Fe_2O_3 , and Fe_3O_4 . Decomposes steam, forming Fe_2O_3 and H. Dissolves in dilute acids, almost in-	Unchanged in ordinary air, heated in air or O burns to NiO or Ni_2O_3 , according to the temperature. Decomposes steam at red heat. Dissolves in dilute acids, but is unacted on by	Unchanged in ordinary air; heated in air or O burns to Co_2O_3 . Decomposes steam at red heat, decomposes NH_3 to N and H. Dissolves in dilute acids, but is unacted on by

TABLE—continued

	MANGANESE	IRON	NICKEL	COBALT
	<p>cold water slowly. Dissolves in dilute acids, hardly acted on by cold conc H_2SO_4. Combines directly with Cl and Br. No simple cyanide of Mn is known, but the acid H_2MnCy_2 and salts derived from it, and also salts of the hypothetical acid H_2MnCy_2, have been isolated. With acids forms two series of salts, MnX_2 and MnX_3, of which the manganese salts ($X = Cl, SO_4, \&c$) are the more stable. Besides the oxides corresponding to the salts (MnO and Mn_2O_3) other acidic oxides, MnO_2 and Mn_2O_7, have been isolated, the acid $HMnO_4$, and numerous salts of this acid, are known, salts of the hypothetical acid H_2MnO_4 are also numerous. Atom of Mn is divalent in the gaseous molecule $MnCl_2$.</p>	<p>sol in cold conc H_2SO_4, unacted on by cold conc HNO_3. Combines directly with Cl, Br, and I, also with C, and probably with H and N. No simple cyanide is known, but a large number of ferro and ferric cyanides, and also the acids H_2FeCy_2 and H_2FeCy_3, have been isolated. Some ammonio compounds are known. Reacts with acids to form two series of salts, FeX_2 and FeX_3, of which the ferric salts ($X = Cl, SO_4, \&c$) are the more stable. Ferrates, M_2FeO_4, are also known, but neither the corresponding acid nor anhydride has been isolated. Atom of Fe appears to be divalent and trivalent in gaseous molecules ($FeCl_2$ and $FeCl_3$), perhaps also tetravalent (Fe_2Cl_4 and Fe_2Cl_6).</p>	<p>cold conc HNO_3. Combines directly with Cl, Br, and I. Probably forms a nitride by heating NiO in NH_3, N Cy_2 and many double cyanides are known, but no acids or salts corresponding with ferro, ferri, manganese, and manganese cyanides have been isolated. Some ammonio compounds are known. With acids forms two series of salts, NiX_2 and NiX_3, of which the nickelous salts ($X = Cl, SO_4, \&c$) are much the more stable. No salts in which Ni forms part of the acidic radicle have yet been certainly isolated.</p>	<p>cold conc HNO_3. Combines directly with Cl, Br, and I, also probably with C. $CoCy_2$ is known, also the acids H_2CoCy_2 and H_2CoCy_3, and many salts derived therefrom. Very many ammonio compounds are known. With acids form two series of salts, CoX_2 and CoX_3, of which the cobaltous salts ($X = Cl, SO_4, \&c$) are much the more stable. By the action of molten KOH on CoO a salt is obtained in which Co seems to form part of the negative radicle.</p>

General formula and character of compounds—Oxides— MO, M_2O_3, M_2O_5 , also MnO_2 , MnO_3 , Mn_2O_3 , Sulphides— MS, M_2S, M_2S_3, MS_2 . Haloid compounds— MX_2 or M_2X_3 , also FeX_2 or Fe_2X_3 , $?MnCl_2$, $?MnCl_3$. Salts— MX_2 and MX_3 , $X = NO_2, \frac{SO_4}{2}, \frac{PO_4}{3}, \&c$ Salts

containing M in the acidic radicle— Ni_2MO_4 , where $M = Mn$ or Fe , also Ni_2MnO_4 ($?NaFeO_4$), also $\alpha MO yFe_2O_3$, where $M =$ decidedly positive metal. Acids— $HMnO_4$.

The oxides MO are basic. They react with acids to form salts MX_2 . MnO and FeO are readily oxidised by standing in air, FeO very rapidly. NiO and CoO , on the other hand, are obtained by heating the higher oxides in air. The oxides M_2O_3 are basic, forming salts MX_3 . In the case of Mn only a few salts corresponding to M_2O_3 are known, e.g. $Mn_2(SO_4)_3$. In the cases of Ni and Co the salts of M_2O_3 are hardly known, the oxides dissolve in cold conc acids, probably forming salts, but on warming salts of MO are obtained. Both Fe_2O_3 and Mn_2O_3 show feebly acidic properties, as they combine with some more basic oxides, e.g. CaO, BaO, ZnO .

The oxides M_2O_3 react with acids, for the most part, as compounds of MO with MO_2 . Fe_2O_3 , however, is said to produce a few corresponding salts, e.g. Fe_2Cl_4 and $Fe_2(SO_4)_3$. It is doubtful whether MnO_2 form any corresponding salts, with acids it usually evolves O , and forms managanous salts MnX_2 . It combines with several more positive oxides, e.g. CaO or Br_2O , to form compounds $\alpha M^{II}O yMnO_2$. MnO_2 and Mn_2O_3 are extremely unstable acidic oxides. The acid of Mn_2O_3 , viz. $HMnO_3$ or $H_2Mn_2O_3$, is known, and from it a large series of salts has been obtained. The acid of MnO_2 (H_2MnO_4) has not been isolated, but the manganates $M^{II}MnO_4$ are well known salts. Ferrates, M_2FeO_4 , corresponding to the manganates, are known although neither the acid H_2FeO_4 nor the anhydride FeO_3 has been isolated.

The sulphides MS are basic, but MnS shows slightly acidic properties. MnS forms a compound with the very positive K_2S ($K_2S \cdot 3MnS$), CoS , on the other hand, combines with the slightly negative As_2S_3 ($2CoS \cdot As_2S_3$). Fe_2S_3 forms compounds with Ag_2S , CuS , &c, which may be regarded as sulphoferrites.

The *haloid compounds* MX_2 or IX_3 , are generally formed by direct union of the elements, also by dissolving the oxides MO in the haloid acids HX and evaporating. Fe is the only member of the group which certainly forms haloid compounds containing more halogen than MX_2 . There are, however, indications of the existence of $MnCl_3$ and $MnCl_4$. The haloid compounds which have been gasified are $MnCl_2$, $FeCl_2$ and $FeCl_3$. There are no indications of the existence of gaseous molecules Mn_2Cl_4 , but the V D s of ferrous and ferric chlorides point to the existence of the gaseous molecules $FeCl_2$ and $FeCl_3$ at high temperatures, and Fe_2Cl_4 and Fe_3Cl_8 at lower temperatures.

The *salts* of the metals of the iron group belong to the series MX_n and $MX_n \cdot xH_2O$, where $X = NO_3, ClO_4, SO_4, CO_3, PO_4$, &c. Fe forms many salts of both series, Mn forms chiefly many ganous salts MnX_n , but a few manganic salts are known, e.g. $Mn_2(SO_4)_3$, Ni forms only nickelous salts NiX_2 . The cobaltic salts are represented by a few double salts, e.g. $Co(NO_2)_2 \cdot 3KNO_2$, but the cobaltous salts are numerous. Ferrous salts are perfectly definite, but are fairly easily oxidised to ferric. Ferric and manganic sulphates form alums, $K_2SO_4 \cdot Fe_2(SO_4)_3$ or $Mn_2(SO_4)_3 \cdot 24H_2O$. Many of the salts MX_2 are isomorphous with the corresponding salts of Cu, Zn, Mg, and Cd.

The only member of the series which forms *acidic oxides* that have been isolated is Mn (*v supra*). Permanganic acid, $HMnO_4$, has also been isolated, but no oxyacid of Fe, Ni, or Co.

The iron elements exhibit analogies with several other families of elements. Their relations with the elements of Group III, especially with Al, Ga, and In, are shown in the composition of the salts MX_3 , in the existence of alums, e.g. $K_2SO_4 \cdot M(SO_4)_3 \cdot 24H_2O$, where $M = Al, Ga, In, Fe, or Mn$, also in the existence and dissociation of the gaseous molecules Fe_2Cl_4 , Al_2Cl_6 , and Ga_2Cl_6 . The analogies shown by Ni and Co to the Al family are very slight.

The iron family is distantly connected with the halogens. This is shown in the existence of the salts $NiMO_4$, where $M = Mn$ (?), $Cr, Cl, or I$.

The relations between the iron family and the chromium family of Group VI are fairly well marked, the MX_3 salts are similar to the CrX_3 salts, MnO_2 resembles CrO_3 , in being acidic and forming salts M_2MnO_4 , analogous to and isomorphous with the chromates. Ferrates, M_2FeO_4 , are also known, although neither FeO , nor H_2FeO_4 , has been isolated. MnO_2 is not unlike CrO_3 , in some of its reactions. The sulpho ferrites, e.g. $CuFe_2S_4$, resemble the sulpho chromites, e.g. $ZnCr_2S_4$. The relations of Ni and Co to the Cr family are chiefly shown in the formulae of the nickelous, cobaltous, and chromous salts, MX_2 , in the existence of many ammonio-cobalt and ammonio-chromium compounds. The formation of the cyanogen acid $H_2M_2Cy_4$, and of salts of $H_2M_2Cy_4$, where $M = Mn, Fe, Co, or Cr$, is a point of similarity between Cr and the iron family.

The composition of the salts MX_2 is similar to that of the salts of Group II, so far as properties go these salts most resemble those of the odd series members of Group II, Mg, Zn, and Cd, the ammonio-compounds of Co also recall

the ammonio-compounds of Hg which is an odd series member of Group II.

There are some resemblances between Cu which belongs to Group I and the iron elements, thus the salts MX_2 resemble in many respects the cupric salts CuX_2 , there are many ammonio-copper compounds, numerous double cyanides of Cu exist and some of those are probably derived from a cyanogen acid $H_2Cu_2Cy_4$.

Finally some of the physical properties of the iron family resemble the properties of the two other families of Group VIII, viz Rh, Ru, Pd, and Os, Ir, Pt, this resemblance is carried out in the formulae of some of the salts, and in the existence, and in some cases the composition and properties, of numerous complex cyanides; these complex cyanides are indicated in the following table —

$H_2M_2Cy_4$, and salts.	$H_2M_2Cy_4$, and salts.
$M = Mn, Fe, Co, Os, Ru$	$M = Fe, Co, Ir$
Salts of $H_2M_2Cy_4$, (acid not isolated)	
$M = Mn, Rh$	

The acid H_2PtCy_4 and salts of this acid exist, salts of H_2PdCy_4 are known. It should be noted that Ni does not form nickelous or nickelic cyanides analogous to any of the complex cyanides formulated above, but only ordinary double cyanides.

The position of the iron family of elements in the classification based on the periodic law is somewhat peculiar (*v Table on p 204 of vol II*). Mn is placed in the even series of Group VII, the only other members of this group as yet known are the halogens Fe, Ni, and Co form a division, or family, of Group VIII, the other families of this group are (1) Rh, Ru, Pd, and (2) Os, Ir, Pt. The iron family should therefore, strictly, include only Fe, Ni, and Co, Mn has been included in this family because of its close relations to Fe, but because of its position in Group VII, Mn has also been included in the halogen family (*v HALOGEN ELEMENTS*, vol II p 664). Each of the three families which together constitute Group VIII is separated from the others by many elements, the analogies between these families cannot be very close. The iron family is preceded in order of atomic weights by a series of elements, which begins with the very positive element K and ends with the element Cr which is both metallic and non metallic, the iron family is succeeded by series 5 which begins with the undoubtedly metallic Cu and ends with the no less undoubtedly non metal Br. The iron family forms one of the turning points in the swing of properties from very positive to very negative, the next similar turning point is marked by the second family of Group VIII, Rh, Ru, Pd, and the third turning point is marked by the last family of Group VIII, Os, Ir, Pt. Recent researches show that Co and Ni are perhaps separable each into two other elements, *v NICKEL* in this volume. In connection with this article *v CHROMIUM GROUP OF ELEMENTS*, vol II p 168, *COPPER GROUP OF ELEMENTS*, vol II p 250, *EARTHS, METALS OF THE*, vol II p 424, *HALOGEN ELEMENTS*, vol II p 664 *NOBLE METALS*, in this vol. For detailed properties of the members of the iron group *v COBALT*, vol II p 217, *IRON*, this vol p 51, *MANGANESE* and *NICKEL*, in this vol.

M M P M

ISAMIC ACID *v* ISATIN.ISATANE *v* ISATYDEISAMIC ACID $C_8H_5NO_3$, *is*

$C_8H_5(NH_2)COCO_2H$ *o* Amido phenyl glyoxylic acid. Formed, as potassium salt, by boiling isatin with conc KOHAq (Laurent, *A Ch* [3] 3, 371, Erdmann, *J pr* 24, 13). Obtained synthetically by reduction of an alkaline solution of *o*-nitro-phenyl-glyoxylic acid with ferrous sulphate (Claisen & Shadwell, *B* 12, 353). By decomposing the lead salt with H_2S and evaporating at atmospheric temperature *in vacuo* isamic acid may be obtained as an amorphous white powder, soluble in cold water. The acidified solution deposits, after a while, crystals of isatin, which is its anhydride, and the separation may be hastened by warming. With acetone in alkaline solution isamic acid forms (*Py* 3)-methyl quino line (*Py* 1) carboxylic acid [241°]. Its silver salt forms microscopic needles (W Pitzinger, *J pr* [2] 33, 100, Beyer, *J pr* [2] 33, 416). Acetophenone gives, in like manner, phenyl-quino line carboxylic acid.

Salts—KA' faint yellow crystals, which dissolve in conc KOHAq, forming a deep violet red solution, turned yellow on dilution with water. Its solution gives a yellow flocculent pp with lead acetate, and, when concentrated, it is also ppd by $BaCl_2$ —BaA', (at 150°) scales.—AgA' beautiful yellow prisms, sol water.

Acetyl derivative $C_8H_5(NHAc)COCO_2H$ [160°]. Obtained from acetyl-isatin by dissolving in cold dilute NaOHAq and ppg with dilute H_2SO_4 (Suida, *B* 11, 586). Needles (from alcohol). Sl sol cold water, m sol alcohol, ether, and benzene. Boiling HClAq converts it into isatin. Sodium amalgam reduces it to the acetyl derivative of *o* oxy *o* amido phenyl acetic acid $C_8H_5(NHAc)CH(OH)CO_2H$, which forms colourless needles [142°] converted by HI into oxindole.

Chloro isamic acid $C_8H_5Cl(NH_2)COCO_2H$. From chloro isatin by warming with KOHAq (Erdmann, *J pr* 19, 339, 24, 5, *A* 33, 129, Laurent, *A Ch* [3] 3, 378). Not known in the free state, since, when its salts are acidified, chloro-isatin is formed—KA' light yellow flattened needles (from alcohol), v sol water—BaA', aq pale yellow needles—BaA', 3aq brilliant deep yellow laminae—PbA', 2aq brilliant yellow gelatinous pp, which in a few minutes changes to scarlet PbA', aq—AgA' light yellow pp, sol boiling water—The cupric salt is ppd as a brownish red powder, which changes to blood red.

Di-chloro-isamic acid $C_8H_3Cl_2(NH_2)COCO_2H$. Formed by dissolving di chloro isatin in hot KOHAq. Separates on addition of HCl as a yellow pp, which, even when exposed over sulphuric acid *in vacuo* in the cold, splits up into water and di chloro isatin. It dissolves in water forming a light yellow solution, which becomes turbid at 60° and deposits di chloro isatin—KA' aq yellow laminae—BaA', 2aq golden needles—CuA', reddish brown pp changing to greenish yellow and crimson—AgA'. small yellowish needles (from hot water).

Bromo-isamic acid $C_8H_5Br(NH_2)COCO_2H$. Formed by warming bromo isatin with aqueous caustic potash (Gericke, *J pr* 95, 176, 257). The free acid splits up at once into isatin and bromo-

isatin—LA' easily soluble cauliflower like crystals—NaA' warty crystals—BaA', 3aq yellow prisms—CuA', 2aq red granular pp—PbA', 2aq yellow pp, changing to a scarlet crystalline powder—ZnA', 2aq brownish pp, changing to red granular powder—AgA' light yellow needles (from hot water).

Di-bromo-isamic acid $C_8H_3Br_2(NH_2)CO_2H$. From di bromo isatin and hot KOHAq (Erdmann, *J pr* 19, 360). The free acid is ppd by adding HCl to a conc solution of the K salt as a light yellow powder, soluble in a large quantity of water. By desiccation, even at 15° *in vacuo*, it is decomposed into di bromo isatin and water—KA' aq pale yellow needles, v sl sol water.

Ethylether EtA' [105°]. From the silver salt and EtI (Baeyer & Oekonomides, *B* 15, 2099). V sol ordinary solvents. At 110° it gives di bromo isatin.

Sulpho-isamic acid

$C_8H_5(SO_3H)(NH_2)COCO_2H$. From isatin sulphonic acid and excess of alkali (G & A Schlieper, *A* 120, 12). Only known in its salts, which are converted by mineral acids into isatin sulphonic acid—KA' aq waxy yellow prisms, v sol water—BaA', 3aq long silky lemon yellow needles, v sol boiling water, insol alcohol—PbA', 14aq yellow needles, v sol water—AgA', 14aq pale yellow needles, sl sol water.

Isomeride of isamic acid *v* m AMIDO-PHENYL GLYOXYLIC ACID

p Methyl isamic acid *Acetyl derivative* $C_8H_5(CH_3)(NHAc)COCO_2H$ [172°]. Small white needles, soluble in alcohol and hot water, very sparingly in ether, benzene, ligroin, and CS₂. Formed by the action of cold aqueous alkalis upon acetyl *p* methyl pseudo isatin.

Ethyl ether of the acetyl derivative $C_8H_5(CH_3)(NHAc)COCO_2Et$ [79°]. White glistening plates, insol water. Formed by boiling acetyl *p* methyl pseudo isatin with dilute alcohol (Duisberg, *B* 18, 198).

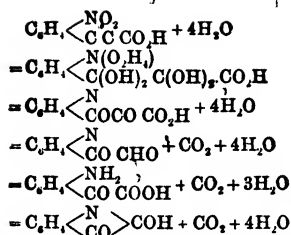
Amide of the acetyl derivative

$C_8H_5(CH_3)(NHAc)COCONH_2$ [141°]. Formed by the action of alcoholic NH_3 on the acetyl derivative of methyl ψ isatin (Panaotović, *J pr* [2] 33, 72). Trimetric prisms (from alcohol), insol water.

ISATILIM *v* ISATINISATIMIDE *v* ISATINISATIN $C_8H_5NO_2$, *is* $C_8H_5\langle\begin{smallmatrix} CO \\ N \end{smallmatrix}\rangle COH$

(stable form) or $C_8H_5\langle\begin{smallmatrix} CO \\ NH \end{smallmatrix}\rangle CO$ (transition form) or pseudo isatin. *Anhydride of isamic acid* *o*-Amido phenyl glyoxylic lactum [201°].

Formation—1 Discovered simultaneously by Laurent (*A Ch* [3] 3, 371) and by Erdmann (*J pr* 24, 11) as a product of the oxidation of indigo by nitric acid (L) or chromic acid (E)—2 By oxidising amido oxindole with $FeCl_3$, nitrous acid, or $CuCl_2$ (Baeyer, *B* 11, 1228). The oxindole may be obtained from phenyl acetic acid by nitration and reduction (Baeyer, *B* 11, 583), and converted successively into nitroso oxindole and amido oxindole—3 Prepared by boiling *o* nitro phenyl propionic acid with alkalis (Baeyer, *B* 13, 2259). The reaction possibly takes place as follows—

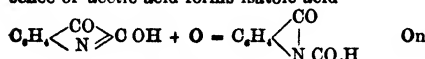


(Michael, *J pr* [2] 25, 255) —4 Together with $\text{N}_2(\text{C}_6\text{H}_7\text{N}, \text{CO}_2\text{H})_2$ by dissolving isatogenic ether in aqueous Na_2CO_3 (Baeyer, *B* 15, 55) —5 From isatogenic acid by dissolving in H_2SO_4 and adding water (Baeyer *B* 14, 1742) —6 By oxidising carbostyryl with alkaline KMnO_4 (Friedlander & Ostermaier, *B* 14, 1921) —7 By boiling nitroso (γ) oxy carbostyryl with conc HClAq (Baeyer & Homolka, *B* 16, 2217) —8 By heating anthroxanic acid with FeSO_4 and dilute ammonia (Schillinger & Wleügel, *B* 16, 2224)

Preparation —1 Finely powdered indigo (50g of 65 p c) is made into a thin paste with boiling water, and a dilute solution of CrO_3 (30 g) is added. The liquid is boiled till it begins to froth strongly, and is then filtered hot, when isatin separates on cooling. The yield is moderate (9 g) (Sommaruga, *A* 190, 369) —2 A mixture of indigo (100 pts), boiling water (300 pts), and nitric acid (70 pts of *S G* 135) is boiled for two minutes, diluted with boiling water (2000 pts), boiled for five minutes more and filtered. The isatin separates on cooling. It may be purified by dissolving in aqueous KOH , adding HCl as long as it forms a black or brown pp, filtering adding more HCl to the clear yellow filtrate, and washing the red pp of isatin (Forrer, *B* 17, 976, Hofmann, *A* 53, 11)

Properties —Yellowish-red monoclinic prisms, $a, b, c = 42.5 : 1.503, \beta = 85^\circ 18'$ (Bodewig, *Z K* 4, 65). Has no odour, but a bitter taste. Sl sol cold, m sol boiling, water, forming a reddish yellow solution. V c sol alcohol, sl sol ether. The alcoholic solution imparts an unpleasant odour to the skin.

Reactions —1 It is not attacked by dilute nitric acid, but a stronger acid forms nitro-oxy benzoic acid, while conc HNO_3 forms oxalic acid —2 Nitrous acid, acting upon isatin suspended in water, forms nitro-oxy-benzoic acid (Hofmann, *A* 115, 280), while, in presence of alcohol, nitrous acid forms benzoic acid (Baeyer & Knop, *A* 140, 4) —3 Chromic acid in presence of acetic acid forms isatoic acid



the other hand, acetyl isatin $\text{C}_6\text{H}_7\text{N} \begin{array}{l} \text{CO} \\ \text{N} \end{array} \text{---} \text{NAC} \text{---} \text{CO}$, being a derivative of pseudo isatin, yields acetyl-o-amido benzoic acid $\text{C}_6\text{H}_7(\text{NHAc})\text{CO}_2\text{H}$ on similar treatment (E v. Meyer & J Bellmann, *J pr* [2] 33, 30) —4 Chlorine and bromine form products of substitution —5 Cold aqueous caustic potash forms a red solution, which on boiling immediately becomes yellow, and then contains potassium isatate. If the solution be concentrated by distillation, decomposition suddenly takes place, aniline and hydrogen being

given off —6 Aqueous ammonia forms isamic acid. Alcoholic NH_3 gives a variety of products, called by Laurent imesatin, imasatin, amisatin, isatimide, and isatilum. By heating isatin with alcoholic NH_3 in sealed tubes Sommaruga (*A* 190, 371) obtained di-imido isatin, oxy-di-imido isatin, and deoxy imido isatin —7 Isatin combines with alkaline bisulphites —8 SO_2 has no action —9 H_2S gives di-thio isatide —10 Soda amalgam reduces it to isatide $\text{C}_6\text{H}_7\text{N}_2\text{O}$, and dioxindole $\text{C}_6\text{H}_7\text{NO}_2$ —11 Ammonium sulphide reduces it to isatide —12 Zinc and dilute H_2SO_4 form isatide —13 Zinc-dust and a little dilute HCl form isatide and dioxindole (Baeyer, *B* 12, 1309) —14 Zinc-dust added to a cold solution of isatin in HOAc forms a colourless solution (? of isatin hydride) which becomes coloured again on exposure to the air, isatin being reproduced —15 Aqueous HI (*S G* 14) at 100° forms isatide. At 140° it forms a dark-green insoluble mass, whence boiling alcohol leaves a dark grey residue, isatochlorin ($\text{C}_{12}\text{H}_{11}\text{N}_2\text{O}_2$), whilst the alcoholic solution, mixed with water, gives a pp, whence ether extracts red isatopurpurin ($\text{C}_{12}\text{H}_{11}\text{N}_2\text{O}_2$), while white isatone ($\text{C}_{12}\text{H}_{11}\text{N}_2\text{O}_2$) remains undissolved (Schutzenberger, *Bl* [2] 4, 170) —16 PCl_5 forms 'isatin chloride' (Baeyer, *B* 12, 456) —17 Hydroxylamine forms nitroso oxindole or isatoxim —18 Alcoholic KCl has no action —19 A solution of isatin, warmed with phenyl hydrazine, forms a pp of the phenyl hydrazide —20 In presence of H_2SO_4 or ZnCl_2 isatin forms condensation products, with (2 mols of) toluene, phenol, di-methyl aniline, thiophene, &c, by elimination of water (1 mol). Thus, phenol and H_2SO_4 give di-oxy-di-phenyl oxindole

$\text{C}_6\text{H}_7\text{N} \begin{array}{l} \text{C(C}_6\text{H}_5\text{OH)}_2 \\ \text{NHCO} \end{array}$, toluene gives di-tolyl-oxindole, while di-methyl aniline gives tetramethyl di-amido di-phenyl oxindole [234°] (Baeyer, *B* 18, 2642) —21 When heated with phenyl cyanate, for three hours at 130° , it forms the anilide of isatin-o-carboxylic acid, which crystallises from alcohol in needles [180° - 185°] (Gumpert, *J pr* [2] 31, 120, 32, 283) —22 Phenyl mercaptan, added to a hot alcoholic solution of isatin, forms silky needles of a compound ($\text{C}_6\text{H}_7\text{NO}_2$)($\text{C}_6\text{H}_5\text{SH}$), insol water, but decomposed by hot benzene into its components (Baumann, *B* 18, 890) —23 By boiling with *m*-amido-benzoic acid and alcohol there is formed an acid which is termed isat-amido-benzoic acid $\text{C}_6\text{H}_7\text{N} \begin{array}{l} \text{CNC}_6\text{H}_4\text{CO}_2\text{H} \\ \text{NH} \end{array} \text{---} \text{CO}$ [253°] *m*-Amido-benzamide gives, in like manner, the corresponding amide [c 280°] (Schiff, *A* 218, 192) —24 Heating with tolylene diamine forms $\text{C}_{12}\text{H}_{11}\text{N}_2$ [290°], which is probably a quinoxaline (Hinsberg, *A* 237, 344)

Metallic derivatives — $\text{C}_6\text{H}_7\text{AgNO}_2$ obtained by mixing isatin with water at 0° , adding KOH (1 mol) followed immediately by AgNO_3 (1 mol) (Baeyer & Oeconomidis, *B* 15, 2095) Red pp — $\text{C}_6\text{H}_7\text{AgNO}_2\text{NH}_3$ Formed in presence of ammonia (L) — $\text{Cu}(\text{C}_6\text{H}_7\text{NO}_2\text{NH}_3)_2$ From cupric acetate and an ammoniacal solution of isatin (Laurent) Brown pp —Potassium isatin is present in the violet red solution obtained by dissolving isatin in conc KOHaq

Indophenine reaction —A solution of isatin

in conc H_2SO_4 , shaken with benzene containing thiophene, is coloured blue, through formation of indophenine (*q v*). A similar reaction is given by most derivatives of thiophene

Combinations with bisulphites $\text{C}_6\text{H}_4\text{NO}_2\text{KHSO}_3 \cdot 2\text{aq}$ Formed by boiling isatin with aqueous KHSO_3 , or by saturating a solution of potassium isatate with SO_2 (Laurent) Large, pale yellow tables, v e sol water and boiling alcohol, sl sol cold alcohol Gives with lead acetate a pp of isatin and lead sulphite — $\text{C}_6\text{H}_4\text{NO}_2(\text{NH}_4)\text{HSO}_3$, pale yellow tables, sl sol cold water — $\text{C}_6\text{H}_4\text{NO}_2(\text{NH}_4\text{C}_2\text{H}_5)\text{HSO}_3$ (Schiff, *A*, 144, 49) — $\text{C}_6\text{H}_4\text{NO}_2(\text{NH}_4\text{Ph})\text{HSO}_3$ Crystals, v. sol water (S)

Acetyl- ψ -isatin $\text{C}_6\text{H}_3\text{N}(\text{CO})_2\text{CO}$ [141°]

Prepared by heating isatin (1 pt) with Ac O (2 pts) for 4 hours (Suida, *B* 11, 584) Yellow needles (from benzene) V sol benzene and alcohol, sl sol cold water Resolved into isatin and acetic acid by boiling with water, or, more readily with HClAq Cold NaOHAq dissolves it, forming $\text{C}_6\text{H}_3\text{N}(\text{NHAc})\text{CO CO}_2\text{Na}$ and, on warming, isatic acid Chromic acid in HOAc oxidises it to $\text{NHAc C}_6\text{H}_3\text{CO}_2\text{H}$

Benzoyl- ψ -isatin $\text{C}_6\text{H}_3\text{N}(\text{CO})_2\text{CO}$

From isatin and BzCl (Schwartz, *C R* 56, 1050)

Oxim $\text{C}_6\text{H}_3\text{N}_2\text{O}_2$ *re* $\text{C}_6\text{H}_3\text{N}(\text{C}(\text{NOH}))_2$ Isat

oxim Nitroso oxindole [c 202°] Formed by passing nitrous acid into a l p c aqueous solution of oxindole (Baeyer & Knop, *A* 140, 34) Formed also by boiling diazo nitroso oxindole chloride with alcohol and HCl (Gabriel & R Meyer, *B* 14, 2332) Obtained by treating isatin with hydroxylamine (Gabriel, *B* 16, 518, Baeyer & Comstock, *B* 16, 1706) Very slender golden needles Decomposed by fusion V sl sol water, sl sol alcohol Dissolves in KOHaq , forming a dark reddish brown solution Not decomposed by boiling aqueous KOH — $\text{C}_6\text{H}_3\text{AgN}_2\text{O}_2$ orange pp, got by adding dilute NH_3 to an alcoholic solution of isatin and AgNO_3 Dries up to a brick red powder

Mono-ethyl-ether of the oxim

$\text{C}_6\text{H}_3\text{N}(\text{C}(\text{NOEt}))_2\text{C}(\text{OH})$ *Isato-ethyl oxim* [188°] Fine yellow needles, soluble in caustic alkalis, formed by the action of ethyl iodide on the silver salt of isatoxim By successive treatment with zinc-dust and HOAc and with FeCl_3 it is converted into isatin

Di-ethyl-ether of the oxim

$\text{C}_6\text{H}_3\text{N}(\text{C}(\text{NOEt}))_2\text{C}(\text{OEt})$ *Ethyl-isato-ethyl-oxim* Crystalline solid, formed by the action of ethyl iodide on the silver salt of the mono-ethyl-ether (Baeyer & Comstock, *B* 16, 1706)

Oxim of ψ -isatin $\text{C}_6\text{H}_3\text{N}(\text{CO})_2\text{C}(\text{NOH})$

Isomytroso- ψ indoxyl Formed by the action of nitrous acid on ethyl indoxyllic acid

$\text{C}_6\text{H}_3\text{N}(\text{C}(\text{OEt}))_2\text{C}(\text{CO}_2\text{H})$ Flat yellow needles Decomposes at about 200° Sol alkalis and reppd by CO_2 With phenol and H_2SO_4 it does not give Liebermann's reaction Reduction with zinc-dust followed by oxidation with Fe_2Cl_3 yields isatin.

Ethyl ether of the oxim of ψ isatin

$\text{C}_6\text{H}_3\text{N}(\text{CO})_2\text{C}(\text{NOEt})$ From ψ isatoxim, KOH , and EtI yellow plates, dissolves in sodium ethylate with a blue colour, reduction with zinc dust followed by oxidation with Fe_2Cl_3 yields isatin Forms a violet solution when warmed with aqueous NaOH

Di-ethyl-derivative of the oxim of

ψ -isatin $\text{C}_6\text{H}_3\text{N}(\text{CO})_2\text{C}(\text{NOEt})_2$ [99°], yellow needles, sublimable, easily soluble in alcohol and ether, sparingly in hot water, insol alkalis On reduction with zinc dust and oxidation with Fe_2Cl_3 it yields ethyl-pseudo isatin. (Baeyer, *B* 15, 782, 16, 2191)

Phenyl hydrazide $\text{C}_{11}\text{H}_7\text{N}_3\text{O}$ [211°]

Fine yellowish red needles Formed as a yellow crystalline pp by boiling a solution of isatin in 1,000 pts of water with phenyl hydrazine hydrochloride (*q v*), the pp is quite distinct with a solution of 1 in 20,000 (E Fischer, *B* 17, 577)

Chloride $\text{C}_6\text{H}_3\text{ClNO}$ *re* $\text{C}_6\text{H}_3\text{N}(\text{CO})_2\text{CCl}$

[c 180°] Formed by warming isatin (5 g) with PCl_5 (7 g) and benzene (9 g) (Baeyer, *B* 11, 1296, 12, 456) Brown needles Decomposed on fusion V sol ether, forming a blue solution Decomposed by moist air Potash converts it into isatin HI gives indigo, zinc dust and HOAc do the same

Chloro-isatin $\text{C}_6\text{H}_3\text{ClNO}_2$ [243°] S 1 at 0°, c 5 at 100° Obtained by passing chlorine in sunlight into boiling water containing isatin in suspension The mono and di chloroisatin are separated by recrystallisation from alcohol, in which the former is much less soluble (Hofmann, *A* 53, 12, Laurent, *A Ch* [3] 3, 462, Erdmann, *A* 33, 129, Dorsch, *J pr* [2] 33, 49) Orange prisms (from alcohol) V sl sol water and alcohol Its solution imparts an unpleasant odour to the skin Decomposed on fusion Hot KOHaq forms potassium chloro isatate Distillation with NH_3 gives *p* chloro aniline An ammoniacal solution of AgNO_3 added to an alcoholic solution of chloro isatin ppts $\text{C}_6\text{H}_3\text{ClAgNO}_2\text{NH}_3$

Di chloro isatin $\text{C}_6\text{H}_2\text{Cl}_2\text{NO}_2$ [186°] Obtained from the alcoholic mother liquors that have yielded chloro isatin Small, reddish yellow needles or short laminae (from alcohol) M sol alcohol When thrown upon solid potash, moistened with alcohol, a red solution is formed which solidifies to a violet black magma of $\text{C}_6\text{H}_2\text{KCl}_2\text{NO}_2$, the solution of which gives a pp with AgNO_3 Hot aqueous KOH forms potassium di chloro isatate Distillation with KOH forms di chloro aniline Chlorine does not act on its aqueous solution, but with its alcoholic solution it forms tetra chloro quinone and other bodies KHSO_3 forms light yellow needles of $\text{C}_6\text{H}_2\text{Cl}_2\text{NO}_2\text{KHSO}_3$, sl sol boiling water Di chloro isatin (10 g) oxidised by CrO_3 (15 g) in glacial acetic acid (60 g) as described under *Bromo isatin* forms di chloro isatoxo acid (*q v*)

Bromo isatin $\text{C}_6\text{H}_3\text{BrNO}_2$ *re*

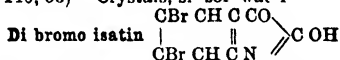
$\text{CBr CH}_2\text{C}(\text{CO})_2\text{C}(\text{OH})$ [255°] Formed, to gether with di bromo isatin, by the action of bromine and water on isatin (Erdmann, *J. pr.*

19, 558, Hofmann, *A* 53, 40) The product is exhausted with boiling water and the crystals that separate on cooling are recrystallised from alcohol whence bromo-isatin crystallises first. Orange prisms. Aqueous KOH converts it, even in the cold, into potassium bromo-isatate. Distillation with KOH yields *p* bromo aniline. When mixed with alcoholic ammonium sulphide and exposed to the air it yields bromo-indigo. Bromo isatin (10 g) is oxidised by a mixture of glacial acetic acid (90 g) and CrO_3 (15 g) to bromo-isatoic acid. The mixture must first be kept ice-cold but after 12 hours it may be slowly raised to 10° . The product, when cold, is poured into cold dilute H_2SO_4 and the yellow sandy powder crystallised from a mixture of alcohol and acetone (Dorsch, *J pr* [2] 33, 45).

cetyl derivative $\text{C}_{18}\text{H}_{15}\text{BrAcNO}_2$. [172°] Formed by boiling bromo isatin (5 pts) with Ac_2O (8 pts) (Baeyer & Oeconomidis, *B* 15, 2096). Long straw-yellow prisms (from benzene).

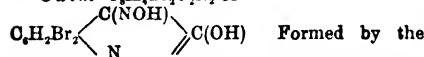
Chloride $\text{C}_8\text{H}_5\text{BrClNO}$. Formed by treating bromo isatin with PCl_5 (Baeyer, *B* 12, 1315). Reddish brown needles, sol hot benzene and ether. HI gives bromo-indigo.

Oxim $\text{C}_8\text{H}_5\text{BrNO}(\text{NOH})$. From isatoxim and bromine water in the cold (Baeyer & Knop, *A* 140, 35). Crystals, sl sol wat r.



[250°] Formed as above or by digesting bromo isatin or isatin with bromine in sunshine. Formed also by heating bromo isatin (1 mol) with bromine (2 mols) in HOAc for 20 hours (Baeyer & Oeconomidis, *B* 15, 2098). Orange prisms (from alcohol). Gives di bromo aniline when distilled with KOH— $\text{C}_8\text{H}_5\text{Br KNO}_2$, bluish violet scales, sl sol water. Very stable but converted into di bromo isatate by warming with KOH aq — AgA' brownish violet powder— $\text{C}_8\text{H}_5\text{Br}_2\text{NO}_2\text{KHSO}_4$, yellow solid, v sl sol water.

Oxim $\text{C}_8\text{H}_5\text{Br}_2\text{O}_2\text{N}$, or



Formed by the action of hydroxylamine on di bromo isatin (Baeyer & Comstock, *B* 16, 1708). Yellow pointed needles. Carbonises without melting at about 255° . Sol caustic alkalis but precipitated by CO_2 .

Mono-ethyl-ether of the oxim

$\text{C}_8\text{H}_5\text{Br}_2 \begin{array}{c} \diagup \text{C}(\text{NOEt}) \diagdown \\ \text{N} \end{array} \begin{array}{c} \diagup \\ \diagdown \end{array} \text{C}(\text{OH})$ *Di-bromo isato-ethyl oxim*. [252°] Yellow needles, formed by the action of ethyl iodide on the silver salt of di bromo isatoxim.

Di-ethyl ether of the oxim

$\text{C}_8\text{H}_5\text{Br}_2 \begin{array}{c} \diagup \text{C}(\text{NEt}) \diagdown \\ \text{N} \end{array} \begin{array}{c} \diagup \\ \diagdown \end{array} \text{C}(\text{OEt})$ *Di-bromo isato-ethyl oxim* [116°] Long yellow, silky needles, formed by the action of ethyl iodide on the silver salt of the mono ethyl ether.

Tri bromo isatin Oxim

$\text{C}_8\text{H}_5\text{Br}_3 \begin{array}{c} \diagup \text{C}(\text{NOH}) \diagdown \\ \text{N} \end{array} \begin{array}{c} \diagup \\ \diagdown \end{array} \text{C}(\text{OH})$ [162°] From the oxim of isatin and bromine in excess (*B* & *K*). Dirty violet needles (from alcohol). Insol

water, v sol boiling alcohol. At 190° it sublimes as red needles.

Nitro isatin $\text{C}_8\text{H}_5\text{NO}_4(\text{NO}_2)$ [226°–230°] Prepared by nitration of isatin dissolved in H_2SO_4 , by addition of powdered KNO_3 (Baeyer, *B* 12, 1312). Sparingly soluble in water, more easily in alcohol.

Bromo nitro isatin

$\text{C}_8\text{H}_5(\text{NO}_2)\text{Br} \begin{array}{c} \diagup \text{CO} \diagdown \\ \text{N} \end{array} \begin{array}{c} \diagup \\ \diagdown \end{array} \text{COH}$ [237°] From bromo-isatin, KNO_3 and H_2SO_4 (Dorsch). Clumps of orange crystals (from alcohol). Sol acetone, glacial acetic acid and alcohol, sl sol benzene and chloroform, v sl sol water and ether. Forms a dark red solution in NaOH aq , whence an orange powder presently separates.

ALKYL DERIVATIVES

Alkyl derivatives of isatin are either derived from the stable form $\text{C}_8\text{H}_5 \begin{array}{c} \diagup \text{CO} \diagdown \\ \text{N} \end{array} \begin{array}{c} \diagup \\ \diagdown \end{array} \text{COH}$

or from the transition form $\text{C}_8\text{H}_5 \begin{array}{c} \diagup \text{CO} \diagdown \\ \text{NH} \end{array} \text{CO}$ (pseudo isatin). Alkyl iodides acting on silver-isatin form alkyl derivatives of stable isatin $\text{C}_8\text{H}_5 \begin{array}{c} \diagup \text{CO} \diagdown \\ \text{N} \end{array} \begin{array}{c} \diagup \\ \diagdown \end{array} \text{COR}$

Alkyl derivatives of pseudo-isatin

$\text{C}_8\text{H}_5 \begin{array}{c} \diagup \text{CO} \diagdown \\ \text{NR} \end{array} \text{CO}$ are formed by the action of an alkaline solution of bromine or chlorine followed by alcoholic NaOH on the methyl-, ethyl-, phenyl-, &c indole carboxylic acids which are obtained by the action of HCl on phenyl methylhydrazine pyruvic acid, phenyl-ethyl hydrazine pyruvic acid, &c (Fischer & Hess, *B* 17, 559).

Methyl derivative of isatin $\text{C}_8\text{H}_5\text{O}_2\text{N}(\text{CH}_3)$

$\text{C}_8\text{H}_5 \begin{array}{c} \diagup \text{CO} \diagdown \\ \text{N} \end{array} \begin{array}{c} \diagup \\ \diagdown \end{array} \text{COMe}$ [102°] Formed by the action of methyl iodide on the silver compound of isatin (Baeyer & Oeconomidis, *B* 15, 2093). Red trimetric prisms. Sol ether, acetone, benzene, and CS_2 , less easily in alcohol, v sl sol. ligroin. It changes on keeping into methyl isatoid. Slowly dissolves in dilute KOH aq , forming a solution from which acids ppt isatin. Its solution in alcoholic ammonium sulphide forms indigo when exposed to the air.

Methyl-isatoid $\text{C}_8\text{H}_5\text{N}_2\text{O}_2$ (?) [219°] Formed by spontaneous change by keeping methyl isatin (Baeyer & Oeconomidis, *B* 15, 2094). Small yellow needles, sparingly soluble in all solvents. Dissolves in dilute NaOH on boiling, and on adding acid isatin is precipitated.

Methyl derivative of bromo-isatin

$\text{C}_8\text{H}_5\text{BrO NMe}$ [147°] Formed by the action of MeI on the silver compound of bromo-isatin (Baeyer & Oeconomidis, *B* 15, 2095). Red needles.

Bromo-methyl-isatoid [231°] Formed by spontaneous change of methyl bromo-isatin by keeping (*B* & *O*).

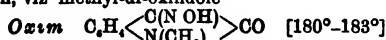
Methyl-pseudo-isatin $\text{C}_8\text{H}_5\text{NO}_2$, i.e.

$\text{C}_8\text{H}_5 \begin{array}{c} \diagup \text{CO} \diagdown \\ \text{NMe} \end{array} \begin{array}{c} \diagup \\ \diagdown \end{array} \text{CO}$ [134°] Red needles.

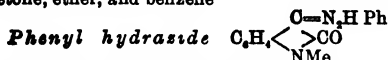
Formation—1 By the action of a cold alkaline solution of bromine or chlorine followed by hot alcoholic NaOH on methyl indole-carboxylic acid [212°], which is obtained by heating phenyl methyl hydrazine pyruvic acid with HCl (Fischer & Hess, *B* 17, 563)—2 By boiling di-

bromo-methyl oxindole with water (Colman, *C J* 55, 5, *A* 248, 118)

Reactions — Dissolves in alkalis with a yellow colour. With H_2SO_4 and benzene containing thiophene it gives the indophenine reaction. Gives a crystalline compound with phenyl-hydrazine. Yields the same product on oxidation in alkaline solutions as in acid solution, viz methyl-di-oxindole.

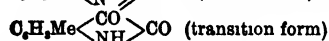
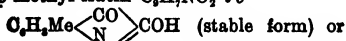


From methyl-pseudo isatin and hydroxylamine sulphate in hot aqueous solution (Colman, *C J* 55, 6). Tufts of small pale yellow needles, m sol. hot water, sl sol. cold water, v sol. alcohol, acetone, ether, and benzene.



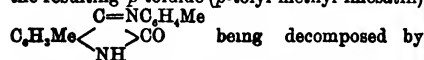
[146°] Formed by the action of phenyl-hydrazine hydrochloride and NaOAc on an aqueous solution of methyl-ψ isatin. A less pure product is produced by the action of phenyl-hydrazine hydrochloride on dibromomethyl-oxindole (Colman). Fascicular group of small yellow needles, insol water and petroleum ether, sl sol. ether, v sol. hot alcohol or benzene.

p-Methyl-isatin $C_6H_4Me \cdot NO_2$ *see*



Tolysatin Anhydride of amido tolyl-glyoxylic acid [184°] (P), [187°] (M)

Formation — From *p* toluidine and di chloroacetic acid by digestion in alcoholic solution, the resulting *p* toluidine (*p*-tolyl methyl imesatin)



HCl, and the crude product purified by boiling with aqueous KOH, ppg with HCl, and crystallising from alcohol and afterwards from water (P J Meyer, *B* 16, 2261, Fr Baeyer & Co, *B* Ref. 17, 867, Panacotovic, *J pr* [2] 33, 57)

Properties — Red crystals or glistening red plates. Sol. hot water and alcohol, sl sol. cold water, v sol. hot HCl aq. Dissolves in cold alkalis with a deep-violet colour, and on heating or standing it then takes up water, becoming yellow and forming methyl isatinic acid. Dissolves in conc. H_2SO_4 with a red colour. Gives the indophenine reaction. CrO_3 in acetic acid forms methyl isatinic acid. Boiling with Ac_2O for three hours forms the acetyl derivative of methyl-ψ isatin.

Acetyl derivative of methyl-ψ

isatin $C_6H_4Me \cdot \langle \begin{smallmatrix} CO \\ NAc \end{smallmatrix} \rangle CO$ [172°] Formed by boiling *p*-methyl-ψ isatin with Ac_2O for 8 hours (Panacotovic, *J pr* [2] 33, 71). Formed also by the action of HCl upon the acetyl derivative of *p*-methyl-ψ isatin *p* toluidine (Duisberg, *B* 18, 197). Lemon-yellow needles (from benzene), sol. chloroform and benzene, sl sol. water, alcohol, ether, ligroin, and CS_2 . Converted by cold dilute alkalis into acetyl-*p*-methyl-isatinic acid. Alcoholic NH_3 converts it into the amide of acetyl-*p*-methyl-isatinic acid $C_6H_4Me(NHAc)COCONH_2$, which crystallises from alcohol in trimetric columns [141°]. CrO_3 ,

in $HOAc$ oxidises it to methyl isatinic acid, which crystallises from boiling alcohol in trimetric plates and, when heated, decomposes at 245° with great increase in bulk, finally melting at 300°.

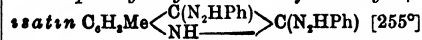


methyl oxindole [226°] Formed by the action of hydroxylamine (base) on *p* methyl isatin (P J Meyer, *B* 16, 2268). Long yellow prisms, sol. alcohol and NaOH, sl sol. water.

Phenyl hydrazide of methyl isatin

$C_6H_4Me \cdot \langle \begin{smallmatrix} C(N \cdot HPh) \\ N \end{smallmatrix} \rangle COH$ From *p* methyl isatin and phenyl hydrazine (P). Golden needles (from chloroform). Nearly insol water, sol. alcohol. May be sublimed at 240°, but melts above 300° with evolution of gas.

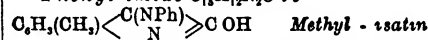
Di phenyl-hydrazide of methyl ψ



Formed by warming the acetyl derivative of methyl-ψ isatin with phenyl-hydrazine (P). Yellow trimetric columns (from alcohol) decomposed by fusion. Nearly insol water, sol. chloroform.

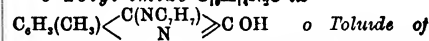
Imide $C_6H_4N_2O$ or $C_6H_4N_2O_2$ *p* Methyl imesatin. Formed by heating *p* methyl isatin *p* toluidine with alcoholic NH_3 at 100° (P J Meyer, *B* 16, 2264). Nearly colourless fine silky needles. V sl sol. hot alcohol and hot water. Insol. cold alcohol and water. Is not reconverted into *p* methyl isatin by acids or alkalis.

Phenyl imide $C_6H_4N_2O_2$ *see*



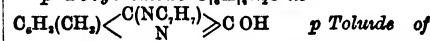
anilide Phenyl-methyl imesatin [240°] Formed by heating *p* methyl isatin with an absolute alcoholic solution of aniline (Meyer, *B* 16, 2267). Thick yellowish red tables or prisms. Sol. hot alcohol, sl sol. cold alcohol and water. Heated with acids or caustic alkalis it is resolved into its constituents.

o Tolyl imide $C_6H_4N_2O_2$ *see*

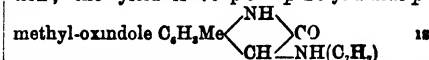


methyl-isatin *o*-Tolyl *p*-methyl imesatin [191°] Red prismatic crystals. Formed by heating *p*-methyl isatin with an absolute alcoholic solution of *o* toluidine. By HCl or hot NaOH it is resolved into its constituents (Meyer, *B* 16, 2268).

p-Tolyl imide $C_6H_4N_2O_2$ *see*



methyl isatin *p* Tolyl *p*-methyl imesatin [259°] Formed by heating di chloroacetic acid (1 mol) with *p* toluidine (4 mols) at 100°C, either alone or in aqueous or alcoholic solution, the yield is 70 p.c. *p* Tolyl amido *p*



is first formed and then undergoes oxidation by absorption of oxygen from the air. Formed also by heating di chloroacetamide (1 mol) with *p* toluidine (8 mols), and by heating *p* methyl isatin with an absolute alcoholic solution of *p*-toluidine (Meyer, *B* 16, 2261). Glistening yellow needles or plates. Sol. ether and hot alcohol, sparingly sol. cold alcohol, insol water. By cold

HCl it is resolved into *p* toluidine and *p* methylisatin. Hot HCl or hot NaOH gives *p* methylisatinic acid—Sodium salt crystallises in red prisms, decomposed by water.

Acetyl derivative

$C_6H_5Me \begin{smallmatrix} \text{NaO} \\ > \text{CO} \\ \text{C}=\text{N}(\text{C}_6\text{H}_5) \end{smallmatrix}$ [122°] Glistening red needles, insol water. By HCl it is split up into *p* toluidine and the acetyl derivative of *p*-methyl pseudo isatin (Duisberg, *B* 18, 190).

Bromo *p* toluidide $C_6H_4BrN_2O$ [210°] Red needles or prism. By heating with acids or alkalis it is split up into its constituents (P J Meyer, *B* 16, 2267).

o Di methyl ψ -isatin $C_6H_4Me \begin{smallmatrix} \text{CO} \\ < \text{NMe} > \end{smallmatrix} \text{CO}$

Methyl ψ -o tolsatin [157°] Formed by adding pyruvic acid to a solution of methyl-*o* tolylhydrazine in dilute HCl, and warming the resulting oil with a large quantity of phosphoric acid (S G 117). The resulting di methyl indole carboxylic acid is then warmed with NaOCl in slightly alkaline solution, when a product [152°] is obtained, which when warmed with water yields the di methyl isatin (Hegel, *A* 232, 221). Brick red needles (from water).

***p* Di methyl ψ isatin** $C_6H_4Me \begin{smallmatrix} \text{CO} \\ < \text{NMe} > \end{smallmatrix} \text{CO}$ [148°] Formed in like manner from methyl *p* tolylhydrazine and pyruvic acid, the di methyl indole carboxylic acid giving with alkaline hypochlorite a chlorinated compound crystallising in yellow needles [135°] which yield the di methyl isatin on dissolving in water (H). Red needles.

Ethyl pseudo isatin $C_6H_4 \begin{smallmatrix} \text{CO} \\ < \text{NEt} > \end{smallmatrix} \text{CO}$ *Lactam of ethyl isatinic acid* [95°]

Formation—1 By the action of a cold alkaline solution of chlorine followed by hot alcoholic NaOH on ethyl indole carboxylic acid [183°] which is obtained by the action of HCl on phenyl ethyl hydrazine pyruvic acid (E Fischer & Hess, *B* 17, 566).—2 By reduction of the di ethyl derivative of pseudo isatin- ω -oxim.

$C_6H_4 \begin{smallmatrix} \text{CO} \\ < \text{NEt} > \end{smallmatrix} \text{C}(\text{NOEt})$ with zinc dust and oxidation of the product with Fe_2Cl_6 (Baeyer, *B* 16, 2193). Large red plates, sol hot water, alcohol and ether. With thiophene and H_2SO_4 it gives a blue colouring matter soluble in ether. It dissolves in alkalis with a yellow colour, at once forming a salt of ethyl isatinic acid $C_6H_4 \begin{smallmatrix} \text{CO CO}_2H \\ < \text{NHEt} > \end{smallmatrix}$.

Ethyl pseudo isatin ω -oxim

$C_6H_4 \begin{smallmatrix} \text{C}(\text{NOH}) \\ < \text{NEt} > \end{smallmatrix} \text{CO}$ [162°] Yellow four-sided prisms. Formed by the action of hydroxylamine on ethyl pseudo-isatin. On reduction with zinc dust followed by oxidation with Fe_2Cl_6 it yields ethyl pseudo isatin. It does not yield indigo on treatment with ammonium sulphide (Baeyer, *B* 16, 2196).

Ethyl pseudo-isatin ω -oxim *Ethyl ether*

$C_6H_4 \begin{smallmatrix} \text{CO} \\ < \text{NEt} > \end{smallmatrix} \text{C}(\text{NOEt})$ [99°] From the ethyl ether of the ω -oxim of ψ -isatin by further ethylation (Baeyer, *B* 16, 2191). Yellow needles, v.

sol alcohol and ether, sl sol hot water. May be sublimed. Insol alkalis. After reduction with zinc dust, on oxidation with $FeCl_3$, it yields ethyl ψ isatin.

Bromo-ethyl-isatin $C_6H_4BrO_2NEt$ s.a.

$C_6H_4Br \begin{smallmatrix} \text{CO} \\ > \text{COEt} \end{smallmatrix} \text{N}$ [109°] From the silver compound of bromo isatin and EtI (Baeyer & Oeconomidis, *B* 15, 2095). Long red prismatic needles (from benzene). On keeping it slowly changes into bromo ethyl isatoid.

Bromo-ethyl isatoid $C_6H_4Br_2N_2O$ [245°] Fine needles. Sparingly soluble in all solvents, most easily in hot alcohol and acetone. Formed by spontaneous change of ethyl bromo isatin by keeping, also by the action of acetic anhydride on ethyl bromo isatin (Baeyer & Oeconomidis, *B* 15, 2095).

p Ethyl-isatin $C_6H_4NO_2$ s.a.

$C_6H_4Et \begin{smallmatrix} \text{CO} \\ > \text{COH} \end{smallmatrix} \text{N}$ [137°] Long red needles (Paucksch, *B* 17, 2805). Formed by heating with $HClAq$ the product of the action of di chloro acetic acid on *p*-amido phenyl-ethane $C_6H_4Et(NH_2)$ [14].

Di-bromo-ethyl-isatin $C_6H_4O_2NBr_2Et$ s.a.

$C_6H_4Br \begin{smallmatrix} \text{CO} \\ > \text{COEt} \end{smallmatrix} \text{N}$ [89°] From silver di-bromo isatin and EtI (B & O). Red crystals. *p* Methyl ψ ethyl- ψ isatin.

$C_6H_4Me \begin{smallmatrix} \text{CO} \\ < \text{NEt} > \end{smallmatrix} \text{CO}$ [110°] **Ethyl ψ -*p*-toluatin**. Formed from the *p*-toluidide by treatment with conc $HClAq$ (Duisberg, *B* 18, 197). Formed also from methyl ethyl indole carboxylic acid by treatment with a weak alkaline solution of NaOCl, and warming the resulting pp with water (Hegel, *A* 232, 219). Red needles or prisms, v sol alcohol, ether, benzene, and CS₂, sl sol water and ligroin. Sol aqueous alkalis. Gives the indophenine reaction.

p-Tolylimide $C_6H_4Me \begin{smallmatrix} \text{CO} \\ < \text{NEt} > \end{smallmatrix} \text{CO}$

***p* Toluidide** [152°] Formed by boiling the *p* tolylimide of *p* methyl isatin with NaOEt and EtBr (Duisberg, *B* 18, 198). Large orange-red prisms, v sol alcohol, acetic acid, benzene, and CS₂, sl sol ether, insol water. Split up by HCl into *p* toluidine and methyl ethyl- ψ isatin.

Isobutyl-bromo-isatin

$C_6H_4Br \begin{smallmatrix} \text{CO} \\ > \text{COC}_4H_9 \end{smallmatrix} \text{N}$. Crystallises with difficulty. Is converted by Ac_2O into iso butyl-bromo isatoid $C_6H_4Br_2O_2$ [210°] which crystallises in slender needles, sl sol all solvents (Baeyer & Oeconomidis, *B* 15, 2097).

Benzyl ψ -isatin $C_6H_4 \begin{smallmatrix} \text{CO CO} \\ < \text{N}(\text{CH}_2\text{Ph}) > \end{smallmatrix} \text{CO}$ [181°].

Formed by adding a solution of sodium benzyl indole carboxylate to one of NaOCl. The pp is dissolved in alcoholic NaOH, warmed gently, diluted with water, freed from alcohol by distillation, and ppd with water (Antriok, *A* 227, 364). Long silky red needles (from water). Almost insol cold water, v sol alcohol and ether.

Phenol-isatin v Di OXY-DI PHENYL-OXINDOLE. Tolu-isatin v Di TOLYL-OXINDOLE

(ϵ)-Naphth-isatin $C_{10}H_7 \begin{smallmatrix} \text{NH} \\ < \text{CO} > \end{smallmatrix} \text{CO}$ or

$C_6H_5 \cdot \begin{smallmatrix} N \\ \diagup \diagdown \\ CO \end{smallmatrix} > COH$ [255°] Formed by dissolving (s) naphthoxindole in alcohol and adding HOAc and $NaNO_2$. The isonitroso derivative so obtained is reduced with Sn and HCl, and subsequently oxidised with $FeCl_3$ (Hinsberg B 21, 117) Red needles Forms a compound with phenyl hydrazine [270°]

(*B*) - Naphth-isatin $C_{10}H_6 \cdot \begin{smallmatrix} N \\ \diagup \diagdown \\ CO \end{smallmatrix} > COH$ [248°]

Formed by dissolving (*B*) naphthoxindole in HOAc and adding sodium nitrite The isonitroso-body formed yields on reduction with Sn and HCl, and subsequent oxidation with $FeCl_3$, the isatin (Hinsberg, B 21, 115) Red needles, v sol ordinary solvents

AMMONIACAL DERIVATIVES OF ISATIN

Imesatin $C_8H_6N_2O \cdot \begin{smallmatrix} CO \\ \diagup \diagdown \\ N \end{smallmatrix} > CNH_2$ or

$C_8H_5 \cdot \begin{smallmatrix} CO \\ \diagup \diagdown \\ NH \end{smallmatrix} > CNH(?)$ or $C_8H_5 \cdot \begin{smallmatrix} C(NH) \\ \diagup \diagdown \\ N \end{smallmatrix} > COH$

Isatin imide (?) Obtained by Laurent (*J pr* 25 457) by passing dry NH_3 into a boiling alcoholic solution of isatin containing a little isatin in suspension Could not be obtained by Sommaruga (*B* 10, 432) Rectangular prisms insol water, v sl sol ether, m sol boiling alcohol Readily decomposed by heating with alcohol and HClAq into isatin and NH_3 KOH acts in like manner

Chloro-imesatin C_8H_5ClNO From chloro-isatin and alcoholic NH_3 (Laurent) Yellow six-sided prismatic tables, v sl sol boiling alcohol, insol ether Dissolves in KOH, giving a red liquid

Bromo-imesatin C_8H_5BrNO From bromo-isatin and boiling alcoholic NH_3 (Gercke, *Z* 1865, 593) Yellowish brown crystalline mass

Iso-amyl-imesatin $C_8H_5(C_4H_9)NO$ $\cdot \begin{smallmatrix} CO \\ \diagup \diagdown \\ N \end{smallmatrix} > CNHC_4H_9$ or

$C_8H_5 \cdot \begin{smallmatrix} CO \\ \diagup \diagdown \\ N \end{smallmatrix} > CNHC_4H_9$ or

$C_8H_5 \cdot \begin{smallmatrix} C(NC_4H_9) \\ \diagup \diagdown \\ N \end{smallmatrix} > COH(?)$ Formed by heating isatin with isoamylamine (Schiff, *A* 144, 53, *Z* [2] 4, 18) Yellow laminae, sl sol ether, v sol alcohol Decomposed by dilute acids or by prolonged treatment with water into isatin and isoamylamine

Phenyl-imesatin $C_8H_5(C_6H_5)NO$ *Amide of isatin* From isatin and aniline in boiling alcoholic solution (Engelhardt, *J* 1855, 541) Formed also by heating the compound of isatin with bisulphite of aniline, and crystallising from ether alcohol

Chloro-phenyl-imesatin $C_8H_5(C_6H_4Cl)NO$ From isatin and chloro aniline (*E*) Yellowish crystals

Bromo-phenyl-imesatin $C_8H_5(C_6H_4Br)NO$ Resembles the preceding (*E*)

Phenyl-chloro-imesatin $C_8H_5Cl(C_6H_5)NO$ Formed by adding aniline to a boiling solution of chloro isatin in alcohol (Engelhardt, *J pr* 65, 260) Yellow needles, insol water, v sol hot alcohol Resolved by boiling dilute acids into chloro-isatin and aniline Potash gives aniline and potassium chloro-isatate.

Phenyl-bromo imesatin $C_8H_5Br(C_6H_5)NO$ Resembles the above (*E*)

Di-phenyl-di-ethyl-di-amide of isatin $C_{24}H_{18}N_4O \cdot \begin{smallmatrix} CO \\ \diagup \diagdown \\ NH \end{smallmatrix} > C(NPhEt)$, or

$C_6H_5 \cdot \begin{smallmatrix} C(NPhEt) \\ \diagup \diagdown \\ N \end{smallmatrix} > COH(?)$ So-called phenyl

ethyl imesatin From isatin by heating with ethyl aniline (Schiff) Yellow laminae, sl sol ether, v sol alkalis Split up by acids, alkalis, or hot water into isatin and ethyl aniline

p-Methyl-imesatin v *Imide of p* Methyl isatin (*supra*)

Phenyl-*p*-methyl-imesatin v. *Amide of p*-methyl isatin (*supra*)

Imasatin $C_{11}H_{11}N_2O$ Formed by passing NH_3 into a boiling saturated solution of isatin in dilute alcohol (Laurent, *A Ch* [3] 8, 493) Greyish yellow granules, insol water and ether, v sl sol boiling alcohol Not attacked by boiling aqueous HCl Caustic potash dissolves it, and the solution is pptd by dilute HClAq

Di-chloro-imesatin $C_{10}H_8Cl_2N_2O$ From chloro isatin and alcoholic NH_3 (*L*) Slightly reddish powder

Di-bromo-imesatin $C_{10}H_8Br_2N_2O$ Formed by heating an alcoholic solution of bromo isatin with ammonia (Gercke, *Z* 1865, 593) Brownish yellow crystalline granules

Tetra-bromo-imesatin $C_{10}H_8Br_4N_2O$ From tetra bromo isatin and alcoholic NH_3 Reddish yellow scale. (*L*)

Amasatin $C_{10}H_8N_2O$ *Isamide Amide of Isamic acid?* One of the products of the action of ammonia on isatin Prepared by heating ammonium isamate till water is given off, and washing the residue with water (Laurent, *A Ch* [3] 3, 488, *J pr* 35, 117) Yellow powder, insol water, nearly insol ether, v sl sol alcohol, m sol boiling alcoholic NH_3 Cold KOHAq dissolves it, giving off NH_3 , and forming a yellow liquid containing potassium isamate HClAq forms a violet solution containing isamic acid

Di-chloro-amasatin $C_{10}H_6Cl_2N_2O$ Yellow powder, formed by evaporating a solution of ammonium chloro isatate (*L*)

Tetra-chloro-amasatin $C_{10}H_4Cl_4N_2O$ Formed by heating ammonium di chloro isatate

Di bromo amasatin $C_{10}H_6Br_2N_2O(?)$ Formed by evaporating an aqueous solution of ammonium bromo isatate and treating the pasty residue with water (Gercke) Orange yellow substance, sl sol water, insol alcohol and ether Dissolves with violet colour in acids

Amisatin $C_{10}H_8N_3O$ A product of the action of dilute alcoholic ammonia on isatin and pptd by adding water to the solution after isatimide and isatium have crystallised out (*L*) Minute needles, insol alcohol, sol alcoholic potash

Isamic acid $C_{10}H_8N_2O$ Produced by the action of warm ammonia on isatin, but is best prepared by dissolving isatin to saturation in aqueous KOH, evaporating the solution to dryness, dissolving in alcohol, adding a very concentrated solution of ammonium sulphate, filtering, and evaporating to dryness The residue contains amasatin and ammonium isamate, and the latter may be dissolved in alcohol, whence, after addition of HCl, isamic acid crystallises (Laurent, *A Ch* [3] 8, 490) Ruby coloured hexagonal tables or scarlet trimetric laminae. Sl. sol. boiling water, forming a yellow

low solution, v sol hot alcohol, m sol ether. Aqueous HCl dissolves it with violet colour, and it separates again from the solution in violet crystals turned red by water. Boiling dilute acids split it up into isatin and ammonia. Bromine forms 'indelbrome', a yellow substance $C_{12}H_7BrN_2O_2$, insol water.

Salts— NH_4A small needles, or very acute minute rhombs. When strongly heated it gives off water and forms amasatin. Its solution does not ppt salts of Ba, Ca, or Mg, but gives a yellow pp with lead acetate and with $AgNO_3$, and a red pp with $HgCl_2$.—The potassium salt may be boiled without decomposing— AgA .

Di chloro isamic acid $C_8H_4Cl_2N_2O_4$. Formed by adding HCl to a solution of di chloro amasatin in dilute KOH and crystallising the brick red pp from alcohol (L). Bright red elongated hexagonal laminae. More sol alcohol and ether than isamic acid. Forms yellow solutions. Decomposed by distillation. HClAq forms a violet solution, but on boiling it is split up into NH_3 and chloro isatin.

Tetra chloro isamic acid $C_8H_2Cl_4N_2O_4$. By boiling tetra chloro amasatin with alcohol and adding silver nitrate a flocculent precipitate of $C_{12}H_4AgCl_4N_2O_4$ is formed (L).

Di bromo isamic acid $C_8H_4Br_2N_2O_4$. Formed by dissolving di bromo amasatin in KOHAq and neutralising with dilute HCl (Gericke). Red powder, nearly insol water, v sol alcohol and ether. HClAq gives a violet solution. Boiling KOHAq gives bromo isamic acid— KA yellow needles, sl sol water— BaA .

Isatimide $C_8H_5N_2O_3$. Formed by passing dry NH_3 over isatin moistened with alcohol (86 to 100 p c). When absolute alcohol is used, imasatin first crystallises out, and the filtrate deposits isatimide as a yellow crystalline powder. Insol water, scarcely sol boiling alcohol or ether, v sol boiling alcoholic NH_3 . Potash dissolves it with yellow colour and evolution of NH_3 , the liquid then containing isatin.

Isatilm $C_8H_5N_2O_3$ (?) Sometimes formed when dry NH_3 is passed over isatin moistened with alcohol, separating from the alcoholic filtrate from which isatimide has separated (Laurent, *J pr* 35, 121). Yellow amorphous flocks, easily decomposed by KOH.

Di isatin diamide $C_{12}H_7N_2O_6$. Formed by saturating an alcoholic solution of isatin with dry NH_3 , and heating to 100° for 24 hours. The product is filtered boiling, the filtrate containing deoxy imido isatin, while the di isatin diamide and oxydiamidosatin diamide which remain on the filter are separated by treatment with water in which the di isatin diamide is the less soluble (L von Sommaruga, *A* 190, 867, *B* 11, 1082, 12, 980). Pale yellow crystals, sl sol alcohol and water. Nitrous acid forms some di isatin amide.

Salts— $BHCl$ yellow crystalline powder, almost insol cold water— $BHNO_2$, yellow needles— $BHSO_4$, yellow needles. Not decomposed by heating with water at 100° — $BHCrO_4$, orange powder.

Di isatin amide $C_{12}H_{11}N_2O_5$, [252°] Formed by warming di isatin diamide with dilute KOH and ppg with HCl (Sommaruga, *M* 1, 579). Small yellowish needles (from alcohol). Scarcely sol. water, v. sol alcohol— $C_{12}H_{11}(NH_2)_2N_2O_5$.

silvery scales— $C_{12}H_9KN_2O_5$, 14aq silvery plates or needles.

Dihydrate $C_{12}H_{11}N_2O_5$, [213°] Formed by treating di isatin amide or di isatin diamide with sodium amalgam (Sommaruga, *A* 194, 88). Slender needles (from alcohol), scarcely sol water and ether. Oxidised by boiling with HgO or aqueous $FeCl_3$ to di isatin amide. Not affected by aqueous KOH (S G 127) at 100° — $NaC_{12}H_{11}N_2O_5$, long colourless needles, v sol hot water— $KC_{12}H_{11}N_2O_5$, broad needles with silky lustre.

Oxy diamido di isatin diamide $C_{12}H_9N_4O_7$, *Oxydimidodiamidosatin* [295°–300°] Formed as above (S). Large colourless needles. Sol water, v e sol alcohol. Dissolves readily in acids, and is not reppd from these solutions by NH_3 . Boiling with water and sodium amalgam converts it into di amido di hydrindic acid. The solutions of its salts show intense red fluorescence— $BHNO_2$, granules, v sol hot water— $BHSO_4$, prisms.

Di amido di hydrindic acid $C_{12}H_9N_4O_7$, [217°] Formed as above. Crystalline granules. Decomposed on fusion. V sol hot water. Oxidised by chromic acid mixture to 'di imido di hydrindin dicarboxylic' acid $C_{12}H_7N_4O_9$, which crystallises from hot water in needles.

Deoxyimido disatin $C_{12}H_9N_2O_5$, [210°] Formed as mentioned under disatin diamide (S), and purified by solution in aqueous KOH and ppg by an acid. Yellowish powder. Decomposed by fusion. V sol alcohol, hot water, and alkalis.

Oxy amido hydro-isatin, so called, $C_{12}H_9N_2O_6$. Formed by treating the preceding body with sodium amalgam or by heating it with KOHAq in sealed tubes at 100° (S). Yellowish amorphous powder, insol water, v sol alcohol. Decomposes at 188° without fusion.

ISATIN CARBOXYLIC ACID *Anilide* $C_{12}H_9NO_4$, i.e. $C_{12}H_9NO$ CONPhH, probably CO

$C_6H_5 < \begin{smallmatrix} \diagup CO \\ \diagdown NCONPhH \end{smallmatrix}$ *Carbanilido isatin* [180°–

183°] Formed by heating isatin for 8 hours at 130° with phenyl cyanate (Gumpert, *J pr* [2] 32, 283). Crystallises from benzene in canary yellow needles. Sol alcohol, ether, and glacial acetic acid.

Reactions—1 When fused it forms isatin and phenyl cyanate (compare the formation of isatin from acetyl isatin).—2 With H_2SO_4 and crude benzene it gives the indophenol reaction. 3 Dissolves in warm dilute KOH forming potassium carbanilido isatate. HCl throws down sparingly soluble carbanilido isatic acid, $NPhH \cdot CO \cdot NH \cdot C_6H_5 \cdot CO \cdot CO \cdot H$, [170°–180°, with decomposition]. The acid is very stable, but by heating at 110° for a long time it loses H_2O , changing back to carbanilido isatin. Carbanilido isatic acid, when heated with ethyl or methyl alcohol, splits off CO_2 and H_2O forming $C_{12}H_{11}N_2O_5$, [175°], and $C_{12}H_{11}N_2O_5$, [197°], respectively.—4 Converted by amines into amides of carbanilido-isatic acid, thus alcoholic NH_3 at 100° forms $NPhH \cdot CO \cdot NH \cdot C_6H_5 \cdot CO \cdot CO \cdot NH_2$, [229°], crystallising from alcohol in needles. It dissolves in dilute NaOH, but is reppd unaltered by HCl. When dissolved in glacial acetic acid it is converted by N_2O_5 into an indifferent body, $C_{12}H_{11}N_2O_9$, [270°].—5. NEt_3 ,

forms $\text{NPhH CO NH C}_6\text{H}_5$, CO CO NEtH [210°] It forms needles (from alcohol) and is a weak acid — 6 *Phenyl hydrazine* forms $\text{NPhH CO NH C}_6\text{H}_5$, $\text{CO CON}(\text{NH}_2)\text{Ph}$ [193°] — 7 *Hydroxylamine* forms

$\text{NPhH CO NH C}_6\text{H}_5$, CO CO NH OH [225°]

ISATIN DIHYDRIDE ν DIOXINDOLE

ISATIN INDOGENINE ν INDIGO

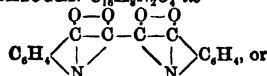
ISATIN SULPHONIC ACID $\text{C}_6\text{H}_4\text{NSO}_3$, $\pm e$

$\text{SO}_3\text{H C}_6\text{H}_4 \begin{smallmatrix} \text{CO} \\ \diagup \text{N} \diagdown \end{smallmatrix} \text{CO OH}$ *Isatosulphuric acid*

Preparation—Commercial indigo carmine (18 pts) is stirred up into a thin paste with water (18 pts), H_2SO_4 (1½ pts) is added, and the mixture heated to boiling while $\text{K}_2\text{Cr}_2\text{O}_7$ (1 pt) is gradually thrown in, as long as decolourisation ensues. The hot solution is filtered and KNO_3 added, which facilitates the deposition of potassium isatin sulphonate. The K salt is mixed with resinous matter, which can be got rid of by dissolving in hot baryta water, ppg excess of baryta by CO_2 , and then ppg by K_2SO_4 (G & A Schlieper, A 120, 1). Yellow radio crystalline mass (containing 2aq) Insol ether and benzene, m sol alcohol, v sol water. Powerful acid, separating HCl from its salts. Not decomposed by H_2SO_4 or HNO_3 , even on boiling. Aqua regia slowly forms tetra-chloro quinone. Alcoholic NH_3 forms dark brown bodies. Hydric iodide does not reduce it. Ammonium sulphide forms $\text{SO}_3\text{H C}_6\text{H}_4 \begin{smallmatrix} \text{CH(OH)} \\ \diagup \text{N} \diagdown \end{smallmatrix} \text{COH}$ Hot potash forms sulpho isatic acid.

Salts— NaA 2aq deep-red tables, ppg from its aqueous solution by NaCl — KA aq small golden needles S 5 in the cold. Insol alcohol. Ppd by KNO_3 from its aqueous solution— NH_4A aq deep yellow needles, v sol water, sl sol NH_4Cl — BaA 4aq scarlet powder, insol alcohol, sl sol water— CaA 2aq small golden needles, m sol water— AgA aq. yellow needles, sl sol water.

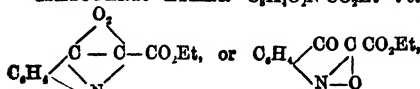
DIISATOGEN $\text{C}_6\text{H}_4\text{N}_2\text{O}_4$, $\pm e$



$\text{C}_6\text{H}_4 \begin{smallmatrix} \text{CO CO CO CO} \\ \diagup \text{N} \diagdown \end{smallmatrix} \text{C}_6\text{H}_4$ (Michael, J pr [2]

85, 256) Prepared by the action of fuming H_2SO_4 on di-*o*-nitro di-phenyl-diacetylene (Bayer, B 15, 52). Red needles. Sol nitro benzene, sl sol chloroform, insol alcohol and ether. With H_2SO_4 and FeSO_4 it gives indom. It is very readily reduced to indigo by cold NH_4HS , by zinc-dust, and NH_4NaOH or AcOH , by glucose, and alkalis, &c. It combines with ammonium bisulphite.

ISATOGENIC ETHER $\text{C}_6\text{H}_4\text{O}_2\text{N CO}_2\text{Et}$, $\pm e$.



or $(\text{CO}_2\text{Et CO CO C}_6\text{H}_4)_2\text{N}_2$ (Michael, J pr [2] 85, 255) [115°] Formed by an isomeric change from *o*-nitro phenyl-propionic ether by the action of cold H_2SO_4 (Bayer, B 14, 1741, 15, 780). Yellow needles. By most reducing agents it is reduced to indoxyl ether, but ferrous salts give indoxanthic ether (Bayer, B 15, 780). Isato-

genic acid is perhaps formed by shaking *o*-nitrophenyl propionic acid with H_2SO_4 , but if so, it is decomposed on diluting with water into CO_2 and isatin.

ISATOIC ACID $\text{C}_6\text{H}_4\text{NO}_3$, $\pm e$ $\text{C}_6\text{H}_4 \begin{smallmatrix} \text{CO} \\ \diagup \text{N} \diagdown \end{smallmatrix} \text{CO}_2\text{H}$

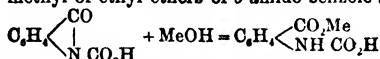
Anthranil carboxylic acid [230°] S (acetone) 4 & 4 at 55°

Formation—1 By the oxidation of isatin by CrO_3 in HOAc (Kolbe, J pr [2] 80, 469).—2 By heating anthranil with chloro formic ether at 30° (Friedlander & Wleügel, B 16, 2227).—3 By boiling *o*-amido benzoic acid with excess of chloro formic ether (Niementowski & Rozanski, B 22, 1672).

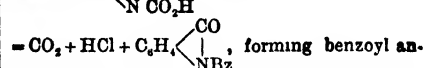
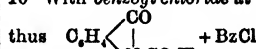
Preparation—(H Kolbe, J pr [2] 30, 469) Isatin (100 g) is powdered and mixed with glacial acetic acid (600 g). A solution of CrO_3 (200 g) in glacial acetic acid (600 g) is slowly added, the temperature being kept below 50°. The flask is left for 12 hours in cold water, and then in water at 50°, and finally at 60°. Isatoic acid then begins to separate as a yellow powder, the separation is completed by pouring into 500 c.c. cold dilute H_2SO_4 . Yield 72 per cent.

Properties—Nearly rectangular prisms, which are yellow (through some impurity) when prepared from isatin. Decomposed at 230°. Sparingly soluble in alcohol, ether, and glacial acetic acid, rather more soluble in chloroform and benzene.

Reactions—1 Boiled with water it forms *o*-amido benzoic (anthranilic) acid $\text{C}_6\text{H}_4\text{NO CO}_2\text{H} + \text{H}_2\text{O} = \text{C}_6\text{H}_4(\text{NH})\text{CO}_2\text{H} + \text{CO}_2$.—2 Conc HCl , H_2SO_4 , and dilute HNO_3 behave in the same way.—3 Gaseous HCl passed into an alcoholic solution forms the hydrochloride of *o*-amido benzoic ether.—4 Aqueous baryta forms, on warming, *o*-amido benzoic acid.—5 *Ammonia* (aqueous or dry) forms *o*-amido benzamide.—6 *Aniline* at 60° forms the anilide of *o*-amido benzoic acid.—7 HNO_3 (SG 148) forms nitro isatoic acid.—8 *Nitrous acid gas* passed into water, containing isatoic acid suspended, forms (a) nitro salicylic acid. Anthranilic acid is doubtless first formed, and this is then nitrated and exchanges NH_2 for OH .—9 Heated with methyl or ethyl alcohol it unites, forming crystalline compounds, carboxylic acids of the methyl or ethyl ethers of *o*-amido benzoic acid.



The acid $\text{CO}_2\text{H NH C}_6\text{H}_4\text{CO}_2\text{Me}$ crystallises in needles [176°], while the corresponding $\text{CO}_2\text{H NH C}_6\text{H}_4\text{CO}_2\text{Et}$ forms needles [126°].—10 With benzoyl chloride at 210° it partly reacts



thranyl [123°].—11 *Bromine* forms bromo isatoic acid, which gives with HCl (β) bromo anthranilic acid [208°].—12 It dissolves in cold dilute NaOH with a blue fluorescence, but on standing the fluorescence vanishes and the solution then contains anthranilic acid (Friedlander & Wleügel, B 16, 2227).—13 *Phenol* at 180° gives phenyl

o-amido benzoate crystallising in needles [70°] (G Schmidt & E v Meyer, *J pr* [2] 36, 370) — 14 Long boiling with glacial *acetic acid* forms an amorphous compound $C_{10}H_7N_3O_6$, v sl sol ordinary solvents, which yields *o* amido benzoic acid when heated with HCl or dilute H_2SO_4 (G Schmidt, *J pr* [2] 36, 380) — 15 Heating with Ac_2O gives acetyl *o* amido benzoic acid [180°] (S) — 16 With *hydroxylamine* it forms *o* amido-benzoyl hydroxylamine (E v Meyer & T Bell-

mann, *J pr* [2] 33, 19) $C_6H_4 \begin{smallmatrix} \diagup CO \\ | \\ N \\ \diagdown CO_2H \end{smallmatrix} + H_2NOH$
 $= C_6H_4(NH_2)CO NH(OH) + CO_2$ — 17 With *phenyl hydrazine* it forms *o* amido benzoyl phenyl-

hydrazine $C_6H_4 \begin{smallmatrix} \diagup CO \\ | \\ N \\ \diagdown CO_2H \end{smallmatrix} + PhNH NH_2$
 $= C_6H_4(NH_2)CO NPh NH_2 + CO_2$ — 18 It does not react with boiling *di-methyl aniline* — 19 *Pyrocatechin* at 180° reacts thus (M & B).

$C_6H_4(OH)_2 + C_6H_4 \begin{smallmatrix} \diagup CO \\ | \\ N \\ \diagdown CO_2H \end{smallmatrix}$
 $= CO_2 + C_6H_4(NH_2)CO O C_6H_4OH$, forming *o* oxyphenyl *o* amido benzoate — 20 Boiled with *formic acid* it forms formyl *o* amido benzoic acid and a weak base, $C_8H_7N_3O_6$. This crystallises in rhombohedra (from alcohol) It melts at [280°], with decomposition Sparingly soluble in water, alcohol, benzene, and chloroform S (ether) 03 at 15°, S (alcohol) 11 at 19° Readily soluble in NaOH, Aq, forming a crystal line salt Its hydrochloride forms prisms, but is decomposed by water Heated with conc HCl at 140° it forms CO_2 , formic acid, and the hydrochloride of anthranilic acid (M & B) — 21 By treatment with PCl_5 in presence of $POCl_3$ a product is got whence methyl alcohol and ethyl alcohol respectively form crystalline products The former gives $C_{10}H_7N_3O_6$ [210°], small needles The latter gives $C_8H_7N_3O_6$ [170°], needles With conc HCl at 140° the latter gives EtCl and anthranilic acid (M & B) — 22 *Blackening powder* suspended in chloroform changes some of the isatoic acid into an isomer [240°], soluble in alcohol, acetone, and benzene It dissolves in NH_3 , Aq without forming anthranilamide (unlike isatoic acid), but when HCl is added to the solution an acid [260°] is ppd (M & B) — 23 Heated with glacial *acetic acid* and *bromine* (1 mol) it forms bromo isatoic acid, but with more bromine CO_2 is evolved, and di-, tri-, and tetra-bromo amido benzoic acids are got.

Chloro-isatoic acid $C_6H_4Cl \begin{smallmatrix} \diagup CO \\ | \\ N \\ \diagdown CO_2H \end{smallmatrix}$

[α 268°] From chloro isatin (10 g), CrO_3 (20 g) and glacial *acetic acid* (120 g) as described under bromo isatin (Dorsch, *J pr* [2] 33, 49)

Properties — Pearly plates (from alcohol-ether) Sl sol alcohol, acetone, and glacial *acetic acid*, insol benzene, chloroform, ether, and water

Reactions — 1 Conc HCl forms chloro-*o*-amido-benzoic acid [204°] — 2 Hot NH_3 , Aq forms the amide of chloro amido benzoic acid

Di-chloro-isatoic acid $C_6H_2Cl_2 \begin{smallmatrix} \diagup CO \\ | \\ N \\ \diagdown CO_2H \end{smallmatrix}$

[256°]

Formation. — By oxidation of di chloro-isatin.

Properties — Yellow prisms (from alcohol-acetone) Melts at 254°-256° with decomposition V sol acetone and glacial *acetic acid*, sol alcohol and chloroform, v sl sol ether and benzene Boiled with water it partly changes to di chloro *o* amido benzoic acid, as shown by its violet fluorescence (Dorsch, *J pr* [2] 33, 51).

Reactions — 1 Conc HCl slowly converts it, on boiling, into di chloro amido benzoic acid [224°] — 2 With NH_3 , Aq it gives di-chloro amido benzamide

Bromo isatoic acid $C_6H_4Br \begin{smallmatrix} \diagup CO \\ | \\ N \\ \diagdown CO_2H \end{smallmatrix}$

[α 275°]

Formation — 1 By oxidising bromo isatin. 2 From Br and isatoic acid suspended in glacial *acetic acid* at 90° (R Dorsch, *J pr* [2] 33, 32)

Properties — Pearly plates (from a mixture of alcohol and acetone) Sol acetone Sl sol alcohol and glacial *acetic acid* Insol water, chloroform, ether, and benzene

Reactions — 1 With boiling conc hydrochloric acid it forms (β) bromo amido benzoic acid $C_6H_4Br(NH_2)CO_2H$ — 2 With hot NH_3 , Aq it gives bromo amido benzamide

Di bromo isatoic acid $C_6H_2Br_2 \begin{smallmatrix} \diagup CO \\ | \\ N \\ \diagdown CO_2H \end{smallmatrix}$

[255°] Obtained by oxidising di bromo isatin (10 g) by CrO_3 (10 g) in presence of glacial *acetic acid* (60 g) as described under bromo isatin

Properties — Flesh coloured prisms (from alcohol-acetone) Sol glacial *acetic acid* and acetone, sl sol alcohol, chloroform, and benzene, hardly sol ether, insol water Much more stable than bromo isatoic acid

Reactions — 1 Boiled for a long time with conc HCl it gives off CO_2 , leaving di bromo *o*-amido benzoic acid — 2 It does not dissolve in NH_3 , Aq But if heated with it at 100° for a long time it forms di bromo amido benzamide $C_6H_2Br_2(NH_2)CO NH_2$ [197°]

Nitro-isatoic acid $C_6H_4(NO_2)NO CO_2H$ [220°-230°] Formed by the action of HNO_3 (S G 148) on isatoic acid Insol water and ether, sl sol alcohol Crystallises from equal parts of alcohol and acetone in pearly plates It resembles isatoic acid, splitting up readily into CO_2 and nitro-*o* amido benzoic acid, when boiled with HCl or even with water Nitro-isatoic acid is converted by aqueous NH_3 into nitro amido benzamide By Sn and HCl it is reduced to di-amido benzoic acid, CO_2 escaping Bromine (in glacial *acetic acid*) converts it into mono-, di-, and tri-nitro-amido benzoic acids (q v)

Methyl-isatoic acid $C_6H_4Me \begin{smallmatrix} \diagup CO \\ | \\ N \\ \diagdown CO_2H \end{smallmatrix}$

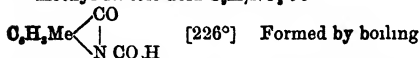
Preparation — Methyl-isatin (10 g) is mixed with glacial *acetic acid* (60 g) and cooled well while a mixture of CrO_3 (21 g) in glacial *acetic acid* (60 g) is slowly added After 12 hours at 0° the mixture is heated for 2 hours at 40° and then to 70° It is poured into water containing H_2SO_4 , and the yellow powder crystallised from absolute alcohol (Panaotović, *J pr* [2] 31, 123, 33, 58)

Properties — Pale-yellow crystalline plates

(from boiling alcohol) or trimetric columns (from acetone) At 245° it suddenly increases in bulk, but it melts above 800°, giving off CO_2 . V sl sol. water, readily sol boiling alcohol, ether, benzene, and chloroform

Reactions — 1 Is more stable than isatoic acid, not being decomposed by dilute mineral acids. — 2 HNO_3 (S G 1 48) dissolves it in the cold After 12 hours water is slowly added, and the pp crystallised from benzene It forms pale yellow trimetric plates of nitro methyl-isatoic acid [175°] This acid is reduced by Sn and HCl to a di-amido-toluic acid — 3 Warmed with HCl (S G 1 2), CO_2 is given off, and the hydrochloride of amido p toluic acid [207°] formed — 4 Warmed with NH_4Aq it gives the amide of amido toluic acid [178°] — 5 With *aniline* in alcohol it forms the corresponding anilide [243°] — 6 With *phenyl hydrazine* it forms the corresponding hydrazone — 7 With MeOH at 180° it forms methylic amido-toluate

Methyl-isatoic acid $\text{C}_8\text{H}_7\text{NO}_4$, \pm



an amido-toluic acid $\text{C}_8\text{H}_7\text{Me}(\text{NH}_2)\text{CO}_2\text{H}$ [5° or 8? 1 2] with chloro formic ether (Niemen towsky a Rozansky, B 22, 1675) Needles or scales, v sl sol benzene and ether, sl sol alcohol Decomposed on fusion Acids and alkalis decompose it into CO_2 and amido toluic acid [177°]

Isomeride of isatoic acid ψ ANTHROXANIC ACID

ISATYDE $\text{C}_{10}\text{H}_7\text{N}_2\text{O}_4$ Produced by the reduction of isatin by zinc and dilute H_2SO_4 by alcoholic ammonium sulphide, or by zinc dust and HOAc (Laurent, A Ch [3] 3, 382, A 72, 285, Erdmann, J pr 24, 15, 25, 438, Baeyer, B 12, 1309) Formed also by atmospheric oxidation of an aqueous solution of dioxindole (Baeyer a Knop, A 140, 10) White powder, with slightly greyish tint, almost insol water, v sl sol boiling alcohol and ether Separates from boiling alcohol in minute scales Decomposed by heat, becoming at first violet-brown Alcoholic potash forms isatoic acid, indin, and other products

Di-chloro-isatide $\text{C}_8\text{H}_5\text{Cl}_2\text{N}_2\text{O}_4$ Formed by the action of ammonium sulphide on chloro-isatin White powder, crystallisable, insol cold, v sl sol hot, water, m sol boiling alcohol, sol. hot aqueous potassium sulphide At 180° it is resolved into chloro-isatin and chloro-indin Boiling aqueous or alcoholic potash forms potassium chloro isatate and the salt $\text{C}_8\text{H}_5\text{Cl}(\text{NH}_2)\text{CH}(\text{OH})\text{CO}_2\text{K}$

Tetra-chloro-isatide $\text{C}_8\text{H}_3\text{Cl}_4\text{N}_2\text{O}_4$ Produced by the action of ammonium sulphide on di-chloro-isatin White powder, insol water Decomposed by heat into di-chloro isatin and di-chloro indin Alcoholic potash forms di chloro isatin and di chloro-o amido-o-oxy phenyl acetic (di-chloro hydrindic) acid

Tetra-bromo-isatide $\text{C}_8\text{H}_3\text{Br}_4\text{N}_2\text{O}_4$ From di-bromo isatin and ammonium sulphide Resolved by heat into di-bromo-isatin and di-bromo-indin

Thio-isatide $\text{C}_8\text{H}_7\text{N}_2\text{O}_3\text{S}$ Formed by slowly adding alcoholic potash to an alcoholic solution of di thio isatide (Laurent, A Ch [3] 8, 463)

White crystalline powder May be crystallised as minute rectangular scales from hot alcohol Insol water, v sl sol boiling alcohol and ether. Cold potash forms indin and other products Hot potash forms the hydride of indin

Di-thio-isatide $\text{C}_8\text{H}_7\text{N}_2\text{O}_2\text{S}_2$ When H_2S is passed into a conc alcoholic solution of isatin the liquid becomes pale yellow and on cooling deposits crystals of sulphur and of isatide The filtrate when mixed with water deposits di thio-isatide (Laurent) Yellowish grey amorphous powder Decomposed by heat Insol boiling water, v sl warm alcohol and ether Ammonium bisulphite converts it into 'ammonium sulphisatamite' $\text{NH}_4\text{C}_8\text{H}_7\text{NSO}_4$, which crystallises in large pale yellow tables v sol water, m sol alcohol

Di-bromo-di-thio-isatide $\text{C}_8\text{H}_5\text{Br}_2\text{N}_2\text{O}_3\text{S}_2$ Formed, together with di bromo tri thio isatide, when H_2S is passed into a boiling alcoholic solution of bromo isatin (Gericke, Z 1865, 595). Yellowish white powder, insol hot water, sol hot alcohol and ether

Di-bromo-tri-thio-isatide $\text{C}_8\text{H}_5\text{Br}_3\text{N}_2\text{O}_3\text{S}_3$ Formed as above Yellowish white powder

Isatane $\text{C}_{10}\text{H}_{11}\text{N}_2\text{O}_4$ Formed, as a white pp, when di-thio isatide is boiled with a solution of ammonium bisulphite (Laurent, J pr 28, 346) Formed also by the action of sodium amalgam on an acid solution of isatin (Knop, J pr 97, 65) Small white cubes (from ether or hot alcohol), insol water Decomposed by hot alcoholic potash into dioxindole and indirubin Its alcoholic solution gives with ammoniacal AgNO_3 a white pp of $\text{Ag}_2\text{C}_{10}\text{H}_{11}\text{N}_2\text{O}_4$

ISETHIONIC ACID $\text{C}_8\text{H}_7\text{SO}_4$, \pm

$\text{HOCH}_2\text{CH}_2\text{SO}_3\text{H}$ *Oxy ethane sulphonic acid*
Sulphonic acid of ethyl alcohol Mol w 126

Formation — 1 Discovered by Magnus in 1833 (P 27, 378, A 6, 163) as a product of the action of SO_2 on alcohol or ether Hence it is found among the residues in the preparation of ether — 2 By boiling ethionic acid with water (Magnus, A 32, 251) — 3 By the action of SO_2 on barium ethyl sulphate (Meves, A 143, 196) — 4 By the action of nitrous acid on taurine $\text{NH}_2\text{CH}_2\text{CH}_2\text{SO}_3\text{H}$ (W Gibbs, Am S [2] 25, 30) — 5 By heating chloro ethyl alcohol (chlorhydrin of glycol) with aqueous Na_2SO_3 at 175° (Collmann, A 148, 101) — 6 By heating ethylene oxide with aqueous KHSO_3 at 100° (Lrilenmeyer a Darmstadter, Z 1868, 342) — 7 Probably formed by boiling ethylene bromide with aqueous Na_2SO_3 (James, C J 43, 44) — 8 E J oxidising thio glycol $\text{HOCH}_2\text{CH}_2\text{SH}$ with nitric acid (Carius, A 124, 260)

Preparation — SO_2 is added, with shaking, to an equal weight of ether at 0° As soon as a sample mixed with water gives a heavy oil the whole is poured into water and the ethyl sulphate washed with water till neutral, dried over H_2SO_4 , treated with its own weight of SO_2 , and then poured into water The two aqueous liquids are boiled for a long time to decompose ethionic acid, and then neutralised by baric carbonate, on evaporating baric isethionate is got (R Hübner, A 223, 212)

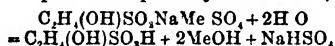
Properties — Viscid, strongly acid syrup, which gradually dries up to a deliquescent radio crystalline mass Is not decomposed at 150°, but blackens at a higher temperature Its

salts are not decomposed by boiling with water, and may even be heated to 200°, above which temperature, however, they lose water and change to di isethionates. Potash fusion gives potassium carbonate, oxalate, sulphate, and sulphite. Berthelot (*Z* 1869, 682) obtained acetylene, K_2SO_4 , and hydrogen PCl_5 yields $CH_2ClCH_2SO_2Cl$, which is converted by heating with aqueous ammonia into taurine (Kolbe, *A* 112, 241). Chromic acid oxidises isethionio acid to sulpho acetic acid $CO_2HCH_2SO_3H$.

Salts— NH_4A' octahedra [130°] (Strecker), [185°] (Seyberth, *B* 7, 391). At 235° it changes to ammonium di isethionate (Carl, *B* 12, 160.)— KA' rhomboidal prisms, melting between 300° and 350° with loss of weight. May be crystallised from alcohol— BaA' , transparent six sided plates [32°]. V sol water S (60 p alcohol) 6 at 14°— CaA' , 2aq pale green prisms— AgA' very hygroscopic pearly needles.

Double salts with Et_2SO , (Engelcke, *A* 218, 270). From NaA' , H_2SO_4 , and alcohol, and conversion into barium salt BaA'_2Et_2SO (?)— $NaATf, SO_4$.

Double salt with Me_2SO ,— $NaA'Me_2SO$, Silky monoclinic tables (from alcohol), very hygroscopic. Decomposed by water at 80°, thus



Ethyl derivative $EtOCH_2CH_2SO_3H$ Sulphonic acid of ether SG ²¹ 1359. The sodium salt is obtained by the action of chloro ethane sulphochloride on excess of $NaOEt$ (R Hubner, *A* 223, 218). Also from $CH_2ClCH_2SO_3Na$ and $NaOEt$. The free acid is got by boiling with water the product obtained by acting with H_2S on the (molecular?) compound of its lead salt with lead ethyl sulphate. It is a syrup.

Salts— NaA' Plates S (alcohol) 27 at 15°— NaA' 3aq Columns— BaA' 3aq— ZnA' 6aq Hygroscopic plates— CaA' 6aq.

Double compounds with the salts of ethyl sulphuric acid $EtSO_3H$. Got by adding H_2SO_4 and alcohol to $EtOCH_2CH_2SO_3Na$, filtering from Na_2SO_4 , and neutralising with a metallic carbonate. The general formula is $C_2H_5SO_3H.C_2H_5OCH_2CH_2SO_3H$ or $C_2H_5S_2O_6H$, e.g. BaA' 3aq Silky scales— $(NH_4)A'_2$, PbA'_2 , NaA'_2 , AgA'_2 , ZnA'_2 , CaA'_2 . These salts are very soluble in water. The free acid splits up on boiling with water, in the following manner $C_2H_5SO_3H.EtOCH_2CH_2SO_3H + H_2O = EtOC_2H_5SO_3H + H_2SO_4 + EtOH$.

Ethyl ether $HOCH_2CH_2CH_2SO_3Et$ From AgA' and EtI (Stempnewsky, *J R* 1892, 95).

Ethyl derivative of the ethyl ether $EtOCH_2CH_2SO_3Et$ SG ¹² 1168 (impure). From $CH_2ClCH_2SO_3Cl$ and $NaOEt$ in ether. Not obtained quite pure.

Benzoyl derivative $BzOCH_2CH_2SO_3H$ From potassium isethionate and $BzCl$ (Engelhardt & Latschnoff, *Z* 1868, 235)— KA' Leaflets, v sol boiling water, sol hot alcohol— BaA' 3aq Large thin tables, m sol cold water, sol boiling alcohol.

Chloride $HOCH_2CH_2SO_3Cl$ Probably formed, together with $CH_2ClCH_2SO_3H$ and $CH_2ClCH_2OSO_3Cl$, by the action of SO_2 on ethyl chloride, or of $ClSO_3H$ on ethylene (Furgold, *B* 6, 504). Excess of SO_2 converts it into the

chloride of ethionio acid (Claesson, *J pr* [2] 19, 253).

Di-isethionio acid $O(CH_2CH_2SO_3H)_2$. **Di sulphonic acid of ether**. The ammonium salt of this acid is obtained by heating ammonium isethionate to 210° (Carl, *B* 12, 1604)— $(NH_4)A''$ [198°] Slender leaflets or scales, v e sol, water— BaA'' 3aq prismatic tables. Formed by heating barium isethionate to 200° (Carl, *B* 14, 65).

ISINGLASS & PROTEIDS, Appendix C.

ISO- Compounds whose names begin with iso are usually described either under the name to which iso- has been prefixed or else under their systematic names as described in the *Introduction to Articles Relating to Organic Chemistry*, vol 1.

ISOMERISM Even a superficial reader of chemical literature will soon become aware that the term *isomeric* and the kindred expressions *allotropic*, *metameric*, and *polymeric* are by no means always used in consistent senses, and he will have considerable difficulty in clearly realising their exact and relative import, it, therefore, appears desirable to discuss the meanings of these terms, especially from the historical side, and as far as possible to define the sense in which they are severally applicable. The following extracts from the article *Isomerism* in the first edition of this Dictionary, vol iii p 415, 1865, by J A Wanklyn, serve to show what views were held at the very outset of the period when the investigation of isomeric substances began largely to engage the attention of chemists—

'Isomerism.—This term is derived from *isos* equal, and *meros* a part, and its employment by chemists is an expression of the fact that very different chemical compounds have sometimes identically the same ultimate composition. Two or more different bodies which are composed of the same elements and of the same proportions of these elements (i.e. which have the same percentage composition) are said to be *isomeric*. Isomerism is sometimes used in a narrower sense, being made to signify equality of molecular weight, as well as identity in percentage composition. When the compounds have the same percentage composition but different molecular weights, the term *polymeric* is employed. Thus there are the terms *isomeric* (in its wide sense), signifying that the different bodies have the same percentage composition, *Polymeric*, signifying that these different bodies have the same percentage composition, but different molecular weights, *Isomeric* (in its restricted sense), sometimes called *Metameric*, signifying that the bodies have the same percentage composition, and likewise the same molecular weight.'

As examples, Wanklyn then cites butyric acid, ethylic acetate, aldehyde, and ethylenic oxide as *isomeric* compounds, using the term in its widest sense of these butyric acid and ethylic acetate are said to be *polymeric* with aldehyde and ethylenic oxide, butyric acid being *isomeric* (in the restricted sense) or *metameric* with ethylic acetate, aldehyde and ethylenic oxide being also metameric compounds. Subsequently, throughout his article, Wanklyn uses the term *isomeric* in its wide sense, substituting the term *metameric* for *isomeric* used in its restricted sense, thus he speaks of methyl, the simplest alcohol radiole, as metameric with ethyl hydride (his article was written at a time when Schorlemmer's investigation was not fully recognised as affording proof of their identity), he points out that several metameric hexanes are possible, and even quotes 'as a very remarkable

example of metamerism' the different varieties of tartaric acid and racemic acid

The term *allotropy* is made use of by Wanklyn in an unusually wide sense. Thus he says—

'Closely related to the term isomerism is the term *allotropy*. Both of them have reference to the same substantial fact, viz that different substances have sometimes the same ultimate composition, but they differ in their manner of stating it. Isomeric and allotropic are in fact complementary terms, "isomeric" being employed to predicate identity of composition between different bodies, whilst "allotropic" expresses difference between bodies of identical composition. Such being the force of these words, there is a certain propriety in their usage. Thus, whilst it is correct to say "butyric acid and acetic ether are isomeric," it should be "there are *allotropic* bodies of the formula $C_4H_8O_2$." The same reason which enjoins the use of allotropic in this case prescribes it in the instance of single elements: thus, for example, we read of "allotropic kinds of sulphur," but never of isomeric kinds. It is worthy of remark that cases of isomerism occurring in inorganic chemistry are usually described by employing the word allotropy or allotropic, while the reverse obtains in organic chemistry. This may be partly ascribed to there being always a very wide difference—or else no difference at all—in the composition of any definite inorganic substances, and hence the fact of identity or non identity of composition being so easily ascertainable. It is implied in the form of expression, whilst the fact of difference of properties alone needs to be made the subject of formal predication. Among organic bodies, on the other hand, it continually happens that the differences of composition are quite decided, and yet so very minute as to tax the utmost powers of chemical analysis for their recognition, and hence the superior dignity which the mere affirmation of identity of ultimate composition acquires in the organic department of the science. The principal examples of allotropy or isomerism, &c. of the co-existence of identical ultimate composition with difference of properties will now be considered.

He then cites the olefines as examples of polymerism, and afterwards discusses numerous cases of metamerism, several of which were referred to above.

Under the heading *Isomers among Inorganic Substances* the following interesting passages occur at the conclusion of the article—

'As before remarked, the instances of inorganic isomerism are usually called instances of allotropy—*isomeric substances and allotropic substances* being nearly equivalent expressions. The elementary substances themselves offer many examples of isomerism. Ozone and oxygen are isomeric bodies. Experiment has shown that the molecular formula for ozone is higher than that for oxygen, but how much higher remains an open question. Sulphur, phosphorus, carbon, and many other elements present somewhat similar examples of allotropy or isomerism. Inorganic compounds, such as the various forms of silicic acid, of sesquioxide of iron, of sesquioxide of chromium, of alumina, must be classed among substances affording examples of isomerism. The explanation of the existence of isomerism will have become sufficiently clear from the course which has been followed in describing the different examples of it. "It is of consequence how the atoms of a compound are arranged, as well as what kind of atoms they are, and hence there may be very many totally different substances composed of the same ultimate atoms. This is in fact the whole philosophy of isomerism."

The definitions given in Kekulé's *Lehrbuch* (1867) are substantially the same as those adopted by Wanklyn, but he specially draws attention to the existence of compounds *isomeric in a restricted sense*, which, according to the state of knowledge of the time, were to be represented by the same rational formulae, although they either were possessed of different properties—as in the case of the C_4H_8 hydrocarbons and of maleic and fumaric acids, or they were in all essential respects chemically identical but physically different—such as the tartaric acids, mucic and saccharic acids, &c. The existence of compounds such as these latter, in fact, gave rise to the re-

cognition of a distinct kind of isomerism, termed *physical isomerism*.

In the latest edition of Watts' *Fownes* by Tilden (1886), polymerism is included under isomerism, but compounds of the same molecular weight are sub-divided into (1) *metameric bodies*, namely, those which exhibit dissimilar transformations under similar circumstances propionic acid, methylic acetate, and ethylic formate are quoted as examples, and (2) *isomeric bodies, strictly so-called*, namely, those which exhibit the same or closely similar decompositions and transformations when subjected to the action of the same reagents, such as the C_4H_{10} hydrocarbons, the glucoses, the tartaric acids, &c.

It will be noted how incompatible are the definitions given by Wanklyn and in *Fownes* of the term *metameric*, it has, however, undoubtedly been customary of late years to employ the term *metameric* in the sense indicated in *Fownes*.

In McGowan's translation of Berntsen's *Organic Chemistry* (1889), the most modern book of its kind, polymerism is not reckoned under isomerism, but the definition given of metamerism is on the whole more in agreement with that quoted from Watts' *Fownes*, after it has been explained that ethers such as methyl amyl ether, ethyl butyl ether, and dipropyl ether are isomeric, we read—

'Such isomerism which depends upon the grouping together by a polyvalent element of alcohol radicals which are individually unequal, but the sum of whose elements taken together are equal, is called *metamerism*. One of the alcohol radicals may here be replaced by hydrogen. Alcohols and ethers containing an equal number of carbon atoms are therefore *metameric*.'

We further learn that the isomerism of the higher paraffins, since it is based upon the dissimilarity of the carbon chains, is often termed *chain isomerism*, that the isomerism between ethylene and ethylidene chlorides, or between primary and secondary propyl alcohols, as it depends upon the difference in position of the substituting halogen or hydroxy in the same carbon chain, is termed *isomerism of place or position*, and that there is the third kind of isomerism, viz *metamerism*. But obviously two different kinds of relationship are thus included under metamerism that of position isomerism, which obtains among the ethers themselves, which are necessarily all compounds of one primary type, and that which obtains between the typically different 'equi molecular' alcohols and ethers, or true metamerism, if the Watts *Fownes* definition be adopted.

If we consider the origin of the four terms under consideration we find that they were all devised by Berzelius. The term *isomeric* is proposed in his *Jahresbericht*, handed in to the Swedish Academy of Sciences, March 31, 1831 (*cf* Wohler's German translation, 1832, pp 44-8), in the following words—

'Da es nothwendig ist, für gegebene Ideen bestimmte, und so viel wie möglich consequent gewählte, Ausdrücke zu besitzen, so habe ich vorgeschlagen, Körper von gleicher Zusammensetzung und ungleichen Eigenschaften *isomerische* zu nennen, vom griechischen *isomeros* (aus gleichen Theilen zusammengesetzt).'

In the next volume of his *Jahresbericht* (Wöhler's translation, 1833, p 63) he gives a

further all important definition of the use he would make of the term, thus —

‘Um jedoch nicht Erscheinungen von nicht völlig gleicher Art mit einander zu verwechseln, ist es nothwendig den Begriff vom Worte *Isomerie* genau zu bestimmen. Ich erwähnte dass ich darunter Körper verstand, die aus einer gleichen absoluten und relativen Atomen-Anzahl derselben Elemente zusammengesetzt sind und gleiches Atomgewicht haben *womit nicht der Fall zu verwechseln ist, wo die relative Anzahl der Atome gleich ist, die absolute aber ungleich.* So ist z. B. die relative Anzahl von Kohlenstoff- und Wasserstoff-Atomen im bildlichen Gas und im Weindol absolut gleich, allein in einem Atom vom Gase sind bloss 1 Atom Kohlenstoff und 2 Atome Wasserstoff enthalten, CH_4 , während dagegen im Weindol 4 Atome Kohlenstoff und 8 Atome Wasserstoff enthalten sind, C_4H_8 . Um diese Art von Gleichheit in der Zusammensetzung bei Ungleichheiten in den Eigenschaften bezeichnen zu können, möchte ich für diese Körper die Benennung *polymerische* (von *πολυς*, mehrere) vorschlagen.’

It will be clear from this quotation, especially from the words which I have italicised, that Berzelius is never intended that polymerism should be regarded as a form of isomerism.

The following passage from the same source clearly exhibits Berzelius's intention as to the use which should be made of the term *metamerism* —

‘Allein es gibt noch andere Verhältnisse, wo Körper, im eigentlichen Sinne des Wortes, isomerisch scheinen, d. h. dieselbe relative und absolute Atomenanzahl derselben Elemente enthalten können, ohne *jedoch vollständig zu sein*. Ein solcher Fall ist wenn Körper aus zwei zusammengesetzten Atomen der ersten Ordnung bestehen, die sich auf verschiedene Weise gegen einander verhalten, und in Folge dessen ungleiche Körper bilden können, z. B. SnS (SnO_2S), schwefelsaures Zinnoxydul, und SnS (SnO_2S_2), basisches schwefelsaures Zinnoxyd, enthalten eine gleiche absolute und relative Anzahl derselben Elemente, und haben dasselbe Atomgewicht, können jedoch nicht als ein und derselbe Körper betrachtet werden. Bei solchen Körpern ist es der Fall, dass wenn sie eine gewisse Zeitlang bestanden haben, oder wenn die Temperatur geändert wird, eine Umlegung der Bestandtheile in ihnen vor sich geht, ohne dass etwas hinzukommt oder davon wegeht, und dass dadurch eine anders beschaffene Verbindung entsteht, welche Veränderung nicht selten von einer Temperatur-Erhöhung begleitet ist. Um solche Fälle bestimmt von Isomerie zu unterscheiden, können wir dafür die Bezeichnung *metamerische* Körper gebrauchen (von *μετα* in der selben Bedeutung wie in Metamorphose).’

Berzelius also cites cyanic and cyanuric acids, which were not then regarded as compounds of different molecular weight, as instances of metamerism, regarding the conversion of the latter into the former on heating as a case in which ‘die Cyanursäure von einem zusammengesetzten Atom der ersten Ordnung, oder einem Oxyd eines ternären Radicals, in ein zusammengesetztes Atom der zweiten Ordnung, nämlich in Cyansäure mit chemisch gebundenem Wasser übergeht’ Cyamelide, into which cyanic acid spontaneously changes, and cyanuric acid, in Berzelius's opinion, were (wenigstens vorläufig) isomeric oxides of the same radicle.

It is clear therefore that the conception involved in the definition of metamerism given in Watts' *Formes*, and in the first edition of this dictionary, is scarcely in conformity with the use of the term by Berzelius, indeed, in 1840 he speaks of ethylic formate and methylic acetate as isomeric. From the example afforded by the two tin compounds, as well as from the explanation given of the nature of the change from cyanuric to cyanic acid, it is to be supposed that the compounds which he intended should be included in the category of metamerism substances were such as we should now term *typically different*, and with this conclusion the modern practice is in

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distinct accord, yet the modern and the original application of the term are far from being equivalent, the *meta-* in metamorphosis denoting change the conversion of the one metameride into the other being expressly referred to, it would appear, in fact, that Berzelius distinctly intended to apply the term to those compounds which are capable, in modern phraseology, of undergoing ‘isomeric change,’ and perhaps to these alone.

It was not until 1840 that Berzelius proposed to substitute the term *allotropic* (‘von *ἀλλότροπος* welches bedeutet, von ungleicher Beschaffenheit,’ *B J* 1841 inorg. sec. p. 13) for isomeric, in the cases of the existence of modifications of elementary substances. He appears, however, to have contemplated its extension to compounds, judging from the following passage —

‘Es kann dann mehr als eine Ursache von dem was wir Isomerie nennen, geben, nämlich (1) Allotropie, wenn nämlich das vorhin angeführte Beispiel von den beiden Schwefelkiesen darauf beruht, dass der eine davon S₂ und der andere S₈ enthält, (2) die ungleiche Lage der Atome in der Verbindung, und (3) kann sowohl Allotropie als ungleiche Stellung der Atome in gewissen Fällen zugleich stattfinden.’

The existence of compounds containing allotropes of one of the constituent elements is distinctly suggested here, but there is nothing to indicate in what way allotropes may be regarded as related, that Berzelius had realised that the relation might be that of polymerides would appear to follow from his reference to Frankenheim's experiments on sulphur, in which it is suggested that the different modifications of sulphur form corresponding gases, and that the dark yellow gas of sulphur, weighing thrice as much as sulphur gas should according to calculation, is not that of the modification occurring in the ordinary sulphur compounds (*v. Allotropie*, vol. 1 p. 128) the view here taken is somewhat wider than that adopted by Prof. L. Meyer in that article).

As the term *allotropic* has a general significance, implying only another condition, and involves no assumption either regarding the molecular weights of, or as to the nature of the relationship which obtains between, the allotropes, it may with great advantage be employed in place of the term isomeric used in a wide sense, this latter term being preferably restricted to those cases in which there is the very closest similarity in structure. Polymerism, metamerism, and isomerism may in fact all be regarded as varieties of allotropy, there is certainly no reason why carbon compounds should be considered apart from those of other elements, or from elementary substances.

The rational formulae which are ordinarily made use of are condensed symbolic expressions affording more or less complete information as to the characteristic chemical properties of the compounds which they represent, especially with regard to the manner in which they are formed, and in which they undergo change when submitted to the action of various agents, and equimolecular allotropes which differ either in their mode of formation or in their behaviour under similar circumstances are, as a rule, necessarily represented by different rational formulae. The formulae devised for any class of compounds, however, will vary according to the views which

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are held as to the valencies of the constituent atoms. At present, formulae are almost invariably constructed on the hypothesis that the several units of affinity—the valencies—of a polyad atom, such as that of carbon, nitrogen, or sulphur, are of the same value and have identical functions, this conclusion being based on the fact that none of the simple derivatives of methane, ammonia, &c., exist in a greater number of modifications than the hypothesis requires. But it cannot be too positively stated that, notwithstanding the extent to which experimental investigation has been carried, we are yet but on the threshold of the temple in which the mysteries of valency are enshrined. The valency of the hydrogen atom is determined *ex hypothesi* to be unity, and when the facts generally are passed in mental review, it would seem that this conclusion is no mere hypothesis, it is not improbable also that the valencies of the atoms of at least the majority of metallic elements are invariable, but the valencies of the atoms of the non metals are apparently variable. By valency is here understood atom fixing power.

Most discussions on valency are dialectical rather than scientific, in consequence of our powerlessness at present to decide what constitutes 'a valency,' the deduction from Faraday's law of electrolysis, to which Helmholtz has directed the attention of chemists (Faraday lect., *C J Trans* 1881 p 277), that definite, as it were atomic, charges of electricity are associated with the atoms of matter—that a monad bears a single charge a dyad two, a triad three—is the only approach yet made to a theory of valency, but hitherto chemists have avoided the discussion of the subject from this point of view.

Oxygen and sulphur, nitrogen and phosphorus carbon and silicon, form gasifiable hydrides, from the composition of which we infer that the atoms of these elements are divalent, trivalent, and tetravalent, respectively, in the case of carbon and silicon there is no reason to suppose that either element ever manifests a higher valency. But both water and ammonia readily combine with other molecules the formation of such compounds from water is rarely interpreted as evidence of the possession by oxygen of the power of acting as a tetrad, but the water molecule is usually supposed to function in some occult manner as a whole, and to enter into a state of 'molecular combination' different from that of 'atomic combination' in which its constituent atoms exist. The formation of ammonium compounds, however, is more usually regarded as due to the manifestation of a higher degree of valency by the nitrogen. But there is no good reason for explaining the behaviour of oxygen in one way, and that of nitrogen in another.

The question to be decided is, whether a given element may possess two or more degrees of valency, and whether the so-called atomic and molecular forms of combination differ merely in degree and not in kind. Or, to put it in

¹ It appears to me that the term valency must at present be used in a perfectly general sense, and that we cannot restrict our attention to the consideration of gaseous compounds (*v. EQUIVALENCY*) in the case of gases, the problems are for the most part of a very simple kind, and rarely excite differences of opinion, in liquids and solids, however, present problems of great complexity

another way—is the number of atomic charges associated with a given atom invariable or variable, is it possible for an oxygen or sulphur atom, for example, to carry more than two, or for a nitrogen or phosphorus atom to carry more than three, charges? The writer has endeavoured to explain the exhibition of varying degrees of valency on the assumption that, while the number of charges which any given atom can carry is invariable, a single charge may operate in promoting the union of more than two atoms (*cf P M* January 1888), that in water, for example, the two charges of the oxygen atom are not fully engaged by those of the hydrogen atoms, and that consequently the oxygen atom is still possessed of a certain amount of *residual affinity*. It may be contended that, according to this hypothesis, a compound formed of say trimethylamine and ethyl iodide, $\text{Me}_3\text{N.IEt}$, would be an allotrope of a compound of ethyl dimethylamine and methyl iodide, $\text{EtMe}_2\text{N.MeI}$. The most careful experimental study of such compounds (*cf V Meyer a Lecco, Ber* 9, 809, Ladenburg, *Ber* 9, 561, 1634, Klinger a Claason, *A* 243, 193) has been made, however, with the result that, in the case of ammonium compounds, it is immaterial in what order or manner the radicals are introduced, and the same is true in the case of sulphine compounds—hence it is supposed that nitrogen has five, and sulphur four, affinities of equal value. But this by no means follows, as the occurrence of 'isomeric change' in such cases is in the highest degree probable—the compound abNId may alone be the stable form into which the allotropes abdNi , acdNi , bedNi , all spontaneously undergo conversion immediately on formation. There is little doubt that such 'isomeric changes' occur far more frequently than is commonly supposed, and it is most important that the possibility of 'isomeric change' should be very carefully kept in view in determining the constitution of compounds from the study of their behaviour in a limited number of interactions. As valency cannot be determined from any *a priori* considerations, and can only be deduced from the knowledge of the structure of the compounds of the elements whose valency is to be determined, it is obvious that the structure of a substance must be inferred from the widest and most careful study of all its properties. The study of the relationships of allotropic substances is in fact inseparable from that of valency, and the converse is equally true.

In the case of 'unsaturated' carbon compounds, it has been customary of late years to represent the affinities not engaged by other elements as saturating each other—thus, ethylene is formulated as $\text{H}_2\text{C.CH}_2$, acetylene as HC.CH . Thomsen's determinations of the heat of combustion of ethylene and acetylene in comparison with those of saturated hydrocarbons, as well as the general behaviour of such unsaturated compounds, may, however, be held to favour the view that the carbon atoms are possessed of free affinities, as expressed by the formulae

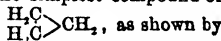
$\begin{array}{c} \text{H}_2\text{C} \\ \text{HC} \end{array}$ — $\begin{array}{c} \text{H}_2\text{C} \\ \text{HC} \end{array}$ —
Although the discussion of this question excited considerable attention a few years ago, it has latterly almost entirely fallen into

oblivion, but as very many of the cases of anomalous isomerism, of which an explanation is required, occur among compounds of the ethylenic type, it is one of considerable importance. The possibility of the two forms of com-

bination pictured by the expressions $\begin{array}{c} \text{H}_2\text{C} \\ \parallel \\ \text{H}_2\text{C} \end{array}$ and

$\begin{array}{c} \text{H}_2\text{C}- \\ | \\ \text{H}_2\text{C}- \end{array}$ should also be taken into account, especially in the case of ethylenic derivatives

Among the more recondite problems of valency requiring mention is that relating to the number of carbon atoms which form closed chains or rings. It is now regarded as well established that, in addition to the six atom ring of benzene, five atom rings also exist, indeed, their formation apparently takes place with peculiar readiness, the existence of both three- and four atom rings is now also generally held to be established, chiefly in consequence of the researches of W H Perkin, jun (cf *C J Trans* 1885 801, *et seq*). The hydrocarbon obtained by the action of sodium on trimethylene bromide, $\text{BrCH}_2\text{CH}_2\text{CH}_2\text{Br}$, is almost universally assumed to be the simplest compound of the kind, trimethylene

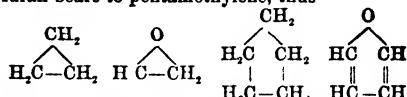


as shown by Freund, its discoverer, this hydrocarbon has the remarkable property of being readily absorbed by a solution of hydrogen iodide, forming normal propyl iodide, although it is acted on with extreme slowness by bromine. By the action of ethylenic bromide on the disodium derivative of ethylic malonate, Perkin has obtained an acid which he regards as a trimethylenedicarboxylic acid $\begin{array}{c} \text{H}_2\text{C} \\ | \\ \text{H}_2\text{C} > \text{C}(\text{COOH})_2, \end{array}$ and by employing trimethylenic bromide in place of ethylenic bromide, he has prepared what he regards as tetramethyl

enedicarboxylic acid $\begin{array}{c} \text{CH}_2-\text{CH}_2 \\ | \\ \text{CH}_2-\text{C}(\text{COOH})_2, \end{array}$ both acids

closely resemble the hydrocarbon in their behaviour with bromine and halohydrates. The easy resolution of closed carbon chains by halohydrates in this manner, however, is altogether without precedent in the case of five and six atom carbon rings, at all events, which, as a rule, cannot be split by means of halohydrates, but are frequently resolvable by bromine, their behaviour is more nearly akin to that of five atom rings, such as those of furfuran and indole, which also resist the action of bromine, but are resolvable by treatment with agents which are commonly regarded as weaker than bromine (the formation of pyridine derivatives from pyrrole and of quinoline derivatives from indole, which appears to involve the resolution at some stage of the operation of the five atom rings, is contemplated in this statement). The alternative formula for trimethylene, if it be not a closed chain hydrocarbon, is $\text{CH}_2\text{CH}_2\text{CH}_2$, which represents it as a compound in which two terminal carbon atoms are each possessed of a *single* free affinity, the possibility of the existence of such compounds has not yet been contemplated by chemists, excepting myself. It is conceivable, I think, that such a compound would be attacked by bromine with difficulty, in consequence of the two un-

satisfied carbon atoms failing to assist each other in separating the constituent atoms of the bromine molecule and the impossibility of a 'conducting chain of molecules' being formed between the carbon atoms, as bromine is a dielectric, as solutions of the halohydrates are electrolytes of low resistance, such a conducting chain might, however, be formed in their case, and the requisite electrolysis of the halohydrate molecule could thus occur. According to Thomsen, trimethylene has a *higher* heat of combustion than propylene, CH_2CHCH_2 , judging from the analogy afforded by benzenoid compounds, it is to be expected, however, that the closure of the chain involved in the formation of the three atom ring would be attended with a considerable loss of energy, and that propylene would, therefore, have the higher heat of combustion. Its behaviour with bromine certainly justifies this view. The confirmation of Thomsen's statement thus becomes of extreme importance. It is a noteworthy fact that ethylenic oxide, according to Thomsen, also has an exceptionally low heat of formation, and that this observer has proposed to represent it as a dimethylenic oxide of the formula CH_2OCH_2 . Ethylenic oxide as represented by the conventional formula bears a similar relation to trimethylene that furfuran bears to pentamethylene, thus




Thomsen's formula for ethylenic oxide is inadmissible, as it indicates a severance of the carbon atoms, if, however, the formula were written $\text{CH}_2\text{CH}_2\text{O}$, it would correspond to that given above to trimethylene, Thomsen's observations that both ethylenic oxide and trimethylene have an exceptional heat of combustion may therefore be regarded as mutually confirmatory. It is also to be noticed that the compound formed from ethylenic bromide and a sulphide is not the corresponding sulphide, but the polymeride $\text{CH}_2-\text{S}-\text{CH}_2$.

thereof $\begin{array}{c} | \\ \text{CH}_2-\text{S}-\text{CH}_2. \end{array}$ If an open chain formula

be assigned to trimethylene, Perkin's tri- and tetramethylene derivatives must also be represented by open chain formulae. Perkin has fully discussed this question, and has pointed out the improbability attaching to such formulae. The evidence does not appear to be sufficient, however, to permit of a final decision being arrived at with regard to so difficult a question.


Finally, it is necessary to refer to a problem closely akin to the two previously considered, viz. that relating to the distribution of the *spare* affinities of the carbon atoms in closed chains, of those affinities, that is to say, which are not engaged in the formation of the ring, or in retaining the hydrogen atoms. This problem is chiefly of importance in discussing the structure of benzenoid hydrocarbons and their derivatives. Of the various formulae proposed for benzene, that of Kekulé always has been, and still remains, the most popular, but it is open to the serious objection that it represents benzene as a compound containing three pairs of carbon atoms in the same condition as the pair in ethylene. Dewar's for-

mula is open to a similar objection. The prism formula of Ladenburg and the diagonal formula of Claus cannot be objected to on this ground, but are open to criticism in many other respects, and in the light of Von Baeyer's recent researches on the reduction products of terephthalic acid (cf. A 245, 103, 251, 257) these formulae are generally regarded as finally disposed of (cf. Miller, *C. S. Trans* 1887 208). A symbol proposed by the writer in February 1887 (cf. *P. M.*), and a year later also by Von Baeyer (A 245, 122), appears to be exempt from the deficiencies which characterise previous formulae, but it embodies somewhat unconventional conceptions, and therefore has not yet attracted attention. The

symbol in question  has been very happily

termed the *centric* formula by Von Baeyer, he expressly states that this formula is to be understood to indicate that 'die 6 Kohlenstoffvalenzen des Benzols sich sättigen, ohne dadurch drei Verkettungen der Kohlenstoffatome zu bewirken' (A 251, 285)—one valency of each atom is directed towards the centre of the ring, and these valencies mutually paralyse each other (A 245, 122). My own words were 'Of the twenty four affinities of the six carbon atoms twelve are engaged in the formation of the six carbon ring, while the remaining six react upon each other, acting towards a centre as it were, so that the affinity may be said to be uniformly and symmetrically distributed. I do not consider that, apart from its connexion with the other carbon atoms owing to their association in the ring, any one carbon atom is directly connected with any other atom not contiguous to it in the ring—each individual carbon exercises an influence upon each and every other carbon atom—there is an excess of affinity beyond what is required to maintain the C_6H_6 ring, but I do not consider that each carbon atom can be considered to have an affinity free'.

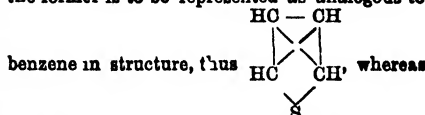
The conclusion here arrived at with regard to benzene, that no direct connexion exists between any but the contiguous carbon atoms in the ring—that para carbon atoms are not and cannot become united—may be regarded as of universal application. Von Baeyer's experiments prove, moreover, that the dihydro terephthalic acid in which an atom of hydrogen is associated with each of the para-carbon atoms to which the carboxyls are attached, unlike terephthalic acid, behaves as an unsaturated compound, forming a tetrabromide, and that it is to

be represented by the formula . From

this it follows that the type changes on conversion of the 'centric' compound into the di addition compound, and probably this is generally the case, for example, when quinol is converted into quinone



It would also follow that in the case of benzenoid compounds four of the six 'spare' affinities can not act 'centrically'. To what extent this is true in the case of other rings remains to be ascertained, from the remarkable similarity of thiophene and benzene, it would appear probable that the former is to be represented as analogous to



the analogous compounds furfuran and pyrrole more nearly resemble the unsaturated compounds in their behaviour, but this is perhaps ascribable to the influence exercised by the oxygen or imidogen.

The foregoing brief discussion will suffice to direct attention to the numerous problem comprised in the philosophy of isomerism, and to show how far from final are the conclusions as to structure which we are at present able to arrive at.

In a large and rapidly increasing number of instances, it is impossible to assign different rational formulae to compounds undoubtedly different so long as the system employed takes into account merely the nature of the radicles and the manner in which they are associated as pictured by disposing them in a single plane. If, while assuming the hydrogen atoms in a compound, such as methane—in other words the four valencies of the carbon atom—to be equal, it be supposed that the radicles introduced in place of the hydrogen atoms occupy relatively different positions in the plane, a variety of isomeric derivatives would appear to be possible, e.g. two of the form $CRRR_1R_1$, two of the form $CRRR_1R_2$, and three of the form $CRR_1R_2R_3$, $R_1R_1R_2R_3$, being different radicles. A special study of methane derivatives from this point of view has been made by Henry, but both his results and our general experience show that isomerides such as are here contemplated do not occur.

By considering the arrangement of the atoms in space, conclusions have, however, been arrived at which are far more in harmony with experience. Such a step was first taken in 1874 by Van't Hoff (*La Chimie dans l'Espace*, Rotterdam, 1875), and independently and almost simultaneously by LeBel (*Bl* [2] 22, 337, cf. *ibid* 23, 295). A German adaptation of Van't Hoff's pamphlet entitled, *Die Lagerung der Atome im Raume*, was published in 1877 by Hermann. A full account of the subject is to be found in the previous edition of this dictionary and in Miller's 'Chemistry,' vol. III.

The fundamental hypothesis of the Van't Hoff system consists, as is well known, in supposing that the carbon atom occupies the centre of a tetrahedron and that its four affinities are directed towards the four solid angles. When four different radicles are associated with the carbon atom, but only in such a case, two isomerides are possible, represented by two irregular non-superposable tetrahedra bearing to each other the relation of an object to its reflected image, and, moreover, these isomerides should be of enantiomorphous crystalline form, as

well as optically active and possessed of equal and opposite rotatory powers, as the molecules are unsymmetrical, such tetrahedra exhibiting, in relation to an axis drawn parallel to the corresponding edges, a screw shaped grouping of the four summits, turning to the right in the one form and to the left in the other. A carbon atom thus situated is termed *asymmetric*, and is represented in a formula by an italicised *C*. The hypothesis serves therefore at once to explain both the existence of isomerides which cannot be represented by formulae written in a single plane, and to account for the optical activity of certain substances. Thus in the case of tartaric acid³, which contains two asymmetric carbon atoms, but is composed of two equal groups, $(\text{COOH})(\text{HO})\text{HC}(\text{CH}(\text{OH})(\text{COOH}))$, the hypothesis indicates the existence of two optically active isomerides of equal but opposite rotatory powers, and a third inactive isomeride in which the optical effect of the one asymmetric carbon atom is balanced and neutralised by the equal opposite effect of the other, it thus accounts for the existence of dextro-, laevo-, and meso tartaric acids, racemic acid, the fourth modification, apparently, is to be regarded as a 'physical' allotrope formed by the conjunction of the two active isomerides. It would seem that it does not exist in solution. Van't Hoff has shown, in a recent much extended new edition of his pamphlet, that every prevision of the hypothesis with reference to the optical characters of isomerides has been fulfilled in the most complete manner possible by the investigations carried out in the interval since its enunciation by Le Bel and himself (cf. *Dix Années dans l'Histoire d'une Théorie*, Rotterdam, P. M. Bazendijk, 1887), this remarkable agreement of practice with theory has naturally led to the almost universal adoption of the hypothesis.

The hypothesis also provides for a greater number of isomerides in the case of compounds of the ethylenic type than is indicated if the space relationship of the radicles be omitted from consideration, if a compound of the form $\text{R}_1\text{R}_2\text{C}=\text{CR}_3\text{R}_4$ be represented by two tetrahedra joined so as to have one edge in common, it will be found that only one such figure can be constructed, if either the four radicles are identical, or if only R_1 differs from R_2 , or R_3 from R_4 , but if R_1 is different from R_2 , and R_3 is also different from R_4 , although R_1 and R_3 , R_2 and R_4 are identical, two such figures may be constructed—this is more readily rendered obvious by the adoption of the simpler plan of writing the symbols of the radicles attached to the two doubly linked carbon atoms on either side of a line representing the plane of their conjunction.

Thus the symbols $\begin{smallmatrix} a & b \\ a & b \end{smallmatrix}$ and $\begin{smallmatrix} a & b \\ b & a \end{smallmatrix}$ represent modifi-

cations in which in the one case the two similar radicles are situated symmetrically with reference to a plane at right angles to the axis of the system, and in the other are symmetrically situated with reference to the axis of the system, such modifications may therefore be termed, as Wislizenus has suggested, the *plane* and *axial* symmetric modifications. The isomerism of maleic and fumaric acids is regarded by Van't

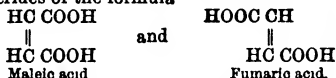
Hoff as dependent on such a difference in structure.

Such a *stereometric* mode of formulation does not afford an increased number of expressions for acetylenic derivatives, the same is true of benzenoid compounds if Kekulé's symbol be adopted (cf. Marsh, *P. M.* Nov 1888, p. 426). It may be noted, however, that the asymmetric carbon atom hypothesis is applicable to the explanation of the optical activity manifested by a variety of closed chain compounds, such as quercitol and quinic acid, which are derivatives of hexamethylene, and conine and its homologues, which are derivatives of piperidine.

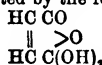
Although the Van't Hoff Le Bel hypothesis has been very generally accepted as affording a sufficient explanation of a very large number of cases of isomerism difficult to account for in accordance with the existing canons of belief, it is as yet by no means certain that it can be always so regarded, in many cases the difference between isomerides is so great that it is somewhat difficult to believe that it depends on so comparatively simple a difference in structure as the presence of one or more asymmetric carbon atoms would involve. The two isomeric tartaric acids are the veritable image the one of the other, but this is rarely the case thus hydrobenzoin differs very considerably from isohydrobenzoin, as do also mannitol and dulcitol, and to a still greater extent the acids, saccharic and mucic, formed from these latter, in all these cases the isomerism is assumed to depend on mere geometrical differences. In the case of mesotartaric acid, the intra-molecular neutralisation of one asymmetric carbon atom by the other already produces a marked effect, as evidenced by the difference in properties of this acid in comparison with either of the active modifications. It may be, therefore, that in the case of mannitol and dulcitol, and the acids formed by their oxidation, the accumulated effect of the several asymmetric carbon atoms is much greater than in the case of tartaric acid, and hence the greater difference in properties between the isomerides. Very little alteration is involved in the formation of racemic acid, and, as above pointed out, this substance appears to be a mere 'physical' allotrope, hardly a polymerde, of its active constituents, in certain cases the formation of the 'racemic form' is attended with a considerable alteration in properties—in the case of leucine and of camphoric acid, for example, their racemic forms being much less soluble and of higher melting points than their optically active constituents. In these cases there would seem to be a more intimate union than in the case of racemic acid, the marked difference between saccharic and mucic acid may be due to some such cause as this. But the isomerism of the hydrobenzoin is more difficult to explain, assuming that they are both compounds of the formula $(\text{C}_6\text{H}_5)(\text{HO})\text{HC}=\text{CCH}(\text{OH})(\text{C}_6\text{H}_5)$. Including a racemic form, four modifications of such a compound appear possible, two of which should be optically active. Neither compound is optically active, and, judging from Zincke's observations, it does not appear probable that, if the one be the meso or inactive modification corresponding to mesotartaric acid, the other is the racemic form. The assumption that the one, perhaps

isohydrobenzoin, contains the two hydroxyl-groups attached to the one carbon-atom, thus $C_6H_5 \cdot H_2C(OH)_2$, would satisfactorily account for its behaviour, but has hitherto been rejected as improbable owing to the general belief that compounds of such a character are excessively unstable. But it may be that the presence of the negative phenyl groups confers stability in such a case, much as in the case of chloral hydrate, and it is well to remember that in not a few instances of late years proof has been given of the incorrectness of views based on general considerations, as in the case of phthalic chloride, for example, and the hydroxamic acids.

Still greater difficulties occur in accepting the conclusion that the Van't Hoff hypothesis affords an explanation of the isomerism of unsaturated compounds such as maleic and fumaric acids, which, according to this view, are respectively the plane symmetric and axially symmetric isomerides of the formula



It is well known how great are the differences between these two acids, both in physical properties and in general chemical behaviour, and that maleic acid alone yields a corresponding anhydride, fumaric acid being converted into the same anhydride when dehydrated. Ostwald also has shown that the electrical conductivity of their solutions is such as to indicate that maleic acid is a weak acid akin to selenious and phosphorous acids, whereas fumaric acid is a well-marked dibasic acid (*J pr* 32, 362). Maleic acid has also a considerably greater heat of combustion than fumaric acid (Thomsen, *J pr* 40, 202). Roser has suggested that, whereas fumaric acid is a normal dicarboxylic acid, maleic acid is to be represented by the formula



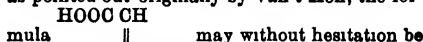
This formula would afford a satisfactory explanation of the great differences observed between the two acids, and it has been strongly advocated by Anschütz (*A* 254, 168). No valid argument has yet been advanced which would prevent its adoption.

It may here be pointed out that no attention is paid in applying the Van't Hoff hypothesis to unsaturated compounds to the peculiarities which are manifest in such compounds, and which apparently must be attributed to the presence of unsaturated carbon atoms, a 'double or ethylenic bond' is represented as the precise equivalent of two single bonds, and a 'treble or acetylenic bond' as the equivalent of three single bonds, which is certainly not in accordance with facts, and especially with Thomsen's observations on the heats of combustion of unsaturated compounds.

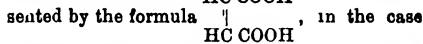
Wislicenus has not only accepted the Van't Hoff Le Bel hypothesis in its entirety, but has in the most ingenious manner possible extended its application, and has endeavoured both to elucidate the structure of geometrically isomeric unsaturated compounds, and to explain the 'isomeric changes' which such compounds frequently exhibit (*Abhandlungen der math. phys. Klasse*

der könig Sächsischen Ges der Wissenschaften, Band xiv Leipzig, 1887).

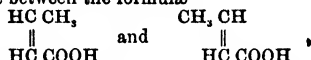
In the case of maleic and fumaric acid—assuming that these are stereometric isomerides—as pointed out originally by Van't Hoff, the formula



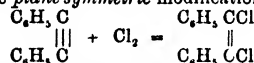
assigned to the latter, as it is incapable of forming an anhydride, whereas maleic acid, being easily convertible into the anhydride, is represented by the formula



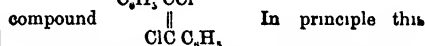
of crotonic and isocrotonic acids however, it is more difficult to find criteria on which to base a choice between the formulae



and similarly in other cases. Wislicenus considers that if a compound of the form $C_2R_1R_2R_3$, consisting of the two systems $R_1R_2C=$, and $=CR_3R_4$, be derivable from a corresponding acetylenic compound, its constitution can at once be inferred from the fact that when one pair of bonds between the carbon atoms in the acetylenic compound is severed, as the two atoms are still united by double bonds, no rotation of the systems can take place, consequently the added radicles both occupy positions on one side of the common axis of the systems. Thus the tolane dichloride melting at 143° , obtained by chlorinating tolane, is necessarily the plane symmetric modification

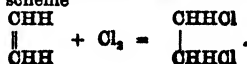


The isomeride of lower melting point (63°) must therefore be regarded as the axially symmetric compound



method appears perfectly sound, but it is based on an assumption with reference to the manner in which the carbon atoms in ethylenic and acetylenic compounds are united, which, as previously pointed out, is perhaps open to question, it also involves the conclusion that the radicles attached to the carbon atoms are incapable of changing their positions, which is also by no means a safe assumption, bearing in mind the extreme readiness with which 'isomeric changes' occur.

A method of more universal application, but involving much more complex considerations, is the following. In saturated compounds in which the carbon atoms are united by single affinities, one carbon system must be capable of rotating about another, moreover, it is to be assumed that the atoms in a molecule—even those which are not directly connected—exercise an influence on each other, and will therefore tend to condition such rotation so that radicles which have the greatest affinity are brought into the closest proximity possible. Thus, on converting ethylene into its chloride, in the first instance the change would take place in accordance with the following scheme



But, in consequence of the superior affinity of hydrogen for chlorine, such a system would be unstable, rotation would set in, and the more

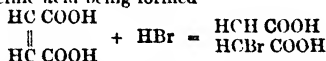
stable system $\begin{array}{c} \text{CHHCl} \\ \text{CClHH} \end{array}$ would result.

To take another case, that of the conversion of tolane tetrachloride into dichloride by reduction, this compound may present three configurations, viz

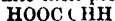


It would probably chiefly consist, especially at low temperatures, of the second and third modifications, as the dissimilar radicles are most proximately situated in these, and as these two modifications would both furnish the axially symmetric dichloride on reduction, it is to be supposed that the chief product of reduction will be this dichloride, actually that melting at 63° is chiefly obtained (Blank, *A* 248, 1), and it is, therefore, to be supposed that this modification is axially symmetric—a conclusion which harmonises with that previously arrived at. To this it may be objected that our knowledge of the relative affinities of the radicles within a compound is purely hypothetical, and that it is by no means certain that dissimilar radicles would in all cases tend to influence and attract each other more than would similar radicles. Thomsen's observations on the heats of combustion of chlorine compounds, in fact, would appear to favour an opposite conclusion. But a more important argument is to be found in the fact that in cases in which the constitution may fairly be regarded as established, the relation is of the obverse order to that required if the contention of Wislicenus be correct: thus, the symmetric or para derivative of benzene always has the highest melting point, the same appears to be true of the symmetric tri- and tetra-derivatives, and, in the case of naphthalene, the axially symmetric isomeride is always that of highest melting point (*Cf C S Proc* 1888 93).

Another use which Wislicenus has made of the argument here criticised may now be mentioned. It is a well known fact that maleic acid is very readily converted into fumaric by the action of acids. Wislicenus supposes that when this conversion is effected, for example, by bromhydric acid, the double bond becomes severed, bromosuccinic acid being formed



but rotation setting in, the radicles are brought into their preferential positions, viz



HC BrCOOH , and when, by the action of the water present, this modification becomes deprived of hydrogen bromide, fumaric acid naturally results. Although in the highest degree ingenious, this conception unfortunately does not appear to be in accordance with the facts, for, as Anschütz points out (*L.* 254, 168), the conversion takes place under conditions under which the succinic derivative is stable, and there is no reason therefore to suppose that such a compound is formed at any stage of the conversion of maleic into fumaric acid, if maleic acid be formulated in the manner advocated by An-

schutz, its conversion into fumaric acid by acids is easily understood.

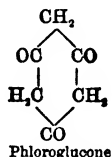
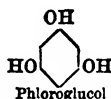
Victor Meyer's researches have led him to carry speculation even further. On submitting benzil to the action of hydroxylamine, Meyer and Goldschmidt obtained an a dihydroxime which they found was converted into a more stable β -isomeride by heating with alcohol to about 180° , a careful investigation of these compounds was subsequently made by Meyer and Auwers, but they were unable to discover any substantial difference in their chemical behaviour, and they came to the conclusion that both must be regarded as compounds of one and the same formula $\text{C}_6\text{H}_5 \cdot \text{C}(\text{N OH}) \cdot \text{C}(\text{N OH}) \cdot \text{C}_6\text{H}_5$. As the existence of two such compounds was incompatible with the assumption that carbon atoms united by single affinities are free to rotate, Meyer and Riecke (*B.* 21, 946) have put forward an hypothesis as to the nature of the carbon atom itself which serves to account for two kinds of union by single affinities, one in which rotation is impossible, the other in which it can freely take place. Having regard, however, to the readiness with which 'isomeric change' takes place, and to the extremely imperfect state of our knowledge of the exact manner in which polyad elements are held in association, there cannot be any doubt that it is premature to conclude that the benzil dihydroximes are necessarily structurally identical, the study of chemical interchanges is after all but an approximate and fallible mode of determining structure.

Hantzsch and Werner (*Ber* 23, 11) have quite recently proposed to extend the Van't Hoff conception to the nitrogen atom itself. They suppose that in some nitrogen compounds the three affinities of the nitrogen atom are directed towards three of the solid angles of an irregular tetrahedron, and that the nitrogen atom itself is located at the fourth. In the case of compounds of the form $\text{CXY} \cdot \text{NZ}$, which are comparable with carbon compounds of the form $\text{CXY} \cdot \text{CH}_3$, it is conceivable that the Z radicle may alter its position in space relatively to the radicles X and Y and thus give rise to isomerides.

Victor Meyer (*B* 23, 567) has, however, pointed out that the views of Hantzsch and Werner are in many respects in conflict with the experimental evidence. This memoir is a noteworthy contribution to the discussion of a number of the more obscure cases of isomerism. (The two papers here referred to have been published since this article was in type.)

It is here the place to refer to the numerous discussions which have taken place during late years in cases in which one set of interactions of a compound appear to be in accordance with one formula, while another set favour a different but closely related formula, which have led, in fact, to the recognition of labile or pseudo-forms capable of passing spontaneously into stable forms. For example, it has long been a question whether ethylic acetoacetate is to be represented as $\text{CH}_3 \cdot \text{CO} \cdot \text{CH}_2 \cdot \text{CO}_2\text{Et}$ or by the formula $\text{CH}_3 \cdot \text{C}(\text{OH}) \cdot \text{CH} \cdot \text{CO}_2\text{Et}$. A discussion of such cases has been given by Laar (*B* 18, 648, 19, 730), who proposes to term such isomerides *tautomeric*. But, as he practically himself admits, the use of such a term savours of tauto-

logy. V. Meyer subsequently brought forward a suggestion of Jacobson's to substitute *desmotropic* for tautomeric. Hantzsch and Herrmann (*B* 20, 2801), while using the term tautomerism when speaking of compounds capable of passing from the one type into the other, proposed to limit the term *desmotropic* to each of the states. As the phenomena in question are the outcome of mobility and not of fixity, the new term also appears to be particularly ill chosen, and bearing in mind the intention of Berzelius in introducing the term *metamerie* (see p 81), it would appear that this old term is a peculiarly appropriate one to use in such cases. Laar suggests that a compound which is capable of an 'alternative' behaviour actually has an alternating structure, the intra-molecular condition being such that the structure is of one kind at one moment and different at another. But, as Hantzsch and Herrmann remark, it is scarcely necessary to make such an assumption. Ethylic succinylsuccinate and its derivatives manifest the peculiarity in question in a high degree, acting sometimes as quinonic compounds and sometimes as quinols, i.e. in modes such as correspond to the presence of one or other of the two forms $\text{CO CH}(\text{CO}_2\text{Et})$ and $\text{C}(\text{OH})\text{C}(\text{CO}_2\text{Et})$. As a rule only one of the forms is stable, the other being developed in the course of the change, thus phloroglucol appears to be a true trihydroxybenzene, i.e. a phenol, but frequently it affords derivatives of 'triketohexamethylene,' which may with propriety be termed phloroglucone. Thus



It is not improbable that in those cases in which the several forms can be obtained in a definite crystalline form, the necessary stability is conferred by union of the fundamental molecules amongst themselves. The formation of such molecular complexes is rendered probable by a large number of observations, one of the most striking is that recently brought forward by Perkin in the case of orthomethoxybenzaldehyde $\text{CH}_3\text{O C}_6\text{H}_4\text{COH}$ (*C J Trans* 1889 549), which is capable of existing in two solid modifications, one unstable melting at 3° , the other stable and melting at 35.5° . A list of similar cases of what is sometimes termed *physical isomerism* is quoted by Perkin.

Having thus briefly touched on the numerous problems which the study of the different kinds of *allotropism* presents, it appears desirable finally to re-direct attention to the terminology of the subject.

The term *allotropic*, as already pointed out, has a perfectly general meaning and is therefore applicable to the phenomena generally, and may be used in all cases in which the nature of the relationship is obscure.

According to our modern conceptions, truly isomeric substances—substances composed of equal parts—are equi-molecular compounds containing identical radicals arranged in relatively

different modes, and on the principle of calling a spade a spade, bearing in mind that it was obviously the intention of Berzelius to limit the scope of the expression, the term *isomeric* should be used only with reference to such compounds. The space relationship of the radicals being the determining cause of isomerism, although it is not always requisite in order that it may become apparent to express their relationship according to stereometric canon, it is scarcely necessary to make any principal distinction between cases, such, for example, as occur among benzene derivatives and between the tartaric acids, but if it be thought desirable to call attention in some way to the finer isomerism which obtains in cases such as the latter, the term *ekisomerism*, from *εἰς*, a likeness or image, may be suggested as not inappropriate.

Typically different allotropes belonging to different classes of compounds might well be termed *heteromeric*, metameric being reserved for those heteromeric allotropes which change their type with exceptional facility in the course of chemical interchanges, but if the use of the term in this restricted sense be objected to, such allotropes might advantageously be spoken of as *isodynamic*. Allotropes belonging to the same class but consisting of different radicals—the butylic alcohols, for example—might be termed *isonomic*. It appears unnecessary to specially distinguish the physical isomerism manifest in the occurrence of several crystalline forms of different melting point. The phenomena of *pleomorphism* generally, as well as those involved in change of state from solid to liquid and gas, and the existence of allotropic forms both of the metallic and non-metallic elements, are now being more and more generally attributed to changes in molecular complexity, and if this be the case, such allotropes mostly partake of the nature of polymeric allotropes. H E A.

ISOMORPHISM (*Isos*, equal to, *μορφή*, form). In the year 1819 Mitscherlich (*A Ch* [2] 14 172) discovered that certain arsenates and phosphates of analogous constitution crystallised in the same form, subsequent investigation (*A Ch* [2] 19, 350, 24, 264, 355) led to the general conclusion that substances of analogous chemical constitution possess the same or nearly the same crystalline forms, and will under suitable conditions crystallise together in all proportions to form homogeneous mixed crystals, such substances are termed *isomorphous*. In spite of numerous attempts to widen or otherwise modify it, this definition has lasted to the present time. Owing, however, to recent researches, especially those of Groth, it will be best not to treat isomorphism as an isolated phenomenon, but as a part of that branch of physical chemistry which studies the relations between the chemical composition and crystalline form of bodies, and which from a knowledge of the constitution and chemical properties of a substance seeks to predict its system, form, and crystallographic constants. We are still very far indeed from doing anything approaching to this, for, although attempts have not been wanting—v. Schrauf (*Physikalische Mineralogie*, 1888 bd 2, 166, *Z K* 9, 265) and Barlow (*O N* 58, 8, 16)—small success has so far attended them,

and at present crystal form cannot be deduced from a knowledge of chemical constitution and properties alone, if, however, we find that in a given case certain atoms arranged in a certain definite way are accompanied by a certain definite form, we may argue that similar atoms similarly arranged will be accompanied by a similar form. This hypothesis is found to be true, and its verification has resulted in the discovery of relations between the forms of substances more or less chemically allied. These relations may be conveniently discussed under the three following heads —

I The same chemical substance possesses two or more forms—*Polymorphism*

II Bodies more or less closely related chemically have more or less analogous forms—*Morphotropy*, including *Isomorphism*

III Bodies not chemically related possess the same form—*Isogonism*

Full references, especially for the early history of the subject, will be found in the article *Isomorphism* by Arrhenius, in Fehling's *Handwörterbuch der Chemie*, 1878 vol. 1, for more recent work Fock's *Einführung in die Chemische Crystallographie*, Leipzig 1888, may be consulted, while full lists of substances considered isomorphous have been given by Topsoe, *Vidtskrift for Physik og Chemi*, 8, 5 193, 321, 9, 225 summarised by Arrhenius above. For the use of isomorphism in determining atomic weights c. ATOMIC AND MOLECULAR WEIGHTS, vol. I, p. 336

POLYMORPHISM Häuy believed that every substance possessed its own characteristic form, and, in spite of numerous observations tending to establish the chemical identity of calcite and aragonite (v. De Lisle, Klapproth, Thenard, and Stromeyer), refused to admit that the same substance could yield two distinct kinds of crystals. In the course of the work on the arsenates and phosphates which led him to announce that two different chemical compounds could possess the same or nearly the same crystalline form, Mitscherlich found that $\text{NaH}_2\text{PO}_4 \cdot \text{H}_2\text{O}$ gave, according to the conditions, two kinds of crystals, while his discovery in 1823 (*A. Ch.* [2] 24 264) that sulphur crystallised both in rhombic and oblique forms readily convertible the one into the other by change of temperature alone, established conclusively that the same substance could exist in two different forms, to express these phenomena Mitscherlich proposed the term *dimorphism*, and in 1828 recognised the possibility of *trimorphism*. The existence of several modifications of the same substance may be generally expressed by the terms *polymorphism* or *heteromorphism*. At the present time these terms are usually applied to compounds only, polymorphism, when exhibited by elements, being termed *allotropy* (q. v., vol. I, p. 128). The change of one dimorphic modification into another has been especially studied by Lehmann, and all known cases have been collected and discussed by him in his *Molekularphysik*, 2 vols. Leipzig, 1888 (vol. I pp. 119–219, vol. II pp. 898–415).

MORPHOTROPY From the time of Mitscherlich onwards the rapid growth of organic chemistry provided a large number of new substances for crystallographic examination, and as the existence of many cases of isomorphism among inorganic substances had been established, numerous attempts were made to introduce the same conception into organic chemistry

In general, however, these efforts were fruitless, and the researches merely resulted in the introduction of new terms and in several attempts at widening Mitscherlich's definition.

Thus Laurent, who investigated the halogen substitution and addition products of naphthalene and salts of the fatty acids, concluded that identity of system is not necessary for isomorphism. Nickles and De la Provostaye collected cases of analogous angles and introduced new terms, such as hemi isomorphism, isomorphism, and hemimorphism, to express those cases in which crystals of two substances had each a zone with nearly similar angles (v. papers by these authors, *C. R.* 11, 15, 20, 23, 26, 27, 29). Pasteur again (*C. R.* 26, 535) found analogies between the tartrates, while other workers in the same direction were Rammelsberg, Marignac, Delafosse, Sella, and Hjortdahl (*J. pr.* 94, 286). Owing to the paucity of results, such researches lost favour till Groth, in 1871 (*P.* 141, 131, *B.* 3, 449), attacked the subject from a new point of view, and set himself to investigate the changes in crystal form which take place when one or more of the hydrogen atoms of an organic molecule, such as benzene, are substituted by other atoms or groups. The relations observed between the form of the parent substance and the form of the substituted body Groth terms morphotropic relations, while he attributes the actual change of form produced by substitution to the morphotropic force of the elementary atom or group. Benzene, which crystallises in rhombic pyramids, was the first substance examined by Groth, he compared its form with the forms of as many as possible of its hydroxy-, nitro-, amido-, and haloid, substitution products, and his researches have led to the following general statement. The less the chemical character of the combination is changed by substitution the less is also the change of crystal form, which depends

1 On the specific morphotropic properties of the substituting atom or group,

2 On the chemical nature and crystalline system of the body in which substitution occurs,

3 On the relative positions of the substituting atoms or groups in the molecule.

1 Morphotropic properties of atoms and groups

(a) The metals Potassium and ammonium, when substituted for H, only change one axis, thus —

		a	b	c
Picric acid	rhombic	0.937	1	0.974
Potassium picrate	"	0.942	1	1.352

As a rule, all the alkali metals produce the same effect

(b) Substitution of OH for H in benzene derivatives only alters the crystalline form slightly, and in rhombic substances the ratios of two of the axes remain unaltered, and only the third axis is materially affected

NO_2 and NH_2 behave much like OH

		a	b	c
Benzene	rhombic	0.991	1	0.799
Resorcin	"	0.910	1	0.540
Nitrophenol 1 2	"	0.878	1	0.6007
Dinitrophenol 1 2 4	"	0.933	1	0.753
Trinitrophenol 1 2 4 6	"	0.937	1	0.974

(c) Cl, Br, and I act more energetically, but less regularly, than NO_2 . The angles of one zone, however, remain like those of the parent substance, while the system is generally altered to one of lower symmetry

Benzene	rhombic	110	$1\bar{1}0 = 96^\circ 30'$
Dichlorobenzene	oblique	"	" $98^\circ 40'$
Tetrachlorobenzene	"	"	" $96^\circ 17'$

(d) The influence of the CH_3 group is extremely variable, depending very much on the nature of the substance into which it enters. Among the substituted ammonias it is often without influence, thus, $2\text{NH}_4\text{Cl PtCl}_6$, $2\text{NH}(\text{CH}_3)_2\text{Cl PtCl}_6$, $2\text{N}(\text{CH}_3)_3\text{Cl PtCl}_6$, all crvs tallise in the cubic system

2 Influence of the nature of the parent substance. When the parent substance belongs to the cubic system, substitution has either no influence on the form, or else causes change to a system of lower symmetry. In other systems the axial ratios may alone alter, or the system itself may be changed, according to the morphotropic force of the substituting element or group. If the hydrogen atoms of like functions are replaced singly the system frequently at first changes to one of lower symmetry, but when the substitution is complete returns to the symmetry of the parent substance. The methyl substitution compounds of $2\text{NH}_4\text{Cl PtCl}_6$ illustrate this well —

$2\text{NH}_4\text{Cl PtCl}_6$	Cubic
$2\text{NH}_3(\text{CH}_3)\text{Cl PtCl}_6$	Rhombohedral
$2\text{NH}_2(\text{CH}_3)_2\text{Cl PtCl}_6$	Rhombic
$2\text{NH}(\text{CH}_3)_3\text{Cl PtCl}_6$	Cubic
$2\text{N}(\text{CH}_3)_3\text{Cl PtCl}_6$	Cubic

As an illustration of the influence of the chemical nature of the parent substance, we may quote the observation of Hintze that, contrary to the general rule, the substitution of the paraffinoid H in $(\text{C}_6\text{H}_5)_3\text{CH}$ by (OH) or Br raises the symmetry, instead of lowering it

3 Influence of position.—The relative position of the substituting atoms in the molecule has a very great influence on the form of the crystal, but we are very far from being able to connect form and constitution

As a rule the crystals of isomerides exhibit very few analogies, and in many cases none at all. In the benzene series, however, a few relations have been observed, though even here they are rare, and often more readily detected by a comparison of the isomerides with the parent substance than directly with each other. Thus a comparison of the angles given by Bodewig (*P* 158, 232) for the three dinitrobenzenes shows that certain analogies do exist, although the compounds do not crystallise in the same system, while *meta* $\text{C}_6\text{H}_4(\text{NO}_2)_2$ is related to benzene itself, as is shown in the following table —

		a	b	c
C_6H_6	rhombic	0.891	1	0.799
$\text{o C}_6\text{H}_4(\text{NO}_2)_2$	oblique	0.8112	1	0.5735 $\beta = 67^\circ 53'$
$\text{m C}_6\text{H}_4(\text{NO}_2)_2$	rhombic	0.9430	1	0.5354
$\text{p C}_6\text{H}_4(\text{NO}_2)_2$	oblique	0.9883	1	1.0432 $\beta = 87^\circ 42'$

The above examples suffice to show very clearly that the great influence exercised by position renders the discovery of the morphotropic properties of an element or group very difficult. Thus we have stated above that the entry of the nitro- group into benzene leaves the

system and axial ratio $a:b$ unaltered, while the c axis is more or less changed. This statement we found held good for $\text{o C}_6\text{H}_4(\text{NO}_2)_2\text{OH}$, $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OH}(1\ 2\ 4)$, $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OH}(1\ 2\ 4\ 6)$, and $\text{m C}_6\text{H}_4(\text{NO}_2)_2$. As soon, however, as we become acquainted with o and $\text{p C}_6\text{H}_4(\text{NO}_2)_2$, the generality of our statement disappears, and the exact morphotropic value of the NO_2 group remains still undetermined. Although we are not in a position to deduce form from a knowledge of constitution alone, we may often from the chemical analogy of two substances, the form of one of which is known, draw conclusions as to the form of the other, or, *vice versa*, knowing the form of two substances, and the constitution of one of them, we may draw conclusions as to the constitution of the other. This method has been chiefly employed in inorganic chemistry, but Friedlander (*Z K* 3 168) has made use of it in order to throw light on the constitution of the trinitrobenzene got by nitrating $\text{m C}_6\text{H}_4(\text{NO}_2)_2$. The trinitrobenzene made in this way may be either of three theoretically possible ones. It crystallises, however, in the rhombic system, and has the axial ratios $0.954\ 1\ 0.733$, and agrees accordingly with $\text{m C}_6\text{H}_4(\text{NO}_2)_3$. On the other hand it shows little or no analogy to the forms of o or $\text{p C}_6\text{H}_4(\text{NO}_2)_3$, and it is therefore probable that this substance is symmetrical trinitrobenzene $1\ 3\ 5$. A comparison of its form with that of picric acid $\text{C}_6\text{H}_3(\text{NO}_2)_3\text{OH}(1\ 3\ 5\ 6)$, rhombic $0.937\ 1\ 0.974$, confirms this view.

ISOMORPHOTROPY AND ISOMORPHISM. When the products obtained by replacing the hydrogen atoms of the parent substance by two closely allied atoms or groups are compared together they are often found to crystallise in identical or nearly identical forms, such products are isomorphous, while the substituting atoms or groups are said to possess the same morphotropic force, or to be isomorphotropic. Thus, K and NH_4 are isomorphotropic, and replace the H of acids to give isomorphous salts.

The term morphotropy was first applied to cases in which an atom of H was substituted, but the conception may very conveniently be extended to include all cases of substitution whatever, thus, KBr may be regarded as derived from KCl by substituting Br by Cl, and since KCl and KBr both crystallise in the same system and possess very similar physical characters, they are isomorphous, while Cl and Br are isomorphotropic. The greater the analogies between the substituting atoms the greater is the correspondence between the forms of their compounds, i.e. the more closely isomorphous they are, but because two elements are isomorphotropic when in combination with one set of atoms or groups it does not necessarily follow that they are always isomorphotropic, for the morphotropic force of any atom is largely conditioned by the compounds in which it finds itself. Again, Kopp (*P* 53, 446) has pointed out that although it is often true that two isomorphous substances combine with a third to form two isomorphous compounds (Schröder), we cannot, as has been too frequently done, deduce the isomorphism of elements from the isomorphism of their compounds, there are few elements which cannot by such a mode of reasoning be made isomorphous. Elements are only truly isomorphous when they

actually crystallise in the same forms, if they merely combine with another element or group to form isomorphous compounds they may be spoken of as isomorphotropic in that particular class of compounds. As we have seen, the conception of morphotropy includes that of isomorphism. Now, ever since the latter term was introduced, great difficulties have been felt in defining it, and in finding tests which might authoritatively decide in any given case whether two substances were isomorphous or not. Mitscherlich's definition is vague, for we at once ask, What constitutes chemical analogy? And again, How widely may the angles of two substances differ and the substances still be considered isomorphous? The truth seems to be that no hard and fast line can be drawn between those substances which are so closely related morphotropically as to be considered undoubtedly isomorphous, and those slightly less closely related whose isomorphism is doubtful, while we may pass by imperceptible stages from truly isomorphous bodies to others to which, although they still exhibit relations, this term is as evidently inapplicable. Although it seems impossible in the present state of our knowledge to give a definition which shall distinctly mark off isomorphous bodies from those which are merely closely related morphotropically, this is not in practice attended with any very great inconvenience, for the difficulty is chiefly one of nomenclature, most cases of isomorphism are at once recognised as such, and it is only in comparatively few instances that we must hesitate.

In doubtful cases much assistance may often be obtained by a comparison of the other properties of the two substances. Besides great similarity of form, agreement in several other physical properties has been thought necessary before two substances could be considered isomorphous, and we must now notice in detail the more important of these properties, pointing out the weight which has been attached to each as a test of isomorphism.

Physical properties of isomorphous substances

(a) *Geometrical properties*. As the name itself implies, near equality of external form is the first and most essential condition for isomorphism. Absolute geometrical isomorphism, however, does not exist except in the case of substances crystallising in the cubic system, in all other systems small differences of angle occur a fact first noticed by Wollaston. These differences are seldom equally distributed, in rhombic oblique and triclinic crystals the anomalies are often confined to one zone, another zone perpendicular to the first having the same angles in both cases. Here a difficulty meets us, for we ask, How great may these differences become and the substances still remain isomorphous? Unfortunately no definite answer to this question is possible, some authors have admitted much wider differences than others, while others still have gone so far as to admit that the boundaries of the systems may be overstepped. It is perhaps best to restrict the term isomorphous to substances crystallising in the same system with nearly the same angles. It is to be noted that though all substances crystallising in the cubic system have exactly the same angles

they are not necessarily isomorphous, to be so they must be either both holohedral or must both possess the same kind of hemihedrisation, and must also agree in other properties, such as cleavage and power of forming mixed crystals.

(b) *Cleavage*. Isomorphous bodies possess as a rule the same cleavages. Variations in the relative perfection of such cleavages are, however, possible. This property is especially useful in assisting us to detect isomorphism in the cubic system.

(c) *Thermal conductivity*. Jannetaz (*C. R.* 75, 1501) has shown that certain isomorphous substances have axes of maximum and minimum conductivity of like direction and magnitude.

(d) *Coefficient of expansion*. The data are not sufficient for general conclusions to be drawn, and no simple relations have been detected at present.

(e) *Etched figures*. Since these stand in a close relation to crystalline form, Baumhauer has sought to use them as a test of isomorphism. He finds that isomorphous substances give, as a rule, the same sort of etched figures, but that they often differ in orientation.

(f) *Optical properties*. These have been specially compared by Senarmont (*A. Ch.* [3] 33, 391), Topsoe, and Christiansen (*A. Ch.* [5] 1, 5). No general law has been established, but as a rule isomorphous bodies if uniaxial have the same sign of double refraction, and if biaxial have the plane of the optic axes similarly oriented, but biaxial crystals, oblique and triclinic ones especially, exhibit frequent exceptions. Optical properties are, moreover, very sensitive to changes of temperature, which often do not influence isomorphous substances to the same degree (Arzruni, *Z. K.* 1, 165).

(g) *Specific volume*. The relations between chemical composition, S.V., and crystalline form are of special importance, since several authors have considered near equality of S.V. an essential condition for isomorphism. The first of these was Kopp (*A.* 36, 1, P. 47, 133, 52, 243, 262, 53, 446), his views may be briefly summarised as follows: (1) The condition for isomorphism is equality or near equality of S.V. The answer to the question, how widely may the S.V.'s differ and the substances remain isomorphous, is given by the empirical formula

$$D = \frac{v - v_1}{\frac{1}{2}(v + v_1)}, \text{ where } v \text{ and } v_1 \text{ are the S.V.'s of the}$$

two substances. If $D=0$ the isomorphism is perfect, but if D is greater than 33 the substances are not isomorphous, thus the value of D for the two substances ZnCO_3 and CaCO_3 , whose S.V.'s are 28.2 and 36.8 respectively, is 264, they are, therefore, isomorphous. (2) Kopp has also pointed out that the nearer are the axial ratios of a series of isomorphous bodies the nearer are their S.V.'s. This statement, though criticised by Schröder, appears to hold good, and has received the support of Tschermak (*Sts. W.* 46 [2] 603) and Schrauf (*P.* 134, 417). The former expresses the law as follows: 'The series of crystal dimensions and specific volumes is in each group of isomorphous bodies of analogous composition the same'. As an illustration he gives the series of rhombohedral carbonates.

	<i>a</i>	<i>d</i>	
ZnCO ₃	1	0.807	28.2?
MgCO ₃	1	0.812	27.9
(MgFe)CO ₃	1	0.814	29.2
FeCO ₃	1	0.819	30.2
MnCO ₃	1	0.822	31.9
CaCO ₃	1	0.854	36.8

Schröder (*P* 50, 553, 95, 441, 562, 106, 226, 107, 118) at first held views much resembling those of Kopp, but in his later papers he comes to the conclusion that the statements made above are erroneous, and expresses himself thus 'The specific volumes of isomorphous compounds differ in general quite as much from one another as the specific volumes of corresponding heteromorphous compounds' (Schröder uses the term heteromorphous as opposed to isomorphous, it is now often considered synonymous with polymorphous), and he believes that his results lead him to a new law, viz 'If two elements or groups, *A* and *B*, enter into combination with other elements or groups *C*, *D*, *E*, &c., to form the compounds *A* *C* and *B* *C*, *A* *D* and *B* *D*, *A* *E* and *B* *E*, &c., belonging to the same type and isomorphous by pairs, then the differences of the specific volumes of *A* *C* and *B* *C*, *A* *D* and *B* *D*, *A* *E* and *B* *E*, are always equal'. If, however, the pairs of compounds are not isomorphous, or if the isomorphous pairs belong to different types, the differences are as a rule unequal. Schröder terms bodies of equal *SV* *isosteric*, and the equality of the differences of analogous pairs *paralleloterism*. Adopting this nomenclature, the above law may be shortly expressed thus 'Isomorphous analogous pairs of like type are also paralleloteristic'.

Tschermak (*l.c.*) finds that, although Schröder's criticism of Kopp is unfounded, his own results support Schröder's law, which he enunciates thus 'Among isomorphous bodies a like difference of composition corresponds to a like difference of specific volume,' thus the difference *Br* - *Cl* is about equal to 6.3

<i>SV</i>	<i>SV</i>	<i>SV</i>	<i>SV</i>
NaBr 33.4	KBr 44.3	AgBr 31.8	AmBr 42.2
NaCl 27.2	KCl 37.9	AgCl 26.0	AmCl 35.2
6.2	6.4	5.8	7.0

Apparent exceptions to Schröder's law are believed by Tschermak to be due to difference of constitution in the substances compared, and he illustrates this by the following table of difference for the change of composition *K* to *Am*

<i>SV</i>	<i>SV</i>
KCl 37.9	KBr 44.3
AmCl 35.2	AmBr 42.2
2.7	2.1
<i>SV</i>	<i>SV</i>
K ₂ PtCl ₆ 134.5	K ₂ SO ₄ 65.6
Am ₂ PtCl ₆ 150.5	Am ₂ SO ₄ 74.6
16.0	9.0

We see from this table that substances of the type *M'*₂PtCl₆ cannot be directly compared with those of the type *M'*₂SO₄ or these again with *M'*₂O

Tschermak has, moreover, endeavoured to trace a connection between the varying differences shown by pairs of elements which occur

in compounds of similar constitution, and the crystal-system to which these compounds belong.

<i>Cubic</i>	<i>SV</i>	<i>Oblique</i>	<i>SV</i>
Am ₂ PtCl ₆	150.5	Am ₂ Cu(SO ₄) ₂ 6aq	218
K ₂ PtCl ₆	134.5	K ₂ Cu(SO ₄) ₂ 6aq	206
	16.0		18
Am ₂ IrCl ₆	156	Am ₂ Mg(SO ₄) ₂ 6aq	209
K ₂ IrCl ₆	138.3	K ₂ Mg(SO ₄) ₂ 6aq	196
	17.7		13
<i>Tetragonal</i>			
Am ₂ CuCl ₄ 2aq	137	Am ₂ Fe(SO ₄) ₂ 6aq	212
K ₂ CuCl ₄ 2aq	133	K ₂ Fe(SO ₄) ₂ 6aq	197
			15
<i>Rhombic</i>			
Am ₂ SO ₄	74.6	Am ₂ Zn(SO ₄) ₂ 6aq	211
K ₂ SO ₄	65.6	K ₂ Zn(SO ₄) ₂ 6aq	201
	9.0		10

Thus he finds that for the change of composition *Am*₂ to *K*₂, the difference of *SV* is greatest in the cubic system, less in the oblique, still less in the rhombic, and least in the tetragonal system.

In connexion with *SV* we may note that at or about maximum and minimum points of the atomic volume curve (*v* PERIODIC LAW) occur ductile metals crystallising in the cubic system, e.g. Na, Mg, Al, K, Fe, Co, Ni, Cu, Pd, Ag, Pt, Ir, Au, Hg, Pb.

Although isomorphism is usually accompanied by similar values for the *SV*'s, we cannot from the near equality of *SV* alone deduce the isomorphism of two substances, neither can we at the present time attach much weight to Kopp's empirical formula as a test of isomorphism, in a general way, however, his results, in common with those of Tschermak, Schröder, and Schrauf appear to hold good, owing, however, to the great discrepancies which exist between the relative densities of many of the commonest substances as given by different observers, we must be cautious in accepting conclusions often resting on doubtful data.

Special properties of isomorphous substances

(a) *Formation of layer crystals*

(b) *Formation of mixed crystals*

(a) *Layer crystals*. These are made by growing a crystal of one substance in a solution of another.

Kopp (*B* 15, 1653) considers regular growth under these circumstances the best test of isomorphism.

Thus a crystal of common alum when brought into a solution of chrome alum continues to increase regularly, again the sulphates of the type *M''*SO₄ 7aq where *M''* = Zn, Mg, Fe, Co, Ni, grow in solutions of one another.

In those cases where the forms of the two substances are the same the new particles have exactly the same orientation as those of the nucleus. Klocke (*Z* K 2, 144) has, however, brought to light differences between the phenomena of growth in such cases, and in those where a crystal grows in its own solution. Having produced etched figures on the faces of two potassium alum crystals, he grew one in its

own solution, the other in a solution of ammonia iron alum, in the first case the etched figures were rapidly filled up from the bottom, in the second very much flattened octahedra of ammonia iron alum were developed at certain points on the crystal of common alum, and perfectly parallel to it, they then increased laterally, and covered up the etched figures, which remained unaltered till the new layer reached them.

Although the power of forming overgrowths seems to be a necessary consequence of isomorphism, the property is not confined to isomorphous substances alone, and from the perfect conformity of form and orientation exhibited by isomorphous substances, such as the alums, we may pass by imperceptible stages to cases of regular orientation among substances possessing but little chemical analogy and crystallising in different systems. Thus rutile TiO_2 , tetragonal, is often found developed on the basal plane of hæmatite Fe_2O_3 , rhombohedral, in such a way that the c axis of the rutile lies in a plane of symmetry of the hæmatite.

A specially interesting and much discussed case of overgrowth is that of calcite CaCO_3 , rhombohedral $rr' = 74^\circ 55'$ and NaVO_3 , rhombohedral $rr' = 73^\circ 27'$. When a rhombohedron of CaCO_3 is placed in a solution of NaVO_3 , this substance is deposited on the former in small rhombohedra, whose morphological axis and planes of symmetry are exactly parallel to those of the CaCO_3 , these rhombohedra grow till they meet, a polysynthetic crystal being the result. Kopp accordingly made a difference between growth in this case and the regular increase of alum crystals, in the light of Klocke's work, and taking into consideration the slight difference of the angles rr' , such a distinction vanishes, and if the formation of overgrowths is a test of isomorphism, these two substances are certainly isomorphous, a view confirmed by the agreement in their other physical characters, the difference in their chemical constitution leads us rather to regard them as an example of isogonism.

Closely connected with the formation of layer crystals are Thomson's researches on supersaturated solutions (*C J* 1879 196), he has shown that supersaturated solutions of certain salts can readily be crystallised by the addition of crystals of isomorphous salts, while crystals of substances not possessing the same form, or fragments of amorphous bodies, such as glass, have no influence. Thus a supersaturated solution of MgSO_4 , 7aq is at once crystallised by ZnSO_4 , 7aq, NiSO_4 , 7aq, and also by CoSO_4 , 7aq and FeSO_4 , 7aq, whose action, however, is less rapid, on the other hand, Na_2SO_4 , 10aq, NaCl , and glass, are quite inactive. Common alum can be crystallised by iron and chrome alums, while other cubic substances, such as NaCl , FeS_2 , and Fe_2O_3 , are inactive.

This appears to be an excellent test of isomorphism as far as it goes, but it unfortunately is of very limited application.

Although at the present time the formation of layer-crystals cannot be considered a conclusive proof of isomorphism, the success or failure of attempts to obtain them may help us to determine whether two substances crystallising in the cubic system are isomorphous or not.

(A discussion of numerous cases of layer-crystallisation will be found in Lehmann's *Molekularphysik*, Bd 1 pp 393-407, v also Wackernagel, *Köstmers Archiv*, 5, 293, and especially Kopp, *B* 12, 901.)

(b) *Mixed crystals* To obtain mixed crystals solutions of two substances are mixed, or in some cases the two substances are simply melted together, and then allowed to crystallise, we thus obtain perfectly homogeneous crystals, which do not, however, contain their components in any fixed ratio.

As a rule only isomorphous substances yield mixed crystals, and their formation is usually considered the best criterion of isomorphism [Klein (*C R* 95, 781), Kopp (*B* 12, 868, 17, 1105), but v also Brugelmann (*B* 17, 2359), and especially Lehmann (*l c* Bd 1 420, 456, 461)].

Numerous researches have been undertaken with a view to the elucidation of the constitution and mode of formation of mixed crystals, while other investigators have endeavoured to trace the connexion between their physical properties and those of their components.

Growth of mixed crystals Mixed crystals are most readily formed when the two substances have nearly equal solubility, and in this case Rammelsberg (*P* 91, 321) has found that the ratio of the two salts in the mixed crystal is about the same as in the solution, mixed crystals of ZnSO_4 , 7aq and MgSO_4 , 7aq afford a case in point, if, however, as more usually happens, the solubilities differ, the first crystals always contain most of the less soluble salt, the last most of the more soluble, such salts are CuSO_4 , 5aq and MnSO_4 , 5aq Thomson (*l c*) has pointed out that in the case of the crystallisation of supersaturated solutions consisting of a mixture of two substances, the composition of the mixed crystal depends very much on the velocity of crystallisation, if this takes place suddenly the composition of the mixed crystal is much the same as that of the solution, if slowly the less soluble salt is deposited first.

The physical properties of mixed crystals

(a) *Geometrical properties* Mixed crystals belong to the same system as their components, but exhibit simpler forms, cf calcite and dolomite, and v also Rammelsberg (*l c*). In some cases the angles of mixed crystals lie between those of their components, this usually holds good for the naturally occurring mixtures of the rhombohedral carbonates, thus

Chalybite FeCO_3 ,	$rr' = 72^\circ 59'$
Pistomeste (FeMgCO_3),	$= 72^\circ 42'$
Magnesite MgCO_3 ,	$= 72^\circ 33'$

But even in this series exceptions are not wanting, and as a rule no simple relation can be detected between the angles of the mixed crystals and those of its components, thus Groth (*P* 133, 193) in the case of the permanganates and perchlorates found that the angles of the mixed crystals often lay outside those of their components.

KClO_4 , rhombic	$0.7819 \ 1.0 \ 0.6396$
$\text{K} \left\{ \begin{smallmatrix} \text{Mn} \\ \text{Cl} \end{smallmatrix} \right\} \text{O}_4$, "	$0.7797 \ 1.0 \ 0.6408$
$\text{K} \left\{ \begin{smallmatrix} \text{Mn} \\ \text{Cl} \end{smallmatrix} \right\} \text{O}_4$, "	$0.7839 \ 1.0 \ 0.6398$
KMnO_4 , "	$0.7974 \ 1.0 \ 0.6492$

In the crystal containing 1Mn 1Cl the ratio $b:c$ lies inside the limits, and the other two outside, further increase in the quantity of Mn causes the form of the mixed crystal to approach nearer to that of $KClO_4$. Similar results have been obtained by Arzruni on the alkaline earths (*B* 5 1048), Memmar on baryto coelestin (*Min Mit* 1875 59), v Lang on Am_2SO_4 and K_2SO_4 (*Sitz W* 1858 81, 85), Arzruni and Baerwald on compounds of FeS_2 and $FeAs_2$ (*Z K* 7, 337), and lastly by Miers in his investigations of proustite and pyrrargyrite (*Min Mag* 8, 37).

(b) *Optical properties*

(i) *Index of refraction* In the year 1878 Dufet (*C R* 86, 881), from an investigation of $MgSO_4 \cdot 7aq$, $NiSO_4 \cdot 7aq$, and $ZnSO_4 \cdot 7aq$, found that the connection between the chemical composition of isomorphous mixtures and their indices of refraction is one of simple proportionality, so that expressing the results graphically by taking, along two axes at right angles, ordinates proportional to the observed values of μ , and abscissae proportional to the percentage of one component in the mixture, the resulting curve is a straight line. Fock (*Z K* 4, 583) has, however, taken exception to Dufet's work, and although he finds that the changes in the values of the ordinary and extraordinary ray are proportional to the changes in composition for mixed crystals of $SrSO_4 \cdot 4aq$ and $PbSO_4 \cdot 4aq$, yet in mixtures of thallium and potassium alums, and again in mixtures of $MgSO_4 \cdot 7aq$ and $MgCrO_4 \cdot 7aq$, he could find no such law, v also Fitz and Sansoni (*Z K* 6, 67). Fock's results have in their turn been questioned by Soret (*Z K* 11, 197) and Dufet (*C R* 99, 990). The former finds that mixtures of thallium and potassium alum and also of ammonium and potassium alum satisfy the law, while the latter has shown that mixtures of $MgCrO_4 \cdot 7aq$ and $MgSO_4 \cdot 7aq$ are not sufficiently homogeneous for investigations of this kind.

Dufet's law probably holds good, but further investigation is needed before it can be considered established.

(ii) *Optic axis angle* Wyruboff (*Bull Soc Min France*, 2, 91, 170) has endeavoured to trace the connection between the angle of the optic axis and the chemical composition in the cases of mixtures of K_2SO_4 and Am_2SO_4 , and of K_2CrO_4 and K_2SO_4 . His results have been expressed graphically by Mallard (*Bull Soc Min France*, 3, 3), who finds that they give a continuous regular curve, not, however, a straight line. If the optical properties of the components are non accordant, those of the mixed crystals will be different from either, v especially Senarmont (*l c*) on mixed crystals of $KNaH_2C_2O_4 \cdot 4H_2O$ and $(NH_4)NaH_2C_2O_4 \cdot 4H_2O$ —

	Optic axis plane	1st mean line	2nd mean line
Potassium salt	010	a	b
Ammonium	100	c	d

Starting from the potassium salt and substituting with NH_4 , we observe the following changes, first the obtuse angle diminishes and more quickly for red than for violet light, then at a certain point the axes for red light coincide, while the axes for violet light still lie in 010, the red axes then begin to diverge in 100, the violet

afterwards coincide and follow the red. Wyruboff (*Z K* 18, 648), on ammonium and thallium tartrates, gives the following example —

	Optic axis plane	$2H$
Ammonium salt	δ	$42^\circ 8'$
Thallium	1 to b	$59^\circ 30'$

The mixture had always the cleavage and optic axis plane of the ammonium salt, and a crystal containing 88.7 p.c. of the thallium salt had $2H = 43^\circ 30'$.

(iii) *Circular polarisation* Bodlander (*Z K* 9, 309) investigated mixtures of lead and strontium thiosulphates, he found that the rotation varied directly with the composition.

(iv) *Angle of extinction* Max Schuster has shown that the angle of extinction of certain oblique and triclinic mixed crystals, especially the feldspars, varies directly with their chemical composition. His results have been confirmed by Mallard (*Z K* 6, 612), who has treated the subject mathematically.

(v) *Specific volume relations* The specific volumes of mixed crystals appear to depend directly on the relative proportions of their compounds. Thus Schroder states (*P* 95, 441) that

$$V(aCaCO_3 + bFeCO_3) = \frac{aV'(CaCO_3) + bV''(FeCO_3)}{a + b}$$

where V is the specific volume of the mixed crystal and V' , V'' are those of its components. These results have been confirmed by Tschermak (*l c*) thus

$BaCO_3$ (witherrite) rhombic	$V = 45.7$
$CaCO_3$ (aragonite) rhombic	33.9
$(BaCa)CO_3$	= 39.8
while alstonite rhombic	= 39.5

The specific volumes of artificial mixed crystals have been especially investigated by Retgers (*Z P C* 3, 497). He has studied with great care the mixed crystals formed by (a) K_2SO_4 and Am_2SO_4 , (b) potassium and thallium alums, he finds that the SG varies directly with the composition, and expressing his results graphically he obtains as his curve a straight line, a result similar to that obtained by Dufet for the values of μ .

Formation of mixed crystals by isodimorphous substances—Two substances, X and Y, are said to be isodimorphous if they each exist in two forms A and B, A' and B', of which A is isomorphous with A' and B with B'.

On crystallising a solution containing two isodimorphous substances two series of mixed crystals are obtained, one set having the form A, the other the form B. It often happens that the form A of one salt, X say, is stable under ordinary circumstances, while of the other salt, Y, the form B is stable, in the mixed crystals of the form A the substance X predominates, in mixed crystals of the form B, Y is present in excess. In such cases as these a continuous series of mixed crystals cannot always be obtained. Crystallisation under these conditions has been especially studied by Rammsberg (*P* 91, 321). Thus he finds that $MgSO_4 \cdot 7aq$ is rhombic and $FeSO_4 \cdot 7aq$ is oblique, from a solution containing both substances two sorts of mixed crystals are obtained, these do not, however, contain their constituents in all possible proportions, but a gap

occurs, the oblique crystals always having more than 1 atom Fe to 2-3 Mg, while the rhombic crystals always contain more than 4 Mg to 1 Fe. Another good example is afforded by BeSO_4 , 4aq tetragonal and BeSeO_4 , 4aq rhombic. The mixed crystals are tetragonal when $\text{S Se} > 7.33$ 1, and rhombic when $\text{S Se} < 4.1$ (Topsøe, *Sitz W* [2] 66, v also Schulze, *A* 125, 49, Wyrouboff, *Bull Soc Min France*, 2, 91).

Cases of crystallisation perfectly analogous to but differing from these have been studied by Rammelsberg (*l c*), thus copper sulphate usually crystallises in the triclinic system with 5aq, ferrous sulphate in the oblique system with 7aq. When solutions of copper sulphate and ferrous sulphate are mixed and allowed to crystallise, two sorts of crystals are got, both of which contain Cu and Fe, as long as the proportion Cu Fe is > 20 1 the crystals possess 5aq and are triclinic, but when there is less Cu the crystals are oblique and contain 7aq.

It is very commonly supposed, when two substances closely allied chemically crystallise in different forms but yield two kinds of mixed crystals, that these substances are isomorphous, although more than one form of each may not be known, v Fock (*Z K* 6, 160), and note the adverse criticism of Wyrouboff (*Bull Soc Min* 5, 32).

The specific volumes of isomorphous substances have been compared by Rammelsberg (*B* 19, 589), while Retgers (*l c*) has found that in those cases where isomorphous substances form mixed crystals, each series obeys the law enunciated above, viz that the relative density varies directly with the composition.

Constitution of mixed crystals—Since the beginning of the century two opposing views have been held as regards the constitution of these substances.

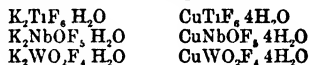
(a) According to the first or physical theory, as we may call it, the most homogeneous mixed crystal really consists of a very intimate mixture of particles of each of the components. This view, held by Haüy, and maintained by Frankenheim in opposition to Berthollet, has recently received the support of Lehmann and Retgers, the latter says that a mixed crystal is a very intimate but purely mechanical mixture of its components, and finds strong confirmation of his view in the fact that the densities, indices of refraction, &c., of mixed crystals vary directly with their composition.

(b) The supporters of the second or chemical theory maintain that molecules of each substance come together in the solution to form a complex 'liquid molecule,' while 'liquid molecules' of the same kind unite on crystallisation and form the mixed crystals. Berthollet held this view, which is perhaps more generally accepted than the other, at the present time its chief exponent is Fock, who believes that the formation of mixed crystals and double salts are phenomena of the same order.

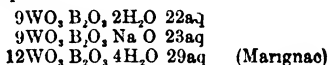
Use of mixed crystals as a test of isomorphism—As we have seen, much weight has been attached to this property by Mitscherlich, Kopp, and others, but at the present time so many instances are known of the formation of mixed crystals by substances which can not be considered isomorphous that the test

has lost much of its value (v Lehmann, *l c*) Retgers (*l c*) has proposed that only those substances should be considered isomorphous, the physical properties of whose mixed crystals vary directly with the chemical composition.

Isogonism—Speaking generally we find that if two substances are to be considered isomorphous they must satisfy the two following conditions (1) they must agree very closely in form and possess similar physical properties, (2) under suitable conditions they must be capable of forming layer and mixed crystals. In Mitscherlich's definition we find, however, a third condition, viz analogy of chemical constitution, and the question arises, How far may two substances which satisfy the two conditions above differ in chemical constitution and still be considered isomorphous? This question has received various answers, thus Scheibler (*J pr* 83, 273) has announced the isomorphism of most meta tungstates, although they do not all contain the same number of molecules of water of crystallisation, Marignac (*C R* 55, 888) considers the two following series of substances to be isomorphous—



And he also makes rhombohedral silicotungstic acid, and the acid silicotungstates of Ba and Ca isomorphous. Some other similar examples are given by Baker (*C J* 1879 760). Klein (*C R* 95, 781) believes that the following substances are isomorphous—



Klein following Marignac, amends Mitscherlich's definition thus 'Isomorphous substances have either an analogous chemical constitution or a very slightly different percentage composition, in the latter case they contain a group of common elements or elements of identical chemical function, this group making up by far the largest portion of the compound.' It seems however, hardly advisable to widen Mitscherlich's definition, indeed, it is perhaps better if possible to narrow it by defining more closely what is meant by the phrases analogous chemical composition and analogous crystalline form.

Substances, such as the double fluorides described by Marignac, are not so much isomorphous as closely related morphologically, while from them we may pass through substances exhibiting less and less of chemical analogy to those which agree in form alone, and which may therefore be considered as examples of isogonism (v Retgers, *l c*). As an instance of this we may quote the remarkable analogy noticed by Hjort dahl (*C R* 88, 584) between the crystals of tin dimethyl and diethyl chlorides and lead chloride, which all crystallise in the rhombic system, and possess the following axial ratios—

	a	b	c
Tin dimethyl chloride	0.8341	1.0	0.9407
Tin diethyl chloride	0.8386	1.0	0.9432
Lead chloride	0.8408	1.0	0.9990

At present it seems that substances which exhibit relations in form and composition may most

conveniently be divided into the following more or less arbitrary groups, between whose limits no absolute line can be drawn —

(1) Substances very closely related morphotopically, or isomorphous substances — Bodies belonging to this class possess great chemical analogy, crystallise in the same system with nearly the same forms and angles, exhibit a general agreement in the physical properties of their crystals, and often possess the same specific volume, they are, moreover, under suitable conditions capable of forming mixed and layer-crystals. Such substances are the sulphates of the group $M''SO_4 \cdot 7H_2O$ ($M'' = Fe'', Zn, Mg, Co, Ni$) and the alums.

(2) Substances closely related morphotopically — To this class belong Marignac's double fluorides, and the substances derived from hydrocarbons by substitution, from them we pass on the one hand to isomerides such as the three dinitro benzenes, which with a minimum difference of chemical constitution possess but little form analogy, and on the other hand to bodies differing but little in form, but showing less and less chemical analogy, till we reach members of the next class.

(3) Isogonous substances, differing greatly in chemical composition but nearly similar in form — To this class belong $CaCO_3$ and $NaNO_3$, and the substances described by Hjortdahl above, while all substances crystallising in the cubic system are strictly isogonous (v. Sohncke, *Entwicklung einer Theorie der Krystalstruktur*, Leipzig, 1879).

ISOPRENE v. PENTINENE

ISOPYRINE An alkaloid obtained by boiling the mashed roots of *Isopyrum thalictroides* with water, filtering, evaporating to a syrup, ppg with ammonia, and extracting with ether (Hartsen, *C* 1872, 523). White powder with bitter taste, forming an amorphous hydrochloride.

Pseudo-isopyrine After the roots have been boiled with water they still contain this alkaloid, which may be extracted with alcohol. After removing the alcohol by evaporation ammonia is added to the aqueous residue, and the ensuing pp treated with ether, whence pseudo isopyrine separates in needles. It is ppd from its solution as hydrochloride by adding very dilute HCl_{aq} and ammonium chloride (difference from isopyrine).

ISUBETIN v. FORMAMIDOXIM

ITABROMOPYROTARTARIC ACID v. BROMO

PYROTARTARIC ACID

ITACHLOROPYROTARTARIC ACID v.

CHLORO-PYROTARTARIC ACIDS

ITACONIC ACID $C_4H_4O_4$, $\pm C_4H_3(CO_2H)_2$, or $CH_2C(CO_2H)CH_2CO_2H$ (?) Mol w 130 [161°] S G 16 (Schröder, *B* 13, 1072) S 5 at 10°, 8 at 20° S (98 p.c. alcohol) 25 at 15° (Baup) R_{∞} 44.25 (in a 4.18 p.c. aqueous solution) (Kanonnikoff, *J* pr [2] 81, 348) H C 476,580 (Lougumine, *O* R 106, 1291) Heat of solution 5,923 (Gal & Werner, *B* [2] 47, 160) Heat of neutralisation 25,725 (G & W) Shown by Raoult's method to be isomeric and not polymeric with citraconic and mesaconic acids (Paternò, *B* 21, 2156).

Formation — 1 By the action of heat on citric acid, acetic acid being first formed (Baup, *A*. 19, 29, Crasso, *A* 34, 68) — 2. By

heating itamallic acid $C_8H_8O_4$ (Swarts, *Z* 1867, 649) — 3 By heating citric acid with water at 160° (Markownikoff & Purgold, *Z* 1867, 264) — 4 By heating acetic acid with water at 180° (Pebal, *A* 98, 94) — 5 By heating citraconic anhydride with water under pressure (Fitting, *A* 188, 72).

Preparation — 1 Coarsely-powdered citric acid (125 g) is distilled from shallow retorts completely filled with it as quickly as the frothing will allow, the oily distillate is separated from the water as completely as possible and left to crystallise (Meilly, *A* 171, 153, cf. Wilm, *A* 141, 28). A temperature of 160°–175° is favourable to the formation of itaconic acid, at a very high temperature it is mostly converted into citraconic anhydride — 2 Citraconic anhydride (2 vols) is heated with water (5 vols) at 150° for 7 hours, and the product poured into flat dishes and left to crystallise (Fitting, *A* 188, 72).

Properties — Trimetric octahedra (from water). Strongly acid. Sol. ether. When heated above its melting point it is resolved into citraconic anhydride and water. Ammonium itaconate gives a brown pp with $FeCl_3$, soluble in excess of $FeCl_3$, but reppd on boiling, and redissolved on cooling.

Reactions — 1 Combines with bromine, forming ita-di-bromo pyrotartaric acid (Kekulé, *A* Suppl. 1, 338, 2, 111, Cahours, *A* Ch [3] 67, 129) — 2 Chlorine forms ita-di-chloro pyrotartaric acid — 3 Heated with aqueous HBr , HCl , and HI , combination takes place with production of ita-bromo-, ita-chloro-, and ita-iodo-pyrotartaric acids. Excess of HI produces pyrotartaric acid (*Bl* [2] 4, 374) — 4 Sodium-amalgam reduces it to pyrotartaric acid (Kekulé) — 5 $HOCI$ gives chloro-itamallic acid $C_4H_3ClO_4$ — 6 Anhydrous HCl at 145° apparently forms an addition-product, which, when distilled, gives citraconic acid (Barbaglia, *B* 7, 465) — 7 Heated with aqueous K_2SO_4 , it forms a gummy salt, possibly sulphopyrotartaric acid $C_4H_3SO_4$ (Wieland, *A* 157, 34) — 8 The electrolysis of the potassium salt yields a mixture of gases, apparently containing propylene and acetylene, but not allylene $CH_2=CH_2$ (Béhal, *A* Ch [6] 16, 366, cf. Aarland, *J* pr [2] 6, 256) — 9 Cold HNO_3 gives off no CO_2 (Franchimont, *R* T C 3, 422).

Constitution — The molecular weights shown by Raoult's method to be the same as that of citraconic and of mesaconic acids. According to Aarland potassium itaconate gives a different allylene on electrolysis from that derived from the potassium salts of citraconic and mesaconic acids, this would lead to the formula $CH_2C(CO_2H)CH_2CO_2H$. Béhal, however, was unable to confirm Aarland's experiments. According to Knops (*A* 248, 228) the molecular refraction of the itaconic ethers indicates one $C=C$ group. Itaconic acid resembles mesaconic acid, but differs from citraconic acid in requiring heat to enable it to combine with HBr and with bromine. Itaconic acid reacts with aniline, but not with dimethylaniline.

Salts — KHA'' aq. lustrous laminae, v. sol. water — K_2A'' does not crystallise — $(NH_4)HA''$ tables S 80 at 12° — $(NH_4)A''$ aq. needles — $BaHA''$ aq. — BaA'' aq. (Petri, *B* 14, 1634) — $CaHA'' \cdot 2aq$ S 77 at 12° — CaA'' aq. needles. S 22 at 18° — $StrA''$ aq. — PbA'' aq. — $PbA''PbO$

(Otto, *A* 127, 181) — $\text{Ag}_2\text{A}''$ almost insol in ling water

Methyl ether $\text{Me}_2\text{A}''$ (212°) SG 15 1 1899 (Anschütz, *B* 14, 2784), Ψ 1 1241 (Knops) R_{20} 60.44 Oil Polymerises, changing to a solid in which $R_{20} = 57.7$

Ethyl ether $\text{Et}_2\text{A}''$ (229°) SG 15 1 051 (Anschütz), Ψ 1 0415 (Knops, *A* 248, 210) $R_{20} = 75.52$ Polymerises $R_{20} = 70.59$ From silver itaconate and EtI (Swarts, *Bull Acad royale Belgique* [2] 36, No 7) Formed also by passing HCl into a solution of itaconic acid in alcohol Colourless oil Changes in a few days into a viscous liquid, and ultimately into a brittle, glassy mass, which is completely decomposed on distillation

Chloride $\text{C}_4\text{H}_4(\text{COCl})_2$ (87°) at 17 mm From the anhydride and PCl_5 Colourless pungent liquid Converted by water into itaconic acid (Petrin)

Anhydride $\text{C}_4\text{H}_2(\text{C}_2\text{O}_3)_2$ [68°] (148°) at 30 mm Occurs among the products of the distillation of citric acid Prepared by the action of AcCl on itaconic acid (Anschütz, *B* 13, 1539) or on silver itaconate suspended in ether (Markownikoff, *B* 13, 1844) Trimeric prisms, $abc = 617.1 \times 455$ Sol chloroform Slowly dissolves in water with formation of itaconic acid When distilled under atmospheric pressure it changes into the isomeric citraconic anhydride

Amide $\text{C}_4\text{H}_4(\text{CONH}_2)_2$ [192°] From $\text{Me}_2\text{A}''$ and aqueous NH_3 (O Strecker, *B* 15, 1639) Small colourless crystals (from water) V e sol hot water, m sol alcohol, insol ether

Mono-anilide $\text{C}_4\text{H}_3(\text{CO}_2\text{H})(\text{CONHPh})$ or $\text{CO}_2\text{H} \text{CH} < \begin{smallmatrix} \text{CH}_2\text{NPh} \\ \text{CH}_2\text{CO} \end{smallmatrix} > (?)$ Itaconanilic acid

[189°] Formed by heating itaconic acid (5 g) with water (50 g) and aniline (3 g) to boiling for half an hour, and depositing on cooling as crystals (Michael & Palmer, *Am* 9, 200, Gotthieb, *A* 77, 265) Formed also by allowing a solution of acid aniline itaconate to stand for several days in the cold Needles (from water) Insol dilute HCl aq, sol dilute alkalis, and reprecipitated unchanged by acids — BaA'_2 (at 170°) — CuA'_2 (at 160°) — AgA'

Mono-anilide $\text{C}_4\text{H}_3(\text{CO}_2\text{H})(\text{CONHPh})$ (?) [151.5°] From itaconic anhydride and aniline (Anschütz, *B* 20, 3215, 21, 958) Not identical with the preceding

Anilide $\text{C}_4\text{H}_3(\text{CONHPh})_2$ [185°] Formed by heating aniline itaconate, or a mixture of its conic acid, with excess of aniline at 180° (Gotthieb, *A* 77, 282, O Strecker, *B* 15, 1639) Large thin plates, sol alcohol and ether, sl sol water A mixture of HNO_3 and H_2SO_4 converts it into an amorphous penta nitro- derivative (Gotthieb, *A* 85, 40)

Chloro-itaconic acid $\text{C}_4\text{H}_3\text{ClO}_4$ Formed by heating conic acid $\text{C}_4\text{H}_3\text{O}_4$ with HCl (Swarts, *J* 1873, 584) Crystalline crusts, sl sol cold water Reconverted by boiling water into conic acid

Bromo-itaconic acid $\text{C}_4\text{H}_3\text{BrO}_4$ [164°] Formed by the dry distillation of its di bromo-pyrotartaric acid $\text{C}_4\text{H}_2\text{Br}_2\text{O}_6$ (Swarts, *Bull Acad roy Belgique* [2] 88, No 1) Also by heating conic acid with HBr Its anhydride is similarly formed from its di bromo-pyrotartaric anhydride (Petrin) Crystals resembling itaconic acid (from water), v sl sol cold water, resolved by boiling

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water or by alkaline carbonates into HBr and conic acid Reduced by tin or zinc to itaconic acid

ITAMALIC ACID v OXY-PYROTARTARIC ACID.

ITAPYRUVIC ACID $\text{C}_4\text{H}_4\text{O}_5$ Obtained, to gether with water and CO_2 , by distilling itatartaric acid at 125°–170° (Wilm, *A* 141, 37) Viscous mass, v sol water and alcohol, volatile with steam — BaA' aq (at 100°) glassy amorphous mass — PbA' , hygroscopic amorphous mass, sl sol water The silver salt is soluble, but very decomposable

ITATARTARIC ACID $\text{C}_4\text{H}_4\text{O}_5$ *Di-oxy-pyrotartaric acid* Formed by gradually adding a solution of HOCl to one of sodium itaconate (2 p c solution) at 0° in the dark, and boiling the resulting solution of chloro oxy pyrotartaric acid The acid is ppd as lead salt, which is subsequently decomposed by H_2S (Wilm, *A* 141, 28, Morawsky, *J pr* [2] 10, 68, 11, 450) Formed also by warming its di bromo pyrotartaric acid with Ag_2O (Kekulé, *A Suppl* 1, 346) Amorphous vitreous deliquescent mass V e sol water and alcohol, not volatile with steam Split up by dry distillation into CO_2 and itapyruvic acid The alkaline salts prevent the ppn of ferric and cupric salts by alkalis — CaA'' aq crystalline mass, sl sol water — BaA'' (at 100°) amorphous, v sol water, and ppd from aqueous solution by alcohol — PbA'' aq monoclinic tablets, v sl sol water — AgA'' bulky pp, sol water, but decomposed on boiling its solution

IVAIN $\text{C}_{20}\text{H}_{32}\text{O}_5$ Occurs in *Iva*, the leaves and stem of *Achillea moschata* gathered before flowering (Von Planta, *A* 155, 150) The volatile oil is first removed by distillation, and the residue is dried and extracted with alcohol, the filtrate is ppd with lead acetate, freed from lead by H_2S , and evaporated Dark yellow resinous mass, insol water, v sol alcohol The alcoholic solution has an extremely bitter taste

IVY *Hedera Helix* When ivy leaves, collected in December, are bruised, exhausted with water, and then treated with alcohol, the alcoholic extract contains a glucoside $\text{C}_{20}\text{H}_{32}\text{O}_{11}$ This may be obtained by evaporating the alcohol, extracting the residue with benzene, and then with acetone, which deposits the glucoside on cooling (Vernet, *C R* 92, 360) It forms mammillary groups of silky colourless needles with slightly sweet taste and neutral reaction. It melts at 233°, and at 22° its alcoholic solution has a levorotatory power of $[\alpha]_D = -47.5^\circ$ It is insol water, chloroform, and petroleum, sl sol acetone, benzene, and ether in the cold, but readily soluble in these liquids when hot. V sol boiling alcohol Sol cold alkalis The glucoside has no action on Fehling's solution, but on boiling with dilute H_2SO_4 a sugar is formed which crystallises by slow evaporation of its alcoholic solution in bulky transparent crystals with decided sweet taste, and reduces Fehling's solution, but is not fermented by yeast The rotatory power of a cold solution of this sugar is $[\alpha]_D = +98.6^\circ$ immediately after its preparation, but some hours afterwards it has been found to fall to $[\alpha]_D = 76.2$ Besides the sugar, there is formed in the hydrolysis of the glucoside a neutral body $\text{C}_{16}\text{H}_{24}\text{O}_5$, [278°–280°], which crys-

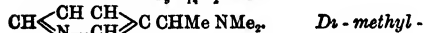
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tallises in slender prismatic needles, without taste, m sol. alcohol, insol. alkalis, and dextro-rotatory $[\alpha]_D = 42.6^\circ$

IXOLITE A fossil resin found in a bed of bituminous coal at Oberhart. Fluid drops, hard-

ening to a hyacinth-red solid, which forms a yellowish powder. SG 1.008. Softens at 78° . Yields pyrocatechin on fusion with potash (Haidinger, *P* 56, 345, Weidel, *Sitz W* [2] 74, 387)

J

JABONINE $C_8H_{11}N_2$, $\epsilon\epsilon$ 

amido ethyl pyridine ($235^\circ - 240^\circ$). Formed by distilling pilocarpine or pilocarpidine with baryta (Hardy a Calmels, *Bl* [2] 48, 231). The crude product is evaporated several times with HClAq and the base set free with KOH. Colourless oil with fetid odour. Its hydro-chloride forms confusedly crystalline masses. The double salts $B'HuCl_2$, $B'AuCl_2$, B_2PtCl_4 , and $B_2H_2PtCl_4$ are all amorphous.

JABORANDI The native name for several drugs of a sudorific and salivating character, consisting of the leaves and twigs of various species of *Pilocarpus* and other trees growing in Brazil, Pernambuco, and Paraguay (Holmes, *Ph* [3] 5, 581, 641, Schelenz, *Ar Ph* [3] 7, 414, Baillon, *Ar Ph* [3] 7, 327). Jaborandi leaves contain pilocarpine (Byasson, *Ph* [3] 5, 826, Hardy, *Bl* [2] 24, 497), which readily changes to jaborine (H Meyer a Harnack, *A* 204, 67). The latter acts like atropine, and is an antidote to pilocarpine. Pilocarpine $C_8H_{11}N_2O_2$ gives with Mel a methyl iodide. Pilocarpidine $C_8H_{11}N_2O_2$ is also present in jaborandi (Harnack, *A* 238, 231). It is crystalline but extremely deliquescent. It is alkaline, and turns yellow in light. v sol. alcohol, m sol. water, sl sol. ether. Its sulphate and hydrochloride are deliquescent, its nitrate forms crystals resembling KNO_3 . Its platinum salt $B_2H_2PtCl_4$ is crystalline. It readily changes when exposed to light into a base jaboridine $C_8H_{11}N_2O_2$, which has an amorphous platinum salt. The jaborandine of Chastaing (*C R* 94, 968) obtained from pilocarpine and HNO_3 is probably jaboridine (Harnack, *A* 238, 238). The jaborandine of Parodi (*Revista farmaceutica*, 1875, 3) from *Piper reticulatum* $C_{10}H_{12}N_2O_2$ may be the same.

JABORIC ACID $C_{10}H_{12}N_2O_2$, $\epsilon\epsilon$ C_8H_9N $CMc(OH)COONMe$, $CMc(C_2H_5N)COH$. Formed, together with NMe_2 , jaborine and pilocarpidine, by rapidly heating pilocarpine to 175° , and keeping it at that temperature for half an hour. If the product be extracted with baryta water and the extract shaken with ether, the ether will contain jaborine, while pilocarpidine and jaboric acid remain in the aqueous solution. In this solution, after removing barium by CO_2 , $AgNO_3$ forms a curdy pp of C_8H_9N , O_2AgNO_3 , whence H_2S removes the silver (Hardy a Calmels, *C R* 102, 1251, *Bl* [2] 48, 225). Resinous mass, v sol. water. Not removed from its aqueous solution by ether. With alkalis it forms gummy salts, sol. water and alcohol, and not decomposed by CO_2 . Hot conc. KOHAq or boiling HClAq converts it into pilocarpidine and oxypyrindyl propionic acid C_8H_9N $CH_2CH(OH)C_2H_5$

Salts — AgA' brown powder, ppd by adding alcohol to a solution of the acid to which a limited quantity of $AgNO_3$ has been added — $AgA'AgNO_3$, curdy pp — $A',PtCl_4$, viscous pp — $A',(PtCl_4)_2$, yellow powder — A',H_2PtCl_4 , sticky pp — $A',(AuCl_3)_2$.

JABORIDINE $C_{10}H_{12}N_2O_2$. Formed by frequently evaporating pilocarpidine with acid (Harnack, *A* 238, 234). Syrup, sol. water, m sol. ether. Acts physiologically like jaborine though weaker. Its hydrochloride is syrupy, and v sol. water — $B_2H_2PtCl_4$ (dried at 100°) [$110^\circ - 120^\circ$]. Amorphous.

JABORINE C_8H_9N , $\epsilon\epsilon$ NMe_2 , $\langle CMc(C_2H_5N)COONMe \rangle$ $COONMe$, (?) Occurs in jaborandi, and in false jaborandi (the leaves of *Piper stictatum*). Prepared as described under JABORIC ACID. Brittle resin. Insol. water, v sol. ether, sol. alcohol. Dissolves in a solution of jaboric acid. Dissolves in HClAq, and is reppd by KOH as a curdy pp which readily agglomerates under warm water. Boiling conc. KOHAq converts it into pilocarpidine. Boiling HClAq does the same. Poisonous, acting like atropine.

Salts — The hydrochloride is amorphous, and v e sol. water and alcohol — B_2PtCl_4 , dirty white gelatinous pp — $B',PtCl_4$, yellowish white pp — $B'H_2PtCl_4$ — $B'(AuCl_3)_2$ (Hardy a Calmels, *C R* 102, 1251).

JALAP The roots and tubers of certain convolvulaceous plants which yield purgative resins. Official jalap consists of the tubers of *Convolvulus Schedanuss*, which contains CONVULVULIN, a strongly purgative resinous glucoside. Jalap wood or jalap stalk (Stiptes Jalapæ), the root of *C. orizabensis*, contains a different glucoside, JALAPIN (Kayser, *A* 51, 100, cf. Mayer, *A* 95, 161).

JALAPIN $C_{24}H_{34}O_{10}$ or $C_{28}H_{42}O_{12}$ [150°]. Occurs in the root stalk of *Convolvulus* (or *Ipomæa*) *orizabensis*, and forms the principal portion (soluble in ether) of the jalap resin prepared therefrom (Johnston, *P T* 1840, 342, *P M* [3] 17, 183, Kayser, *A* 51, 101, Mayer, *A* 83, 122, 95, 129, Haule, *Rep* 48, 365, Planche, *J Ph* 13, 165, 24, 169, Weppen, *N Br Arch* 87, 153, Chevallier, *J Ph* 9, 306). Occurs also in the resin in scammony root from *Convolvulus Scammonia* (Johnston, *P T* 1840, 340, Keller, *A* 104, 63, 109, 209, Spargatis, *A* 116, 289, Planche, *J Ph* 13, 165, 18, 183, Marquart, *N Br Arch* 7, 248, 10, 139).

Preparation — 1 Jalap resin is dissolved in a large quantity of alcohol, water is added until the liquid becomes slightly turbid, and the whole boiled with animal charcoal, filtered, ppd by lead acetate and a little ammonia, filtered, freed

from lead by H_2S , the alcohol distilled off, and the residue kneaded in boiling water, and the jalapin finally extracted with ether (Mayer) — 2 Finely powdered jalap resin is mixed with sand and extracted with ether, which deposits the jalapin on evaporation (Stevenson, *Ph* [3] 10, 644)

Properties — Colourless amorphous resin Softens at 123° , and melts at 150° Tasteless V sol water, v sol alcohol, ether, and chloroform, sol benzene, oil of turpentine, petroleum, and $HOAc$ Conc H_2SO_4 dissolves it slowly in the cold, acquiring in a few minutes a maroon colour, but finally becoming black With K_2CrO_4 , $KMnO_4$, $KClO_4$, or KNO_3 there is a brown colour and a smell of rancid butter, while MnO_2 gives a dark green colour (Stevenson)

Reactions — 1 Boiling dilute HCl splits it up into sugar and jalapinol — 2 Boiling aqueous alkalis convert it into jalapic acid — 3 Nitric acid forms ipomic and oxalic acids

Jalapic acid $C_{12}H_{18}O_{11}$ or $C_{12}H_{16}O_{11}$ [120°] Formed by dissolving jalapin in aqueous alkalis (W Mayer, *A* 92, 125, 95, 129, Spargatis, *A* 116, 289) Prepared by boiling jalapin with baryta-water until the solution is no longer ppd by acids, the baryta is then removed by H_2SO_4 , the excess of sulphate by lead hydrate, and the excess of lead by H_2S Jalapic acid is then deposited on evaporating Translucens, amorphous, yellowish, brittle mass Softens below 100° , and melts about 120° Has no smell, but an irritating taste and strong acid reaction. V sol water, v sol alcohol, m sol ether. In neutral solutions it is ppd by lead subacetate, but not by any other metallic salt

Reactions — 1 fuming HCl splits it up into jalapinol and glucose According to Spargatis the products of hydrolysis of jalapic acid (and jalapin) derived from scammony are jalapinic acid and glucose — 2 Boiling dilute HCl forms (a) jalapic acid, so called (Mayer) — 3 HNO_3 gives ipomic, oxalic, and isobutyric acids — 4 $KMnO_4$ gives isobutyric, oxyisobutyric, and oxalic acids (Poleck & Samelson, *C C* 1884, 813)

Salts — Jalapic acid displaces CO_2 from aqueous Na_2CO_3 — $Ba(C_{12}H_{18}O_{11})_2$ obtained on evaporating a solution of jalapic acid mixed with a slight excess of baryta-water after removing excess of baryta by CO_2 Amorphous — $Ba_2(C_{12}H_{18}O_{11})_2$ obtained by using baryta (2 pts), water (4 pts), and jalapic acid, passing CO_2 through the boiling solution, filtering and evaporating Amorphous mass [100°]

(a) - Jalapic acid, so called $C_{12}H_{18}O_{11}$, Formed, in small quantity, in the hydrolysis of jalapic acid by $HClAq$ or dilute H_2SO_4 (Mayer) On cooling it separates with the jalapinol as a soft brown amorphous mass, which is then boiled with baryta water, and the resulting solution deposits barium jalapinate on cooling When the mother liquor is concentrated silky needles of the Ba salt of (a) jalapic acid are obtained The salt is decomposed by acetic acid White flexible needles, melting below water at 80° Feebly acid in reaction Sl sol cold, m sol hot, water, v sol alcohol and ether Boiling dilute acids split it up into jalapinol and glucose HNO_3 gives ipomic acid and oxalic acid. Potash-fusion forms jalapinic acid and oxalic acid —

$Ba(C_{12}H_{18}O_{11})_2$: White needles, sol. water and alcohol

JALAPINOL $C_{12}H_{20}O_8$ [62°] Formed, together with sugar, by the hydrolysis of jalapin and of jalapic acid (W Mayer, *A* 95, 145) $C_{12}H_{18}O_{11} + 11H_2O = C_{12}H_{20}O_8 + 6C_6H_{12}O_6$ Prepared by allowing a solution of jalapic acid (2 vols) mixed with fuming $HClAq$ (1 vol), to stand for a few days, washing the crystalline product with water, and recrystallising from alcohol, with use of animal charcoal, whence it separates in cauliflower like groups of crystals Insol cold, v sol boiling water, v sol alcohol and ether Aqueous alkalis convert it into jalapinic acid $KMnO_4$ forms isobutyric acid and (by further oxidation) oxyisobutyric acid (Poleck & Samelson, *J* 1884, 1447)

JALAPINOLIC ACID $C_{12}H_{20}O_8$ [64°] Formed by treating jalapinol with aqueous KOH , ammonia, or baryta Formed also by fusing jalapin or jalapic acid with KOH (Mayer, *A* 95, 145, Spargatis, *A* 116, 306) According to Spargatis it is also produced, together with sugar, by the action of mineral acids on jalapin and jalapic acid derived from scammony Poleck and Samelson (*J* 1884, 1447) by the action of alcoholic potash on jalapinol obtained, together with isobutyl alcohol, a jalapinic acid $C_{12}H_{20}O_8$ White tufts of minute thin four sided prisms Insol water, v sol alcohol and ether Lighter than water Has an acid reaction Nitric acid oxidises it to ipomic acid and oxalic acid

Salts — $(NH_4)HA'$, cauliflower heads of minute needles — NaA' (dried at 100°) slender needles — BaA' , (dried at 120°) minute thin white needles, nearly insol cold, sl sol boiling, water, m sol boiling alcohol — CuA' , (dried at 100°) greenish blue pp formed on mixing hot aqueous solutions of sodium jalapinate and $CuSO_4$ (Spargatis) — $Cu_2A'_2(OH)_2$, (dried at 100°) Ppd by adding aqueous cupric acetate to a slightly alkaline solution of ammonium jalapinate Amorphous dark bluish-green mass — PbA' , Ppd by adding lead acetate to an alcoholic solution of jalapinic acid mixed with a little ammonia White amorphous mass — AgA' From aqueous $AgNO_3$, and alcoholic ammonium jalapinate flocculi

Ethyl ether EtA' [82.5°] From the acid (or from scammony) by saturation of the alcoholic solution with HCl , and ppg with water Flat tables

JAMBOSIN $C_{12}H_{19}NO_3$ [77°] Occurs in the bark of jambosa root (the root of *Myrtus jambosa*?), from which it may be extracted with ether (Gerrard, *Ph* [3] 14, 717) White tasteless crystals, sol cold ether, alcohol, chloroform, and hot petroleum ether Insol cold, sol boiling, water Conc H_2SO_4 gives a bright green colour changing to reddish brown Is not a glucoside. Does not form salts with bases Has no medicinal properties

JAPACONINE v ACONITINE ALKALOIDS

JAPACONITINE v ACONITINE ALKALOIDS

JAVANINE v CINCHONA BASES

JECORIN $C_{10}H_{15}N_3SE_2Na_2O_{10}$ (?) A substance obtained from liver (Drechsel, *J pr* [2] 33, 425) Occurs also in ox spleen, human brain, and horse's muscle (Baldi, *Ar Physiol* 1887, Suppl 100) Amorphous substance Hygroscopic, insoluble in dry ether, soluble in wet ether, v sol

sol alcohol Swells up in water forming a sticky mass, which separates into a clear solution and a pp The latter dissolves in much water Not p'd by boiling, p'd by NaCl Prevents p'n of cupric salts by NaOH, forming a blue solution, and on boiling a pp of Cu_2O Boiling HCl decomposes it, forming amongst other products stearic acid

JERVIC ACID An acid occurring in white hellebore root (*Veratrum album*) (Weppen, *Ar Ph* [3] 2, 101, 193), found by E Schmidt (*Ar Ph* [8] 24, 513) to be identical with chelidonic acid

JERVINE $\text{C}_{20}\text{H}_{25}\text{NO}_2$ (Wright & Luff), $\text{C}_{20}\text{H}_{25}\text{N}_2\text{O}_2$ (Tobien) or $\text{C}_{20}\text{H}_{25}\text{N}_2\text{O}_2$ (Will) [237°] (W & L) An alkaloid occurring in the rhizomes of *Veratrum album* (Weppen, *Ar Ph* [3] 2, 101, 193, Simon, *P* 41, 569, Mitchell, *Ph* [3] 4, 796, Bullock, *Ph* [3] 6, 1009, Wright & Luff, *C J* 35, 407), and of *V lobelianum* (Tobien, *Ph* [8] 8, 808) and *V viride* (Bullock, *Ph* [3] 10, 186)

Preparation—The extract obtained with alcohol containing tartaric acid is used several times to macerate fresh portions of the root, freed from alcohol by distillation, mixed with water, fractionally p'd by NaOH and taken up by ether The base first p'd is pseudojervine, and is the least soluble in ether Other fractions contain jervine, rubijervine, and veratralbine (Wright & Luff)

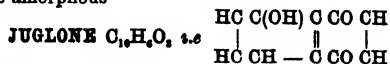
Properties—Loosely coherent crystals (containing $1\frac{1}{2}$ aq or 2aq (from alcohol) Almost insol water, sol alcohol, v sl sol ether (when pure), insol ligroin Conc H_2SO_4 forms a yellow solution changing through brown to greenish-brown Boiling alcoholic KOH has no action

Salts—The hydrochloride is crystalline, sl sol water, v sl sol HClAq (Will, *A* 35, 116) According to Tobien it gives a rose colour when warmed with nitric acid The nitrate and sulphate are sl sol water — $\text{B}^+\text{H}^+\text{Cl}^-$ (Will) — $\text{B}^+\text{H}^+\text{Ac}^-$ —The acetate is sol water

Rubijervine $\text{C}_{20}\text{H}_{25}\text{NO}_2$ [236° cor] Anhydrous crystals (from alcohol) Dissolves in conc H_2SO_4 , forming a yellow solution changing to brownish-red The sulphate is crystalline, v sol cold dilute H_2SO_4 , sl sol cold water The hydrochloride is crystalline and v sol water — $\text{B}^+\text{H}^+\text{Cl}^-$

Pseudojervine $\text{C}_{20}\text{H}_{25}\text{NO}_2$ [209°] Crystals V sl sol ether Reacts with conc H_2SO_4 in the same way as jervine Not affected by boiling alcoholic KOH — $\text{B}^+\text{H}^+\text{Cl}^-$ crystalline, v sl sol hot water, m sol dilute HClAq —The sulphate is sl sol cold, v sol hot, water — $\text{B}^+\text{H}^+\text{Ac}^-$

Veratralbine $\text{C}_{20}\text{H}_{25}\text{NO}_2$ Amorphous resinous base Conc H_2SO_4 forms a yellow solution turning crimson and exhibiting green fluorescence Alcoholic potash does not saponify it Its salts are amorphous



Oxy (a) naphthoquinone *Nucm. Regianum* [154°] Formed by the oxidation of (a) hydrojuglone which occurs in the green shell of the walnut (Vogel & Reischauer, *C C* 1858, 543, *B* 10, 1544, Phipson, *C R* 69, 1872, *C N* 52 89, Mylius, *B* 17, 2411) Formed also by allowing (1,4')-di-oxy-naphthalene to stand for 24

hours mixed with $\text{K}_2\text{Cr}_2\text{O}_7$ and dilute H_2SO_4 (Bernthsen & Semper, *B* 20, 934)

Preparation—Dry ripe walnut shells are repeatedly extracted with cold ether The etheral solution is shaken with a dilute solution of CrO_3 (10 pts $\text{K}_2\text{Cr}_2\text{O}_7$ + 18 pts H_2SO_4 in 500 pts. of water) The ether is distilled off, and the residue, after removing tarry impurities by boiling with a little ether, is crystallised from a mixture of chloroform and ligroin The yield is 150 grms from 100 kilos (Bernthsen & Semper, *B* 18, 204)

Properties—Thin glistening red prisms or needles Sublimable in small quantities Somewhat volatile with steam Easily soluble in chloroform and acetic acid, sparingly in ether, ligroin, and cold alcohol, almost insol water It stains the skin brown Dissolves in dilute NaOH with a fugitive purple colour, in H_2SO_4 with a blood red colour It is decomposed by hot water or hot acids By boiling with HNO_3 it is oxidised to a di nitro oxy phthalic acid On heating with water or HCl it decomposes into insoluble amorphous bodies Its alkaline solution is oxidised by the air to oxy juglone $\text{C}_{10}\text{H}_6\text{O}_5$ With aqueous dimethylamine it gives dimethyl amidojuglone $\text{C}_{10}\text{H}_6\text{O}_4(\text{NMe}_2)_2$ and with alcoholic aniline it gives phenyl amido juglone $\text{C}_{10}\text{H}_6\text{O}_4(\text{NHPh})$ On fusion with KOH it yields m oxy benzic acid (Mylius, *B* 18, 463) Finely powdered juglone gradually added to an alkaline solution of hydrogen peroxide is oxidised to c oxy phthalic acid (B & S) Juglone and its oxim do not dye materials mordanted with metallic oxides (Kostanecki, *B* 22, 1347) Juglone yields naphthalene when distilled with zinc dust Reducing agents yield (a) hydrojuglone

Salt— $(\text{C}_{10}\text{H}_6\text{O}_4)_2\text{Cu}$ nearly insoluble microscopic violet prisms

Acetyl derivative $\text{C}_{10}\text{H}_6\text{O}_4(\text{OAc})$ [155°], flat yellow tables or thin prisms, sublimable in long thin plates, volatile with steam, sparingly soluble in water and cold alcohol, easily in hot alcohol, ether, benzene, chloroform, and CS_2 , yields an intense green colouration with alcoholic KOH

Mono oxim $\text{C}_{10}\text{H}_6\text{O}_4(\text{OH}) \begin{cases} \text{O} \\ \text{NOH} \end{cases}$ [187°]

Glistening red needles or thin prisms, easily soluble in alcohol and acetic acid, very slightly in water, dissolves with a blood red colour in H_2SO_4 and in dilute NaOH (Bernthsen & Semper, *B* 18, 203)

Di-oxim $\text{C}_{10}\text{H}_6\text{O}_4(\text{OH})(\text{NOH})_2$ From juglone (1 mol) and hydroxylamine hydrochloride (2 mols) at 140° (Bernthsen & Semper, *B* 19, 168) Brownish yellow needles (from HOAc) Puffs off at 225° Sl sol alcohol and acetic acid Alkali dissolve it, forming an orange solution

Phenyl-amido juglone $\text{C}_{10}\text{H}_6\text{O}_4(\text{NHPh})$ [230°] Formed by heating an alcoholic solution of juglone and aniline (Mylius, *B* 18, 472) Red tables Sublimable V sl sol alcohol Dissolves in alkalis with a purple red colour

Di-methyl-amido-juglone $\text{C}_{10}\text{H}_6\text{O}_4(\text{NMe}_2)_2$ [150°] Formed by dissolving juglone in dimethylamine solution (Mylius, *B* 18, 464) Brownish-violet tables Soluble in benzene, chloroform, and CS_2 , sparingly in alcohol, ether, and acetic acid, insoluble in water By SnCl_4 it

is reduced to dimethylamido hydrojuglone; On heating with strong HCl it splits off dimethylamine, giving oxy juglone

Oxyjuglone $C_{11}H_7O_3$, s.s.

$CH_3C(OH)C(=O)C(OH)C(=O)CH_3$

Di-oxy-naphthoquinone $C_{10}H_6O_4$

$CH_3CH=CH-C(=O)CH=CH-C(=O)CH_3$

one [c 220°] Formed by heating di methylamido juglone with HCl. Formed also by oxidation of an alkaline solution of juglone by exposure to the air. Prepared by oxidation of juglone with an alkaline solution of potassium ferricyanide, yield 50 p.c. of theory (Mylus, B 18, 466). Small yellow trimetric tables. Sparingly soluble in alcohol, ether, benzene, and CS_2 .

Salts —A'Na₂ easily soluble red needles —A'K₂ sparingly soluble red needles —A'Ag₂ —A'Cl₂ amorphous red pp

Benzoyl derivative $C_{13}H_8O_4(OBz)_2$, [170°], small yellowish white crystals, easily

soluble in benzene, sparingly in alcohol and acetic acid, insoluble in water

JUGLOXIM v OXIM of JUGLONE.

JUGLONIC ACID v DI NITRO-OXY PHTHALIC ACID

JUNIPER OIL The berries of the juniper, which are used for flavouring gin, yield on distillation an essential oil $C_{10}H_{16}$, (155°-163°) S G 25 839 (Blanchet, A 7, 165, Souberran a Capitaine, A 34, 325). The oil is levorotatory, it gives no solid bromide, but after heating the oil to 260° it is partly changed to a product boiling about 180°, and this gives cinenetetrabromide. According to Wallach (A 227, 288) it contains pinene (v TERPENES). The product of distillation of the wood of the juniper contains a sesquiterpene $C_{15}H_{24}$, which forms a hydrochloride $C_{15}H_{21}H_2Cl_2$ melting at 118° (Wallach, A 238, 82).

JUTE v CELLULOSE.

K

KACOTHELINE $C_{11}H_{12}O_4N_2$ (not $C_{11}H_{12}O_4N_2$) s.s. $C_{11}H_{12}(NO_2)_2O_4N_2$. Prepared by dissolving brucine in an excess of cold nitric acid (1, 2 S G) and warming to 50°-60° till the solution has become yellowish red, the kacotheline separates out in small crystals (cf vol 1 p 654). On oxidation with CrO_3 and H_2SO_4 it yields the same product $C_{11}H_{10}N_2O_4$, as brucine. By treatment of kacotheline, suspended in boiling HCl, with bromine water it is converted into an acid $C_{11}H_{12}N_2O_4$. This acid by further oxidation with CrO_3 and H_2SO_4 yields the above oxidation product $C_{11}H_{10}N_2O_4$. On reduction with tin and HCl it yields a base $C_{11}H_{12}(NH_2)_2N_2O_4$ (Hanssen, B 20, 452).

Base $C_{11}H_{12}N_2O_4$, s.s. $C_{11}H_{12}(NH_2)_2N_2O_4$, [232°] Formed by reduction of kacotheline with tin and HCl. Needles. Atmospheric oxidation or $FeCl_3$ produces a blue black colouration. Insoluble in water and alcohol, dissolves in aqueous NaOH with a reddish yellow colour. Its methyl iodide forms glistening plates easily soluble in water —B'H₂Cl₂ large glistening crystals (Hanssen, B 20, 453).

KAIRINE v (B 4) OXY (Py 4) METHYL-QUINOLINE TETRAHYDRIDE

Kairine A v (B 4) OXY (Py 4) ETHYL QUINOLINE TETRAHYDRIDE

KAIROCOLL $C_{11}H_{11}NO_4$, s.s.

$C_6H_5(OH) \begin{cases} CH_2CH_2 \\ | \\ N-CH_2 \end{cases}$ [66°] Long fine needles

CH_2CO_2H
Sol alcohol and ether, al sol water. Prepared by heating (B 4) oxy quinoline tetrahydride with chloro acetic acid (O Fischer, B 16, 718).

KAIROLINE is described as METHYL-QUINOLINE TETRA-HYDRIDE

Amido-kairolin v AMIDO (Py 4) METHYL-QUINOLINE TETRAHYDRIDE

Nitro-kairolin v NITRO-(Py 4)-METHYL-QUINOLINE TETRA-HYDRIDE.

KAIROLINE m CARBOXYLIC ACID v (Py 4) METHYL QUINOLINE TETRA HYDRIDE (B 3)-CARBOXYLIC ACID

KAKOSTRYCHNINE v CACOSTRYCHNINE.

KAMALA A yellow dye, used in India, contained in the seed capsules of *Mallotus phillyensis*, and occurring in commerce as a yellowish brown powder composed of minute resinous globules. It contains MALLOTOXIN (q v) (A. G. a. W. H. Perkin, jun., B 19, 3109).

KAOIRI GUM The product of a coniferous tree (*Dammara australis*) growing in New Zealand. When distilled with steam it yields a terpene $C_{10}H_{16}$, (157°-158°), S G 25 863 (Rennie, C J 39, 240, cf Thomson, A Ch [3] 9, 499; Muir, C J 27, 733). It is feebly levorotatory, and yields a small quantity of cymene when treated with PCl_5 .

KAWAIN A crystalline resin, occurring along with METHYSTICIN (q v) in Kawa kawa, the root of *Macropiper methysticum* (Gobley, J Ph. [3] 37, 19, O'Rourke, C R 50, 598, Cuzent, C R 50, 436, 52, 206, Nölting a Kopp, Monat. Scient [3] 4, 9, 20). It is not a glucoside. On oxidation it yields benzoic acid.

KELLIN A glucoside in *Ammi Visnaga* (Mustapha, C R 89, 442). Small silky needles, v sl sol cold water, sol alcohol, v sol ether. Has a bitter taste. Gives a white pp with Nessler's reagent. Emetic and narcotic.

KEPHIR v MILK

KERATIN v PROTEIDS, Appendix C

KETATES (ethers of ketones) Bodies constructed on the type $\begin{matrix} X \\ | \\ Y \end{matrix} > C < \begin{matrix} O R' \\ | \\ O R'' \end{matrix}$ (Morley a. Green, B 17, 8015). Bodies of the form $\begin{matrix} X \\ | \\ Y \end{matrix} > C < \begin{matrix} SR' \\ | \\ SR'' \end{matrix}$ or $\begin{matrix} X \\ | \\ Y \end{matrix} > C < \begin{matrix} OR' \\ | \\ SR'' \end{matrix}$ may be called thioketates.

KETINE A name given by V Meyer and Treadwell (B 14, 1150, 15, 1059, 1055) to the product of the reduction of nitroso-acetone with tin and HCl. Its di-carboxylic ether is formed

in like manner by reducing nitroso-aceto-acetic ether Ketine C_6H_5O , appears to be di-methyl-
 $CH_3N CMe CH N CMe$

pyrazine $\begin{array}{c} || \\ CH_3N CH \\ || \end{array}$ or $\begin{array}{c} || \\ CH_3N CH \\ || \end{array}$, and will be

described as such (*cf* Oecononimides, *B* 19, 2524, Japp, *C J* 51, 98) Meyer (*B* 21, 19) proposes to use the name aldine instead of pyrazine

for the group $\begin{array}{c} CH N CH \\ || \end{array}$ or $\begin{array}{c} CH N CH \\ || \end{array}$ Other

alkyl-pyrazines are formed in like manner by the reduction of the nitroso-derivatives $HO N CH CO C_nH_{2n+1}$, of ketones of the form $CH_3 CO C_nH_{2n+1}$. As ketine di carboxylic acid does not form an anhydride it has probably the

formula $\begin{array}{c} || \\ CO_2H C - N CMe \\ || \end{array}$ rather than

$\begin{array}{c} || \\ CO_2H C - N C CO_2H \\ || \end{array}$, whence the symmetrical formula

for ketine given above (Oecononimides)

KETIPIC ACID A name given to Di METHYLDIKETONE DI-CARBOXYLIC ACID, as it might perhaps be called di keto adipic acid

KETO This prefix is employed by some chemists to denote the displacement of H_2 in the group C_2H_4 by O . Thus quinone might be called di keto benzene, while aceto acetic acid would be β keto-butyric acid. The prefix 'keto' does not indicate the introduction of a new or ganic group, whereas the corresponding prefix 'aldehydo-' indicates the introduction of the group CHO , and 'carboxy' is used to denote the introduction of CO_2H . In order to avoid the confusion likely to arise from this circumstance the term 'keto' is avoided in the headings of articles in this Dictionary substances which might have such names given them are described as ketones, quinones, or oxy compounds. The prefix keto- has also been used as indicating substitution of hydrogen by the ketonic group $CO CH_3$, thus Erdmann (*B* 21, 635) terms $CH_3 CO C_6H_5 OH$ keto naphthol. In keto compounds the group $CO OH$ may often be regarded with equal propriety as $C(OH)C$, and such compounds are named in accordance with the latter formula. Thus phloroglucin may be described as tri-keto-benzene hexahydrate or as tri-oxybenzene

DI-KETO- HEPTANE *v* METHYL BUTYL DIKETONE

DI-KETO-HEXANE *v* METHYL PROPYL DIKETONE

α - β -DIKETO-HEXYLENE *v* METHYL ALLYLDIKETONE

KETO-HEXA HYDRO-BENZOIC ACID CY ANHYDRIN *v* Nitrile of Oxy ISOPHTHALIC ACID HEXAHYDRIDE

DI-KETO-HYDRINDENE *v* INDONAPHTHOQUINONE

DI-KETO-INDONAPHTHENE *v* INDONAPHTHOQUINONE

KETO-LACTONIC ACID $C_5H_6O_4$, [181°] When α -acetyl- β -ethyl-succinic ether is heated it partly breaks up into alcohol and ethyl ketolactonate, which, on saponification, gives potassium ketolactonate (*S* Young, *C J* 43, 175, *A* 216, 45). It is also formed, together with $CO_2H.CHEt.CH_2Ac$, by boiling the ether

($CO_2H.CHEt.CH_2Ac$) with dilute HCl , and may be extracted from the product by ether. The free acid crystallises from water, and is al sol cold water, m sol hot water

Salts — $BaA', 2H_2O - AgA'$

Reaction — 1 Heated with baryta the following reaction occurs $2C_5H_6O_4 + 3BaO + H_2O = 2BaCO_3 + Ba(C_5H_3O_4)_2$. The new acid, $C_5H_3O_4$, appears to be liquid, it forms a very soluble and unstable silver salt — 2 Cold baryta produces an unstable dibasic acid, $C_5H_3O_4$, by assumption of H_2O

Constitution — Ketolactonic ether is formed from $CO_2Et.CHAc.CHEt.CO_2Et$ by splitting off $HOEt$. Inasmuch as $CO_2Et.CMeAc.CHEt.CO_2Et$ does not form in the same way the ether of a crystalline acid it is probable that the H which is here displaced by methyl goes to form alcohol in the formation of the ketolactonic ether. And since ketolactonic ether appears to be a lactone it should be

$CO_2Et.C.CHEt.CO_2$,

the acid being $\begin{array}{c} CH_3C O \\ || \end{array} CO_2H.C.CHEt.CO_2$

$\begin{array}{c} CH_3C O \\ || \end{array} - -$

KETOLE Another name for INDOL (*Jackson, B* 14, 879)

KETONES The ketones, in their simplest form, contain a carbonyl group CO attached to two monad hydrocarbon radicles. If the two radicles are identical, as in acetone (dimethylketone) $CH_3.CO.CH_3$, the compound is a simple ketone, if different, as in methyl ethyl ketone $CH_3.CO.CH_2CH_3$, it is a mixed ketone

Just as the fatty acids of the series $C_nH_{2n}O_2$ may be derived from the hypothetical carbonic acid $CO(OH)_2$, by replacing one hydroxyl group by a monad hydrocarbon radicle, so the ketones may in turn be derived from the fatty acids by replacing the remaining hydroxyl group by a second radicle

$HO.CO.OH$ $CH_3.CO.OH$ $CH_3.CO.CH_3$
Carbonic acid Acetic acid Acetone

There is therefore a simple ketone derivable from each fatty acid by the introduction of a hydrocarbon radicle, identical with that which is attached to the carboxyl group of the acid, in place of its hydroxyl group

If one of the radicles in a ketone is hydrogen the resulting compound is an aldehyde

$CH_3.CO.H$ $CH_3.CO.CH_3$
Aldehyde Acetone

and owing to this similarity in constitution, the aldehydes and ketones have many reactions in common. In the case of formic acid the radicle attached to carboxyl is hydrogen, and the ketone therefore coincides with the aldehyde

$H.CO.OH$ $H.CO.H$
Formic acid Formaldehyde.

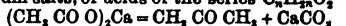
From this point of view the ketones may be regarded as homologues of formaldehyde

Compounds containing two carbonyl groups are termed *diketones*, those containing three, *triketones*

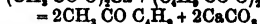
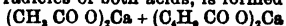
Fatty ketones

Formation. — 1 By the destructive distilla-

tion of the calcium salts, or, better still, of the barium salts, of acids of the series $C_nH_{2n}O_2$.

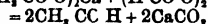
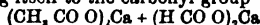


If a mixture of the salts of two acids is employed, a mixed ketone, containing the hydrocarbon radicles of both acids, is formed



(Williamson, *C J* 1852, '38)

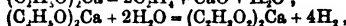
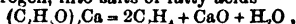
If one of the salts is a formate, an aldehyde is formed, a hydrogen atom from the formate attaching itself to the carbonyl group



(Piria, *Cimento*, 3, 126, *A* 100, 104, Lumprich, *A* 97, 5-8)

In the preparation of ketones by the destructive distillation of salts it is advantageous, if the molecular weight of the ketone is high, to conduct the distillation under reduced pressure (Kraft, *B* 15, 1693)

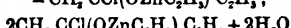
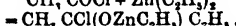
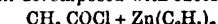
2 By the destructive distillation of the calcium compounds of primary alcohols (Destrem, *A Ch* [5] 27, 7) In this reaction the alcoholates are converted, with evolution of olefins and hydrogen, into salts of fatty acids



and the calcium acetate is then decomposed by the heat, yielding acetone. A portion of the acetate, however, reacts with the calcium oxide, evolving methane

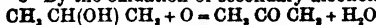
3 By heating a mixture of the sodium compound of an alcohol and the sodium salt of a fatty acid in a current of carbon monoxide, mixtures of higher ketones are formed, along with sodium salts of higher organic acids (Genther, *A* 202, 288) The constitution of these ketones is not known with certainty, and the process in which they are formed is not understood

4 By the action of acyl chlorides on zinc alkyls (Freund, *A* 118, 1, cf Pawlow, *A* 188, 104) The acyl chloride is added to the zinc alkyl, at first gradually, afterwards more rapidly, cooling with ice during the operation. The mixture is then decomposed with excess of water



If the mixture is allowed to stand too long before adding water, the additive compound formed in the first stage of the process reacts with a second mol of alkyl chloride, exchanging its halogen for alkyl and generating a compound which, on treatment with water, yields a tertiary alcohol

5 By the oxidation of secondary alcohols



6 By the oxidation of secondary acids of the lactic series with potassium bichromate and dilute sulphuric acid. Thus hydroxy-isobutyric acid yields acetone

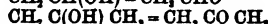
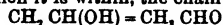


7 Certain ketonic acids part with carbon dioxide, forming ketones. Thus decomposition takes place very readily in the case of the β -ketonic acids. Thus when ethylic acetoacetate is warmed with a caustic alkali it is

hydrolysed, and the liberated acid splits off carbon dioxide, thus

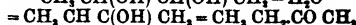
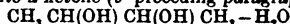


8 Alcohols containing a hydroxyl-group attached to a doubly-linked carbon-atom do not appear to be capable of existing in the free state, and change, at the moment of their formation, into either aldehydes or ketones, the former transformation occurring when the hydroxyl group is situated at the end of, the latter when it is within, the chain

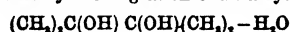


(Erlenmeyer, *B* 13, 809, 14, 320)

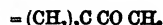
9 When secondary or secondary tertiary glycols are treated with dehydrating agents, such as phosphorus pentoxide or zinc chloride, they part with water, forming ketones. Probably an unsaturated alcohol is first produced, and this changes, at the moment of its formation, into a ketone (v preceding paragraph).



Primary glycols, when thus treated, yield aldehydes. Tertiary glycols (pinacones) are dehydrated even by boiling with dilute sulphuric acid, forming ketones, the reaction being accompanied in this case by the migration of an alkyl group

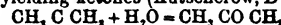


Pinacone



Pinacol

10 When the homologues of acetylene are passed into a solution of mercuric chloride heated to 90°-95° they take up the elements of water, yielding ketones (Kutscherow, *B* 17, 13)



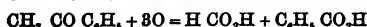
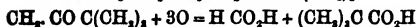
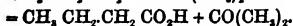
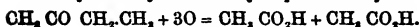
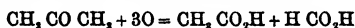
Acetylene itself yields aldehyde

11 Ketones, especially acetone, are formed in the destructive distillation of wood, sugar, citric acid, and various other organic compounds

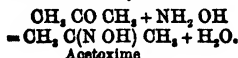
Properties and Reactions—The lower members of the ketone series are liquids, with a peculiar ethereal odour, boiling without decomposition. The higher homologues, beginning with $C_8H_{16}O$, are crystalline solids

The ketones are isomeric with the aldehydes containing the same number of atoms of carbon. They closely resemble the aldehydes in many of their reactions, but are distinguished from them by not reducing ammoniacal silver solutions. Like the aldehydes, many of the ketones combine with the *hydrogen sulphites of the alkalis* to form crystalline double compounds. The rule given by Grimm (*A* 167, 262) that only those ketones combine with hydrogen sodium sulphite which contain the group CH_3, CO , holds very generally an exception is propione (diethyl ketone) $CO(C_2H_5)_2$, which forms a double compound, although with difficulty. These compounds are used in separating and purifying the ketones on treating the double compound with sodium carbonate the ketone is liberated. The ketones, like the aldehydes, unite with *hydrocyanic acid* to form nitriles of hydroxy-acids $(CH_3)_2CO + HCN = (CH_3)_2C(OH) CN$ (Urech, *A* 164, 268, Tiemann & Friedländer, *B* 14, 1971) and these nitriles react with ammonia, yielding nitriles of amido-acids $(CH_3)_2C(OH) CN + NH_3 = (CH_3)_2C(NH_2) CN + H_2O$ (T & F.).—In

presence of caustic potash, acetone unites with *chloroform* to form acetone-chloroform $(\text{CH}_3)_2\text{C}(\text{OH})\text{COCl}$, which when heated with water to 160° yields hydroxy-isobutyric acid and hydrochloric acid (Willgerodt, *B* 14, 2451, 16, 1585) — *Sodium amalgam*, acting on aqueous solutions of the ketones, reduces them to secondary alcohols, thus acetone is converted into isopropyl alcohol $(\text{CH}_3)_2\text{CHOH}$, $\text{CO} + \text{H}_2 = (\text{CH}_3)_2\text{CHOH}$ (Friedel, *C R* 55, 53) At the same time, a second reaction occurs in which 2 mols of the ketone are united during the process of reduction, forming a tertiary glycol, or pinacone $2(\text{CH}_3)_2\text{CO} + \text{H}_2 = (\text{CH}_3)_2\text{C}(\text{OH})\text{C}(\text{OH})(\text{CH}_3)_2$. — The ketones are less susceptible of oxidation than the aldehydes, and, unlike the latter, can never yield by oxidation an acid containing the same number of carbon atoms. They are best oxidised by boiling them with a mixture of potassium bichromate and dilute sulphuric acid. Popoff (*A* 161, 300) has formulated the following rules regarding the products formed in the oxidation of the ketones — rules to which, however, there are occasional exceptions. The ketone is broken up at a point between the carbonyl group and one of the alkyl groups, the combined carbonyl being oxidised to carboxyl, and the separated alkyl being oxidised, if primary, to a fatty acid with the same number of atoms of carbon as the alkyl itself, and if secondary, to a ketone, which may in turn be further oxidised. If a ketone contains two dissimilar alkyls, the carbonyl group will remain attached to that carbon atom which has most hydrogen combined with it, unless in the case of a tertiary alkyl or a radicle of the benzene series, when the carbonyl group will remain in combination with the non hydrogenated carbon atoms. If the two dissimilar alkyls are both primary, or both secondary, or both tertiary, the carbonyl will remain attached to the alkyl of lower molecular weight. The following equations illustrate these rules



In several cases, however, a subordinate reaction, in which the ketone is oxidised in a way the reverse of that predicted by the rule, occurs simultaneously — *Chlorine* and *bromine* give substitution compounds — *Phosphorus pentachloride* replaces the oxygen of ketones by two chlorine atoms — *Ammonia* reacts less readily with ketones than with aldehydes. With acetone it forms diacetoneamine $\text{C}_6\text{H}_{11}\text{NO}$, and triacetoneamine $\text{C}_9\text{H}_{17}\text{NO}$ — With *hydroxylamine* in aqueous solution the ketones yield the ketoximes.

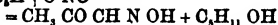
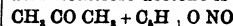


The ketoximes are generally solid crystalline compounds, volatile without decomposition. Concentrated hydrochloric acid decomposes them into hydroxylamine and ketone. — In like

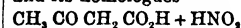
manner *phenylhydrazine* reacts with the ketones, eliminating water and forming ketohydrazones: $\text{CH}_3\text{COCH}_3 + \text{C}_6\text{H}_5\text{NH NH}_2$



The ketohydrazones of the fatty series are for the most part only liquids, which may be distilled under reduced pressure. Acids decompose them into phenylhydrazine and ketones (E. Fischer, *B* 16, 661) — *Nitrous acid* converts the ketones into isonitroso ketones. The reaction occurs more readily, however, with ethereal salts of nitrous acid, thus when a mixture of amyl nitrite and acetone is warmed with hydrochloric acid isonitroso acetone is formed



Sodium ethoxide may be substituted for hydrochloric acid in the foregoing reaction, but in this case the sodium compound of the isonitroso ketone is formed and must afterwards be decomposed by acetic acid (Claisen, *B* 20, 252, and 656, Claisen a. Manasse, *B* 20, 2194). The isonitroso ketones are also readily obtained by the action of nitrous acid on acetoacetic acid and its homologues



Acetoacetic acid

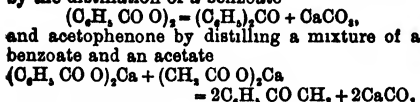


(V. Meyer a. Zublin, *B* 11, 695, Ceresole, *B* 15, 1328) — Concentrated *nitric acid* converts the ketones into dinitro paraffins, thus propione $\text{CH}_3\text{CH}_2\text{COCH}_2\text{CH}_3$ yields dinitroethane $\text{CH}_3\text{CH}(\text{NO}_2)_2$ (Chancel, *Bl* 31, 504) — The ketones react with the *mercaptans*, eliminating water and yielding 'mercaptoles' (thioketates), ethyl thiodimethylketate is formed when gaseous hydrochloric acid is passed into a mixture of acetone and mercaptan $(\text{CH}_3)_2\text{CO} + 2\text{C}_2\text{H}_5\text{SH} = (\text{CH}_3)_2\text{C}(\text{S C}_2\text{H}_5)_2 + \text{H}_2\text{O}$ (Baumann, *B* 18, 887). The mercaptoles are liquids, insoluble in water, not volatile without decomposition, stable towards alkalis and dilute acids. — Under the influence of *dehydrating agents*, such as sulphuric acid, zinc chloride, hydrochloric acid, &c., the ketones readily undergo condensation, two or more molecules combining, with elimination of water, to form more complex compounds. Thus 2 mols of acetone unite to form mesityl oxide $(\text{CH}_3)_2\text{C CH CO CH}_3$, and 3 mols to form either phorone $(\text{CH}_3)_2\text{C CH CO CH CO CH}_3$, or mesitylene (1, 3, 5 trimethylbenzene) $\text{C}_6\text{H}_2(\text{CH}_3)_3$, according as 2 or 3 mols of water are eliminated in the process.

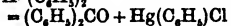
Benzene ketones. Ketones of this class may either contain two benzene radicles directly attached to carbonyl, as in benzophenone (diphenyl ketone) $\text{C}_6\text{H}_5\text{CO C}_6\text{H}_5$, or the carbonyl group may unite a benzene radicle and a fatty radicle, as in acetophenone (phenyl methyl ketone) $\text{C}_6\text{H}_5\text{CO CH}_3$, or one or both of the benzene radicles may be attached to carbonyl by means of a fatty group, as in phenyl benzyl ketone $\text{C}_6\text{H}_5\text{CO CH}_2\text{C}_6\text{H}_5$, or di benzyl ketone $\text{C}_6\text{H}_5\text{CH}_2\text{CO CH}_2\text{C}_6\text{H}_5$.

Preparation. — Many of the methods already described for the preparation of the fatty ketones are also applicable in the case of the benzene

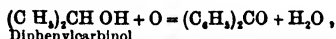
ketones Thus benzophenone may be obtained by the distillation of a benzoate



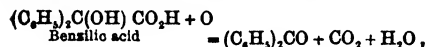
Again, acetophenone may be obtained by the action of benzoyl chloride on zinc methyl, but if it is desired to introduce a benzene radicle in place of the halogen of the acyl chloride the mercury compound of that radicle must be employed $\text{C}_6\text{H}_5, \text{COCl} + \text{Hg}(\text{C}_6\text{H}_5)_2$



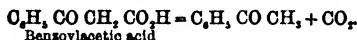
The benzene ketones are also formed, like the fatty ketones, by the oxidation of the corresponding secondary alcohols



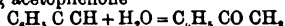
and of secondary glycolic acids containing benzene radicles



further, by splitting off carbon dioxide from ketonic acids of the benzene series



By shaking phenylacetylene with sulphuric acid of 75 p.c. it takes up the elements of water, yielding acetophenone

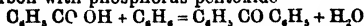


(Friedel & Baisohn, *Bt* [2] 85, 55)

The following modes of formation are peculiar to benzene ketones —

1 By the action of an acyl chloride on a benzene hydrocarbon in presence of aluminium chloride $\text{CH}_3, \text{COCl} + \text{C}_6\text{H}_6 = \text{CH}_3, \text{CO C}_6\text{H}_5 + \text{HCl}$ (Friedel & Crafts, *A Ch* [6] 1, 507) In this reaction the acid radicle always attaches itself to a benzene nucleus, and never to a fatty group. In like manner ketones containing two benzene radicles directly united to carbonyl may be prepared by acting on a benzene hydrocarbon with carbonyl chloride in presence of aluminium chloride $\text{COCl}_2 + 2\text{C}_6\text{H}_6 = \text{CO}(\text{C}_6\text{H}_5)_2 + 2\text{HCl}$

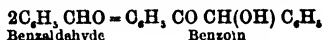
2 The last mentioned class of ketones may also be obtained by heating a mixture of a benzene carboxylic acid and a benzene hydrocarbon with phosphorus pentoxide



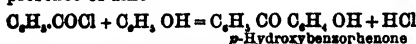
3 By the oxidation of hydrocarbons in which two benzene radicles are united by a methylene group



Ketone alcohols are obtained by polymerising benzene aldehydes with potassium cyanide



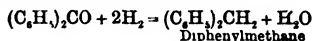
Ketone phenols are formed when acyl chlorides of the benzene series act on phenols in presence of zinc



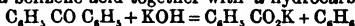
The acid radicle enters the nucleus of the phenol in the para- position to the hydroxyl group

Properties and Reactions — The benzene ketones are liquids and solids, boiling without decomposition and having a pleasant aromatic odour

In most of their reactions they closely resemble the fatty ketones. Popoff's rules (*supra*) apply to the oxidation of the benzene ketones: the carbonyl group always remains attached to a benzene nucleus to which it is directly united. Like the fatty ketones, the benzene ketones react with hydroxylamine and with phenyl-hydrazine eliminating water and yielding hydroximes and hydrazones. On reduction with sodium amalgam they form secondary alcohols, but heating with hydriodic acid, or distillation with zinc dust, reduces them to the corresponding hydrocarbons



By fusion with caustic alkalis they yield a salt of a benzene acid together with a hydrocarbon



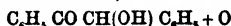
They are less subject than the fatty ketones to undergo condensation with elimination of water, and those which contain two benzene radicles directly attached to the carbonyl group do not exhibit this tendency at all

Only those benzene ketones combine with hydrogen sodium sulphite in which the carbonyl is attached to two fatty groups, one of which is methyl: thus, benzyl methyl ketone $\text{C}_6\text{H}_5, \text{CH}_2, \text{CO CH}_3$, and methyl phenylethyl ketone $\text{C}_6\text{H}_5, \text{CH}_2, \text{CH}_2, \text{CO CH}_3$, form double compounds, acetophenone $\text{C}_6\text{H}_5, \text{CO CH}_3$, does not.

Diketones The diketones contain two carbonyl groups attached to hydrocarbon radicles. If the two carbonyl groups are directly united, the compound is an α -diketone, if they are united by a carbon atom, it is a β -diketone, if by a chain of two carbon atoms, a γ -ketone, and so on.

Some of the diketones, such as benzil $\text{C}_6\text{H}_5, \text{CO CO C}_6\text{H}_5$, have been known for a considerable time, but the majority are of recent discovery. Their chief interest lies in the ease with which the two carbonyl groups may be made to condense with one or more molecules of some other substance to form closed chain compounds. In this way glyoxalines, quinolines, pyrazoles, furfurans, pyrroles, thiophenes, and similar compounds have been prepared.

α -Diketones These may be regarded as homologues of the dialdehyde glyoxal, CHO CHO . They may be obtained by the action of chlorine, or, better, of nitric acid, on a ketone alcohols



Benzoin

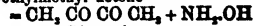


Benzil

(Laurent, *A Ch*. [2] 59, 402, Zinin, *A* 34, 188) By hydrolysing the isonitrosoketones by boiling them with dilute sulphuric acid



Isonitroso-ethylmethyl ketone

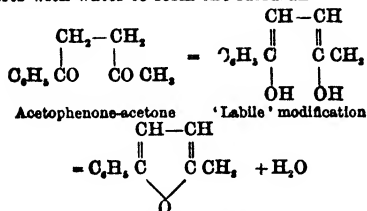


Diacetyl

(Von Pechmann, *B*. 21, 1411)

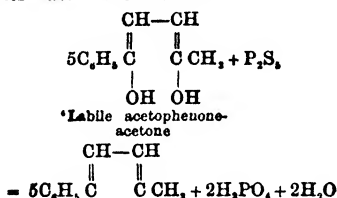
The benzenoid α -diketones react with caustic potash to form secondary glycolic acids, the two

tride, the γ -diketones are converted into furfurans. It is supposed that in this reaction the diketone first forms a 'labile' modification containing two hydroxyl groups, and that this then parts with water to form the furfuran



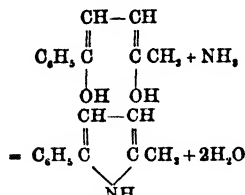
Phenylmethyl furfuran

(Paal, B 17, 2756) An unsaturated ketone isomeric with phenylmethylfurfuran is formed at the same time—*Phosphorus pentasulphide* converts γ diketones into thiophenes, whilst with ammonia they yield pyrroles, the reaction in both cases being supposed to be preceded by the above-mentioned transformation of the diketone into its 'labile' modification



Phenylmethylthiophen

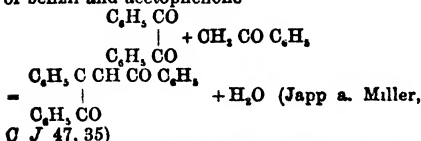
and



Phenylmethylpyrrole

(Paal, B 18, 367)

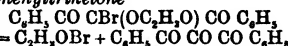
An unsaturated γ diketone, *anhydrazetophenonebenz* ($\alpha\beta$ dibenzoylstyrolene) is formed by the action of caustic potash on a mixture of benzil and acetophenone



Quinones If the two carbonyl groups of a diketone occur in a closed chain of six carbon atoms, the resulting compound belongs to the class of the quinones. This group of diketones will be treated of separately (*v* Quinones).

Triketones A triketone in which the three carbonyl groups are directly united has been obtained in the following manner (von Pechmann, B. 22, 862). Dibenzoylbromomethane

$\text{C}_6\text{H}_5\text{COCHBrCO C}_6\text{H}_5$ (prepared by the action of bromine on dibenzoylmethane, *v supra*) is converted by treatment with potassium acetate into the acetate of dibenzoylcarbinol $\text{C}_6\text{H}_5\text{COCH(OC}_2\text{H}_5\text{O)CO C}_6\text{H}_5$, which, when brominated, yields $\text{C}_6\text{H}_5\text{COCHBr(OC}_2\text{H}_5\text{O)CO C}_6\text{H}_5$. When this compound is heated above its melting-point (101.5°) it breaks up into acetyl bromide and *diphenyltriketone*



Diphenyltriketone, like some other compounds in which a carbonyl-group is situated between two electro negative groups, unites with 1 mol of water to form a hydrate, this has probably the constitution $\text{C}_6\text{H}_5\text{CO C(OH)(CO C}_6\text{H}_5\text{)CO C}_6\text{H}_5$.

The formation of another triketone, *tribenzoylmethane*, in which the three carbonyl groups are attached to the same carbon atom, has already been referred to F R J

DIKETO-PENTANE v METHYL ETHYL DIKETONE

$\alpha\beta$ DIKETO-PROPYL-BENZENE v PHENYL-METHYL DIKETONE

KINIC ACID Synonym of QUINIC ACID

KINO The dried juice, obtained from incisions in the trunk of *Pterocarpus marsupium* (Mala bar) The name has also been given to the dried juice from other trees, e.g. *Eucalyptus resinifera* (Australia) *Pterocarpus ernaceus*, and *Coccoloba uvifera* (America) Kino greatly resembles catechu, and is used in dyeing and in medicine. It occurs in small brittle opaque reddish black fragments, partly soluble in water, almost entirely soluble in alcohol, insoluble in ether. It has no odour, but a very astringent taste. Used in medicine as an astringent. It gives off pyrocatechin when distilled (Eissfeld, A 92, 101). Potash fusion forms phloroglucin and protocatechuic acid (Hlasiwetz, A 134, 122, Stenhouse, A 177, 187). Kino contains kinoin and kino red and other bodies. Kinoin has been described as a 'tannin' (Bergholz, cf Etti, B 17, 2241).

Kinoin $\text{C}_{11}\text{H}_{12}\text{O}_6$. Occurs together with kino-red, &c., in gum kino, from which it is extracted by first boiling with dilute HCl, and then extracting the solution with ether, yield 1½ p.c. (Etti, B 11, 1879). Colourless prisms. Sol cold, readily sol hot, water. On heating to 120° it gives kino red ($\text{C}_{12}\text{H}_{12}\text{O}_{11}$). On heating with HCl to 130° it is decomposed into methyl chloride, pyrocatechin, and gallic acid.

Kino red $\text{C}_{22}\text{H}_{20}\text{O}_{11}$ [160°-170°] A red resin occurring in kino and left undissolved when it is boiled with dilute HCl. It is also formed by heating kinoin. It is sol water, v sol alcohol. FeCl₃ gives a dirty green colouration. It dissolves in alkalis. It ppts a solution of gelatin. On fusion it yields a red amorphous anhydride $\text{C}_{22}\text{H}_{20}\text{O}_{10}$, which also ppts a solution of gelatin. The anhydride is also formed by heating kino-red with dilute HCl aq. Kino-red yields phenol and pyrocatechin when distilled.

KOSIN $\text{C}_{12}\text{H}_{12}\text{O}_8$ [142°] Obtained from cussor or koussor, the dried flowers of the Abyssinian koso tree (*Hagenia Abyssinica*), used by the inhabitants as a remedy again it tape worm (Flückiger a Burri, Ar Ph [3] 5, 193, Ph [3] 5, 562). The anthelmintic properties of koso seem to be due to kosin. Sulphur yellow trimetric crystals. Nearly insol. water, v sol

alcohol, ether, benzene, CS_2 , chloroform, and HOAc . Dissolves in aqueous alkalis and Na_2CO_3 , and is reprecipitated by acids. FeCl_3 colours its alcoholic solution permanently red. Conc. H_2SO_4 does not decompose it at 15° , but on warming it forms isobutyric acid and a red amorphous body $\text{C}_8\text{H}_8\text{O}_4$ (?). Potash fusion gives formic, butyric, and oxalic acids. Sodium amalgam yields a volatile oil $\text{C}_8\text{H}_8\text{O}_2$ and an amorphous yellow substance $(\text{C}_8\text{H}_8\text{O}_2)_x$.

Acetyl derivative $\text{C}_{11}\text{H}_{12}\text{Ac}_2\text{O}_4$. From kosin and Ac_2O

KOUMISS v MILK

KYANETHINE v CYANETHINE

KYANPHENINE v CYAPHENINE

KYANPROPINE v CYANPROPINE

KYNURENIC ACID v OXY QUINOLINE CAR

BOXYLIC ACID

KYNURIC ACID v CYNURIC ACID

KYNURIN v OXY QUINOLINE

L

LABURNINE An alkaloid said to occur along with cytosine in laburnum seeds (Husemann & Marmé, *Z* [2] 1, 161, 5, 677). Light crystalline groups of monoclinic prisms, v sol water, sl sol alcohol, nearly insol ether.

LAC A substance surrounding the eggs of *Coccus lacca* on the twigs of various trees in Bengal and Sumatra. These twigs are called stick lac. The lac is removed by melting and squeezing through canvas, when the hardened drops are known as seed lac. After bleaching by chlorine or charcoal and making into sticks it is known as shell lac or shellac. Lac is used for making sealing wax, varnishes, and a red dye called lac dye. Lac contains 68 to 88 p.c. resin (about half of which is soluble in ether), a small quantity of wax and of a red dye, and a larger quantity of a substance insoluble in alcohol. The wax [60°] contains ceryl and myricyl alcohols and their stearic, palmitic, and oleic ethers (Benedikt & Ulzer, *M* 9, 579). Shellac may be freed from wax by dissolving in boiling dilute Na_2CO_3 , filtering, and ppg with HCl . If the boiling be continued too long, liquid shellac is produced. Shellac freed from wax gives azelaic acid when oxidised by KMnO_4 (Benedikt & Ehrlich, *M* 9, 163).

Liquid shellac $\text{C}_{16}\text{H}_{12}\text{O}_4$. Formed by boiling shellac (1,000 g) with NaOH (300 g) and water (20,000 g) for 2 hours. When cold, sulphuric acid is added and the liquid shaken with ether (B & E). Thick liquid, insol water, v sol alcohol and ether. Forms salts with Pb , Ag , Zn , Ba , and Ca — $\text{C}_{16}\text{H}_{12}\text{O}_4\text{Mg}_2\text{O}_2$, amorphous, v sol water, ppd on heating, but redissolves on cooling.

Laccalic acid $\text{C}_{16}\text{H}_{12}\text{O}_4$ (?). Obtained from lac dye by treating with HClAq , filtering, boiling the residue with water, ppg the solution with lead acetate, and decomposing the pp with H_2S (R E Schmidt, *B* 20, 1285). Brownish red powder composed of minute tables (from alcohol ether), m sol alcohol, sl sol water, nearly insol ether, insol benzene. Decomposes, without previous fusion, at about 180° . Alkalis form a crimson solution which has an absorption spectrum like that of carminic acid. Baryta gives a violet pp. FeSO_4 a black pp. FeCl_3 a black colouration only. It reduces ammoniacal AgNO_3 but not Fehling's solution. Conc. HNO_3 yields picric and oxalic acids. Conc. HClAq at 180° forms a substance $\text{C}_{16}\text{H}_{12}\text{O}_{11}$. Laccalic acid dyes like carminic acid, but its solution in H_2SO_4 shows a different absorption spectrum— $\text{K}_2\text{C}_{16}\text{H}_{12}\text{O}_4$ (dried

at 100°) flocculent pp formed by adding alcoholic potash to an alcoholic solution of the salt $\text{BaC}_{16}\text{H}_{12}\text{O}_4$, brownish red pp formed by adding NH_3 and BaCl_2 .

LACMOID

Preparation—1 A mixture of resorcinol (20 pts), sodium nitrite (1 pt), and water (1 pt) is slowly heated to 110° , at which temperature a vigorous reaction sets in, the melt is then kept at 115° – 120° till it has become quite blue and the evolution of NH_3 has ceased (Traub & Hook, *B* 17, 2615)—2 By the action of H_2O_2 on ammoniacal resorcin solution (Wurster, *B* 20, 2938)—3 By fusing resorcin (15 g) with KNO_3 (18 g) (Benedikt & Juhus, *M* 5, 534).

Properties—Glistening brown powder. It is easily soluble in methyl, ethyl, and amyl alcohols, acetone, acetic acid, and phenol, less easily in ether and water, insoluble in benzene and petroleum spirit. It dissolves in strong HCl or H_2SO_4 with a blue colour. It decomposes below 200° .

Lacmoid differs from litmin, the colouring matter of litmus, in being soluble in strong alcohol. Litmin is insoluble in strong alcohol, though soluble in dilute (50 p.c.) spirit. The solution of litmin in 50 p.c. alcohol is bleached after a time, while that of lacmoid retains its colour for months. The absorption spectra of the two substances do not differ markedly (Hartley, *R. Dublin Soc* 5, 159).

LACTAM The anhydride of an amide acid $\text{R}^-(\text{NH}_2)(\text{CO}_2\text{H})$ containing the group $\text{R}^-\begin{matrix} \text{CO} \\ \diagup \\ \text{NH} \end{matrix}$

(Baeyer, *B* 15, 2102)

LACTAMIC ACID is a AMIDO PROPIONIC ACID

Di lactamic acid is a IMIDO DI PROPIONIC ACID

LACTAMIDE v Amide of LACTIC ACID

LACTARIC ACID $\text{C}_4\text{H}_6\text{O}_4$. Prepared by pressing the juice out of the mushroom (*Lactarius piperatus*) and extracting the residue with ether or boiling alcohol (Chuit, *Bl* [3] 2, 158). White leaflets (from alcohol), v sol hot alcohol, ether, CHCl_3 , CS_2 , and benzene, v sl sol cold alcohol and petroleum spirit, insol water.

Salts—**Naa'** greasy scales, decomposing at 250° —**Ka'** silvery scales, decomposing at 245° without melting—**KHa'**, silky scales. [115°]—**NH₄A'** [92°], white leaflets, decomposed by boiling water—**CaA'**, scales. When distilled it forms the ketone $(\text{C}_4\text{H}_6\text{O})_2\text{CO}$ [82°]—**BaA'**, white amorphous pp. Insol water, alcohol, and ether—**PbHa'**, [114°]. Thin transparent leaflets.

Methyl ether MeA' [J8°] White scales
Sol. ligroin and ether

Ethyl ether EtA' [36°] White scales
V sol warm alcohol, ether, chloroform, and CS₂
Amide C₁₁H₁₅O NH₂ [108°] Trimetric needles

LACTIC ACID C₃H₅O₃ = CH₃.CH(OH).CO₂H
α-Oxy propionic acid Ethyldene lactic acid
Fermentation lactic acid Mol w 90 SG ³⁹
1.2403 (Brühl), ⁴ 1.2485 (Mendeleeff, C R 50,
52) $\mu_D = 1.4469$ $R_{20} = 31.18$

Isomerides—Lactic acid is isomeric with hydracrylic acid, and appears to be chemically identical with sarcosolactic acid (*v infra*)

In 1863 Wislicenus got from ethylene chlorhydrin and KCN, by saponifying the product, a lactic acid which formed a crystalline zinc salt ZnA', 2aq. He called it ethylene lactic acid, thought it was identical with Liebig's sarcosolactic acid (A 128, 4) Dossios (A 146, 168) said that sarcosolactic acid gave, on oxidation, malonic acid. Wislicenus held (A 167, 346) that his ethylene lactic acid was not hydracrylic acid, for by the action of HI he was unable to obtain β iodopropionic acid. He said also that his ethylene lactic acid differed from sarcosolactic, although it accompanied sarcosolactic acid in extract of meat. Erlenmeyer (A 191, 261) showed that Wislicenus's ethylene lactic acid from glycol chlorhydrin is hydracrylic acid, and could be converted by HI into β iodopropionic acid. The acid accompanying sarcosolactic acid is, according to Siegfried (B 22, 2711) the acetyl derivative of lactic acid.

Occurrence—1 Discovered by Scheele in sour milk and first recognised as a peculiar acid by Berzelius—2 In opium (Smith, Ph [2] 7, 50, Buchanan, B 3, 182)

Formation—1 by the lactic fermentation of milk sugar, cane sugar, or glucose (*v* FERMENTATION, vol II p 543)—2 By the oxidation of propylene glycol by the oxygen of the air in presence of platinum black (Wurtz, A 105, 206, 107, 192)—3 By boiling a chloro propionic acid with water and Ag₂O (Wurtz, A Ch [3] 59, 165, Buff, A 140, 156) In like manner from a bromo propionic acid (Friedel & Maohuca, A 120, 286)—4 By the action of nitrous acid upon alanine (Strecker, A 75, 27, 42)—5 By the reduction of pyruvic acid by sodium amalgam (Wislicenus, A 126, 227) or with zinc in presence of dilute acetic acid (Debus, C J 16, 260)—6 From aldehyde by combination with HCN & a d saponification of the resulting nitrile (Wislicenus, A 128, 6, Simpson & Gauther, A 146, 264)—7 By heating dichloroacetone with a large quantity of water at 200° (Linnemann & Zotta, A 159, 247)—8 By boiling glucose with aqueous NaOH (Hoppe Seyler, B 4, 346)—9 When glucose (20 g) is allowed to stand with potash (40 g) and water (200 cc) at 40° in a flask loosely plugged with cotton wool, it is converted within 24 hours into lactic acid (8 g) and another acid, soluble in alcohol (Nencki & Sieber, J pr 183, 499) Lactic acid may possibly be thus formed in animals from sugar. The decomposition occurs even in very dilute solutions, thus even 9 g of glucose and 9 g of potash dissolved in 8 litres of water and kept at 35°–40° are converted in 10 days into potassium lactate Na₂CO₃ does not decompose

sugar at 40°, nor does NH₃, but NaOH, NMe₃OH and neurine do convert it into lactic acid. Creatinine and guanidine produce no lactic acid. Milk sugar and maltose behave like glucose. Lactic acid is formed when glucose (dextrose) (1 pt), water (10 pts) and KOH (2 pts) are left, even in an atmosphere of hydrogen, for 48 hours at 35°. No sugar is left (Nencki & Sieber, J pr [2] 26, 1, Kiliani, B 15, 701)—10 By heating cane sugar with baryta at 150° (Schutzenberger, B [2] 25, 289)—11 Among the products of the distillation of glycerin with KOH (Herter, B 11, 1167), and formed, together with formic and oxalic acids, by boiling glycerin with potash solution (Debus, A 109, 229)

Preparation—1 The filtered solution of sour whey evaporated nearly to dryness, in strong alcohol, is mixed with alcoholic tartaric acid, as long as any precipitate of tartrate of potassium, sodium, and calcium is formed, the liquid decanted after 24 hours, and evaporated, the residue dissolved in water, the solution digested with carbonate of lead, till lead dissolves in it, the filtrate evaporated, neutralised with carbonate of barium, again filtered, and diluted with water, the whole of the barium precipitated with sulphate of zinc, and the filtrate evaporated till lactate of zinc crystallises out (Berzelius, *Lehrb* Aug 5, v 241)—2 Three kilos of cane sugar and 15 g of tartaric acid (which serves to invert the cane sugar) are dissolved in 13 kilos of boiling water, 1½ kilos of levigated chalk added after two days, together with 60 g of stinking cheese, suspended in 4 kilos of sour milk (decaying cheese favours the production of lactic acid and retards its conversion into butyric acid), the mixture set aside at a temperature between 30° and 35°, and well stirred every day till, in the course of six or eight days, it is converted into a stiff paste of lactate of calcium, this paste is boiled for an hour with 15 g of quicklime and 10 kilos of water, the solution strained through a cloth filter and evaporated to a syrup, the crystalline mass, which forms in four days, pressed, first by itself, then three or four times, after having been each time stirred up, with $\frac{1}{2}$ of its weight of cold water, and the lactate of calcium thus purified is dissolved in twice its weight of boiling water. To the solution of every 32 pts of the calcium salt there is then added a mixture of 7 pts oil of vitriol and 7 pts water, the lactic acid, whilst still hot, is strained through linen to separate it from sulphate of lime, the filtrate obtained from 7 pts of oil of vitriol is boiled with 1½ pts carbonate of zinc for a quarter of an hour (by longer boiling a very sparingly soluble basic salt is formed), the liquid is filtered boiling hot, the colourless crystalline grains of lactate of zinc, which separate on cooling, are freed from sulphuric acid by washing with cold water, and additional quantities of crystalline grains are obtained by evaporating the mother-liquor almost to the end. Lastly, 1 pt of the zinc salt is dissolved in 7½ pts of boiling water, sulphuretted hydrogen passed through the solution as long as sulphide of zinc is precipitated, and the filtrate boiled and evaporated on the water bath to a syrup, whereupon 8 pts. of the zinc salt yield 5 pts. of syrupy lactic

acid (Bensch, *A* 61, 174) By this process, 100 pts of cane sugar yield 117 pts of lactate of calcium, which, if the sugar was white, is colourless, and does not require to be purified by pressure (*cf* Engelhardt a Maddrell, *A* 63, 83, 70, 241, Boutron a Frey, *J Ph* 27, 841) 8 Lautemann (*A* 113, 242) recommends the following modification of Bensch's process of preparation—Retaining the proportions of sugar, tartaric acid, milk, and cheese indicated by the latter, he takes one third more water, uses 1,200 g oxide of zinc (commercial zinc white) instead of levigated chalk, and keeps the temperature as constantly as possible between 40° and 45° during the fermentation. After eight or ten days, the inside of the vessel is lined with white crystals of lactate of zinc, which can be obtained pure by one or two crystallisations from boiling water. The lactic acid prepared from the zinc salt generally contains mannite, which does not completely crystallise out from the concentrated acid. To separate this, the aqueous acid is shaken up with ether, and then the ethereal layer is pipetted off and evaporated; it then leaves pure lactic acid—4 The following mixture is recommended by C O Harz (*Vierteljahrsschrift pr Pharm* 20, 501), 3 pts milk sugar, 36 pts ordinary water, 0.5 to 0.75 pt flour containing a large proportion of gluten, a little beer yeast, 6 pts of soda crystals, or 3 pts of sodium bicarbonate. When the fermentation has once been set up by milk sugar, it may be continued by addition of cane sugar—5 By the action of NaOH or KOH on dextrose or levulose, the operation being performed as follows—A solution of 500 grms of cane sugar in 250 cc of water and 10 cc dilute sulphuric acid (3 pts H_2SO_4 to 4 pts H_2O) is heated to 50° in a closed vessel for 3 hrs. After cooling, 400 cc of aqueous NaOH (50 p.c.) is slowly added. The mixture is then warmed to 60°–70° till it no longer reduces Fehling's solution, the calculated quantity of H_2SO_4 (same strength as before) is then added, and the Na_2SO_4 made to crystallise out by cooling and agitation. The mass is extracted with 93 p.c. spirit and filtered, the filtrate is divided into two portions, one half being neutralised with $ZnCO_3$, filtered hot and the other half added. On cooling, the zinc lactate crystallises out in a nearly pure state, the yield is 200 grms (Kihani, *B* 15, 699, *cf* Hoppe Seyler, *B* 4, 346)

Properties—Colourless syrup with very sour taste. Hygroscopic. Miscible with water and alcohol, sl. sol. ether. Does not solidify at –24°

Reactions—1 When gradually heated it gives off water at 130° leaving solid lactide. At 250° to 300° the products are CO , CO_2 , aldehyde, and lactide—2 Dilute H_2SO_4 at 130° gives aldehyde and formic acid (Erlenmeyer, *Z* 1868, 348) $CH_3CH(OH)CO_2H = CH_3CHO + HCO_2H$ 8 When gently heated with conc H_2SO_4 it gives off CO freely (Pelouze, *A Ch* [3] 13, 257)—4 Boiling nitric acid forms oxalic acid—5 Distillation with $NaCl$, dilute H_2SO_4 , and MnO_2 yields aldehyde and chloral (Städeler, *A* 69, 392)—6 Distillation with MnO_2 and dilute H_2SO_4 yields aldehyde and CO_2 —7 Fuming H_2SO_4 gives methane disulphonic acid (Strecker, *A* 118, 291)—8 Chromic acid mixture gives

acetic acid and CO_2 (Chapman a Smith, *Z* 1867, 477)—9 $KMnO_4$ gives pyruvic acid—10 $POCl_3$ acting on dry calcium lactate gives chloro-propionyl chloride $CH_3CHClCOCl$ whence water forms a chloro propionic acid 11 HIAg reduces it to propionic acid (Lautemann, *A* 113, 217)—12 A concentrated solution of potassium lactate submitted to electrolysis yields aldehyde and CO_2 (Kolbe, *A* 113, 244, Brestre, *Z* 1866, 680)—13 Distillation with quicklime yields alcohol C_2H_5O , $= CO_2 + C_2H_5O$ (Hanriot, *C R* 101, 1156, *Bl* [2] 45, 80)—14 Heated at 170° in a stream of gaseous HBr , or at 100° in sealed tubes with conc $HBrAg$, it yields a bromo propionic acid (Kekulé, *A* 130, 11)—15 Bromine at 100° decomposes lactic acid (Beilstein, *A* 120, 227) Br acting on an ethereal solution of lactic acid forms tribromo pyruvic ether (Wichelhaus, *A* 143, 10, Khimenko, *J R* 8, 125)—16 The dry distillation of calcium lactate produces CO_2 , ethylene, propylene (Gossin, *Bl* [2] 43, 49), acrylic acid, phenol (?) (Claus, *A* 136, 287), and other products—17 Distillation of calcium lactate with soda lime yields acetic, propionic, butyric, hexoic, and other fatty acids. Heating calcium lactate with KOH at 280° yields formic, acetic, propionic, butyric, and oxalic acids (Hoppe Seyler, *H* 3, 352)—18 Among the products of the nutritive fermentation of calcium lactate are hydrogen, CO_2 , acetic, propionic, butyric, and n valeric acids and ethyl alcohol (Pasteur, *Bl* 1862, 52, Strecker, *A* 92, 80, Fitz, *B* 11, 1898, 12, 479, 13, 1309)—19 Heating with m amido benzoic acid forms $CH_3CH(OH)CONH C_6H_4CO_2H$ crystallising from water in small prisms [162°] (Pellizzari, *A* 232, 154), which at 240 forms an anhydride CH_3CH

$\begin{array}{c} | \\ >N C_6H_4 CO_2H \\ CO \end{array}$ [243°] The corre

sponding acetyl derivative

$CH_3CH(OAc)CONH C_6H_4CO_2H$ melts at 148°

Estimation—The substance, acidified with H_2SO_4 , is extracted with ether, the ethereal solution evaporated, the residue treated with water and the aqueous solution ppt with lead acetate and filtered. The filtrate is then ppt with alcoholic NH_3 , and the pp of $PbO_2 \cdot 2(C_2H_5O_2)$ washed with alcohol, dried, weighed, ignited, and weighed again (Palm, *Fr* 26, 33)

Salts—The crystalline lactates do not effloresce in the air, but give off water *in vacuo*. They are not decomposed at 150°. They are insoluble in ether and, for the most part, sparingly soluble in cold water—Ammonium salt forms deliquescent prisms, and gives off NH_3 when exposed to air— NaA' (at 140°) amorphous, deliquescent mass, v sol water and alcohol, ppt by ether from its alcoholic solution— $Na_2C_2H_3O_2 \cdot 2e$ $CH_3CH(ONa)CO_2Na$ obtained by the action of sodium on the preceding at 130°. Hard, deliquescent brittle mass. Decomposed by cold water, with evolution of heat, into $NaOH$ and $NaC_2H_3O_2$. Absorbs CO_2 from the air, forming Na_2CO_3 and sodic lactate. Basic sodium lactate appears to dissolve without decomposition in perfectly dry alcohol. With MeI it gives NaI and $CH_3CH(OMe)CO_2Na$ (Wischneus, *A* 125, 49)—Potassium salt crystallises with difficulty— BaA' , $4aq$ (at 100°):

large cauliflower-like bundles of rectangular needles (Hans Meyer, *B* 19, 2454) Deliquescent, v sol dilute alcohol, insol absolute alcohol and ether— $\text{BaH}_2\text{A}'$, crystals, v sol water Not altered by exposure to air— CaA' , 5aq small white mammillated crystals (from water or alcohol) *S* 10.5 in the cold Extremely soluble in boiling water Sol hot, nearly insol cold, alcohol At 100° it becomes CaA'_2 , and at 280° it is converted into $\text{CaC}_2\text{H}_3\text{O}_6$ (Wurtz a Friedel, *A Ch* [3] 63, 134)— $\text{CaH}_2\text{A}'$, 3aq crystals resembling wavelite, sol absolute alcohol (Engelhardt a Maddrell, *A* 63, 119)— $\text{CaK}_2\text{A}'$, octahedra Deposits CaA' when dissolved in warm water (Strecker, *A* 91, 352)— $\text{CaNa}_2\text{A}'$, 2aq granules (*S*) CaCl_2 3aq prisms, v e sol water (*E a M*)— $\text{Ca}_2(\text{CHO})_2(\text{C}_2\text{H}_3\text{O}_6)\text{Cl}$, 10aq from calcium chloride, formate, and lactate (Böttger, *A* 188, 329) Long needles— SrA' , 3aq very soluble— MgA' , 3aq prisms, insol alcohol, less soluble than the corresponding sarcosylactate *S* 3.6 in the cold, 16.7 at 100° — $\text{Al}_2\text{A}'$, triclinic octahedra (Hans Meyer, *B* 19, 2454)— $\text{AlNa}_2\text{A}'$, 5aq rectangular prisms and tables— FeA' , 3aq small greenish crystals *S* 2.1 at 10° , 8.5 at 100° Insol alcohol—Ferric lactate is a brown amorphous deliquescent mass, v sol water— MnA' , 3aq amethyst coloured monoclinic crystals, m sol cold, v sol hot, water— CoA' , 3aq peach blossom coloured needles, nearly insol cold, m sol boiling water, insol alcohol— NiA' , 3aq apple green needles— ZnA' , 3aq shining crusts, or large crystals irregularly grouped *S* 1.07 at 8° (Buff *A* 140, 160), 1.8 at 10° (Wislicenus, *A* 126, 228), 1.9 at 15° (Strecker *A* 105, 316), 16.7 at 100° Almost insol alcohol, which partly converts it into amorphous ZnA' , 3aq, which again takes up 2aq when exposed to the air (Khimenko, *J R* 12, 98)— ZnA N_2H_4 — ZnA' , N_2H_4 (Lutschak, *B* 5, 30)— $\text{ZnNa}_2\text{A}'$, 2aq— CdA' , small needles Anhydrous when deposited from a boiling solution Insol alcohol *S* 10 in the cold, 12.5 at 100° — $\text{BiC}_2\text{H}_3\text{O}_6$ (Brüning, *A* 104, 194)— CuA' , 2aq dark blue monoclinic tables (Schabus *J* 1854, 40c) *S* 16.7 in the cold, 45 at 100° *S* (alcohol) 9 in the cold, 4 at 78° Decomposed at 200° giving aldehyde, lactide, and CO_2 (Engelhardt, *A* 70, 249)— $\text{CuC}_2\text{H}_3\text{O}_6$, v sl sol water— $\text{Hg}_2\text{A}'$, 3aq rose-coloured or crimson crystals, sl sol water, obtained by mixing the boiling solutions of mercurous nitrate and sodium lactate (*E a M*)—Prisms of the mercurous salt $\text{Hg}_2\text{A}'_2$ are also formed by boiling aqueous lactic acid with HgO (Brunner) — $\text{SnC}_2\text{H}_3\text{O}_6$, crystalline powder, insol water— PbA' , gummy, v sol water— $\text{PbC}_2\text{H}_3\text{O}_6$ (Moldenhauer, *A* 131, 333)— $\text{PbA}'_2\text{O}_2$, 3aq heavy granular pp, formed when lactic acid is mixed with lead acetate and alcoholic NH_3 — UroA' yellow crystalline crusts— AgA' 3aq silky needles *S* 5 in the cold *V* sol hot, nearly insol cold, alcohol

Nitrosyl derivative

$\text{CH}_3\text{CH}(\text{ONO})\text{CO}_2\text{H}$ *S G* 135 Formed by dissolving lactic acid in a mixture of conc HNO_3 and H_2SO_4 , and ppg with water (Henry, *B* 3, 532) Thick oil Sl sol water v e sol ether Decomposes in the cold into HCO_2H and oxalic acid (Henry, *B* 12, 1887)

Acetyl derivative $\text{C}_4\text{H}_7\text{O}_5$, *ie*

$\text{CH}_3\text{CH}(\text{OAc})\text{CO}_2\text{H}$ [167°] (Siegfried) Formed by heating ethyl lactate with AcCl and saponifying the product by heating it with water at 150° for 8 hours (Perkin, *Z* 1861, 166, Wislicenus, *A* 125, 60) Formed also by boiling a solution of zinc acetate and sarcosylactate or lactate (Siegfried, *B* 22, 2715) Occurs in extract of meat. Thin needles, v e sol water Inactive Volatile with steam Decomposed by long boiling with water into acetic and lactic acids Bases quickly effect this decomposition When first prepared it is soluble in alcohol, but on keeping it becomes insoluble in alcohol, does not then melt below 300° , and is saponified with great difficulty by alkalis HI produces no β iodopropionic acid— BaA'_2 , 4aq brittle gumlike mass, v sol water, sol alcohol—Copper salt amorphous deliquescent bluish green mass— ZnA' , gummy mass Its solutions become quickly acid, from conversion into acetic acid and zinc lactate

Benzoyl derivative $\text{C}_{10}\text{H}_{11}\text{O}_5$, *ie*

$\text{CH}_3\text{CH}(\text{OBz})\text{CO}_2\text{H}$ [112°] *S* 25 in the cold Obtained by heating lactic acid with benzoic acid at 180° (Strecker, *A* 80, 42, 91, 360) Formed also by the action of BzCl on calcium lactate (Wislicenus, *A* 183, 277) Tables or needles, m sol boiling water, v sol alcohol and ether Converted by boiling water into benzoic and lactic acids When recrystallised from water benzoyl lactic acid is always accompanied by an oily hydrate $\text{C}_{10}\text{H}_{11}\text{O}_5$, 3aq, which in dry air is slowly converted into the crystalline acids— BaA'_2 , 6aq thin six sided plates— AgA'

Amide $\text{C}_4\text{H}_7\text{NO}_5$, *ie* $\text{CH}_3\text{CH}(\text{OH})\text{CONH}_2$, [74°] Obtained by the action of gaseous or alcoholic NH_3 on lactide (Wurtz a Friedel, *A Ch* [3] 63, 108) Formed also by saturating ethyl lactate with ammonia and allowing the liquid to stand (Brunner, *A* 104, 197), and by heating ammonium lactate in a slow current of dry NH_3 at 130° (Engel, *C R* 98, 574) Small prisms, v sol water and alcohol Does not combine with acids or bases, but is decomposed by them on boiling into NH_3 and lactic acid

Benzoyl derivative of the amide $\text{CH}_3\text{CH}(\text{OBz})\text{CONH}_2$, [124°] From benzoyl lactic ether and alcoholic NH_3 (Wislicenus, *A* 133, 257) White needles, may be sublimed, sl sol water, v sol alcohol Resolved by boiling KOH into lactic and benzoic acids and NH_3 Prolonged treatment with alcoholic NH_3 forms benzamide and lactamide

Isomeric of the amide $\text{C}_4\text{H}_7\text{NO}_5$, Formed by heating ammonium lactate in a current of dry NH_3 at 100° (Engel, *C R* 98, 574) Pale amber coloured syrup Decomposes at 200° In contact with water it immediately forms ammonium lactate

Ethylamide $\text{CH}_3\text{CH}(\text{OH})\text{CONHET}$ [48°] (260°) From lactide and NH_3Et Crystalline (Wurtz a Friedel, *A Ch* [3] 63, 610) Decomposed by alkalis into ethylamine and lactic acid

Anilide $\text{C}_9\text{H}_9\text{NO}_5$, *ie*

$\text{CH}_3\text{CH}(\text{OH})\text{CONHPh}$ [58°] Formed by heating lactic ether with aniline at 150° (Leipen, *M* 9, 48) Colourless prisms, sl sol water, insol ligroin, v sol ether, chloroform, and alcohol

o-Toluide $C_6H_4NO_2$ [72°] From *o*-toluidine and lactic ether Crystalline powder (from benzene) Insol lignroin (Leipen, *M.* 9, 50, 51)

p-Toluide $C_6H_4NO_2$ \cdot $\frac{1}{2}$ $CH_3CH(OH)CO_2NHCH_3$ [102°] From *p*-toluidine and ethyl-lactate White needles $\frac{1}{2}$ sol water

Cyanamide $CN NH(CO CH(OH) CH_3)$
Lacto-cyanamide Formed by dissolving lactide (20 g) in a strong alcoholic solution of potassium cyanide (16 g), passing in CO_2 to remove free alkali and evaporating to crystallisation (Mertens, *J pr* 125, 33) It forms a silver derivative

Methyl ether $C_4H_8O_4$ \cdot $\frac{1}{2}$ $CH_3CH(OH)CO_2Me$ (145° \cdot V) SG $\frac{1}{2}$ 1180 (Schreiner, *A* 197, 1, B 12, 179) Colourless neutral liquid Combines with $CaCl_2$. Is immediately decomposed by water

Ethyl ether $C_4H_8O_4$ \cdot $\frac{1}{2}$ $CH_3CH(OH)CO_2Et$ **Ethyl lactate** (154° \cdot V) SG $\frac{1}{2}$ 10546 Formed by distilling calcium lactate with $KEtSO_4$ (Strecker, *A* 81, 247, 91, 355) Formed also by heating lactic acid (dried at 145°) with alcohol (F & W) and by passing alcohol vapour into lactic acid at 175° (Wislicenus, *A* 125, 58) Colourless liquid, immediately decomposed by water (Schreiner) Forms with $CaCl_2$ the compound $CaCl_2 \cdot 4EtA'$ crystallising in granules Chloral gives a liquid compound, whence phosphorus pentachloride forms liquid $CCl_3CHClOCHMeCO_2Et$ SG $\frac{1}{2}$ 142 (Henry, *Bull Acad Roy Belg* [2] 37, No 5) Lactic ether is a weak hypnotic (Pallacan 5) Berton, *C C* 1887, 1149)

Acetyl derivative of the ethyl ether $CH_3CH(OAc)CO_2Et$ (177°) at 733 mm VD 570 (calc 554) SG $\frac{1}{2}$ 1046 From ethyl lactate and $AcCl$ (Wislicenus, *A* 125, 58) Neutral oil, gradually decomposed by water into alcohol and $CH_3CH(OAc)CO_2H$ Miscible with alcohol and ether

Benzoyl derivative of the ethyl ether $CH_3CH(OBz)CO_2Et$ (288° cor) From ethyl lactate and $BzCl$ at 100°, or from silver benzoyl-lactate and EtI (Wislicenus, *A* 133, 272) Oil, miscible with alcohol and ether Resolved by water at 150° into lactic acid and benzoic ether Alcohol NH_3 gives $CH_3CH(OBz)CONH_2$

Nitroxyl derivative of the ethyl ether $CH_3CH(ONO_2)CO_2Et$ (178°) SG $\frac{1}{2}$ 1153 From ethyl lactate, HNO_3 , and H_2SO_4 (Henry, *B* 3, 532)

Butyryl derivative of the ethyl ether $CH_3CH(O C_3H_7O)CO_2Et$ (208°) SG $\frac{1}{2}$ 1024 From a chloro-propionic ether and potassium butyrate (Wurtz, *A* 112, 235)

Isopropyl ether $CH_3CH(OH)CO_2Pr$ (167°) (Silva, *Bl* [2] 17, 97)

Ethylidene ether $CH_3CH<\begin{smallmatrix} O \\ \diagup \diagdown \\ CO \end{smallmatrix}>CHMe$ (151°) Formed by heating lactic acid with acetic aldehyde at 150° (Leipen, *M* 9, 46). Liquid, sol water Quickly decomposed by hot water into its components

Tri-chloro-ethylidene ether

$CH_3CH<\begin{smallmatrix} O \\ \diagup \diagdown \\ CO \end{smallmatrix}>CHCOCl$ [45°] (228°)

Prepared by heating syrupy lactic acid with excess of chloral for a short time at 150°, and

distilling the product with steam (Wallach, *A* 193, 1) Colourless crystals, readily sol alcohol, ether, and CS_2 , insol water

Methyl derivative $C_4H_8O_4$ \cdot $\frac{1}{2}$ $CH_3CH(OMe)CO_2H$ The Na salt is formed, together with the methyl ether, by treating $CH_3CH(ONa)CO_2Na$ with MeI The free acid is a syrup, volatile with steam It forms an amorphous silver salt $C_4H_7AgO_4$, $\frac{1}{2}$ sol water

Methyl ether of the methyl derivative $C_4H_8O_4$ \cdot $\frac{1}{2}$ $CH_3CH(OMe)CO_2Me$ (185°–188°) (Markownikoff & Krestownikoff, *A* 208, 343) From basic sodium lactate $CH_3CH(ONa)CO_2Na$ and MeI (Wislicenus, *A* 125, 53)

Ethyl ether of the methyl derivative $C_4H_8O_4$ \cdot $\frac{1}{2}$ $CH_3CH(OMe)CO_2Et$ (135° \cdot V) SG $\frac{1}{2}$ 9906 From a bromo propionic ether and $NaOMe$ (Schreiner, *A* 197, 1) Colourless liquid, nearly insol water

Ethyl derivative $C_4H_8O_4$ \cdot $\frac{1}{2}$ $CH_3CH(OEt)CO_2H$ **Ethyl lactic acid** (195°–198°) Obtained by decomposing its ether $CH_3CH(OEt)CO_2Et$ with caustic potash or lime Formed also, together with CH_3I , and acrylic acid, by the action of $NaOEt$ on iodoform (Butlerow, *A* 114, 206, 118, 325, *Bl* 1861, 9) Liquid, partly decomposed by distillation Miscible with water, alcohol, and ether, but separated from its aqueous solution by $CaCl_2$ or Na_2SO_4 Decomposes carbonates Alkalis do not convert it into lactic acid and alcohol When heated with conc $HIAq$ there is formed lactic acid and EtI

Salts — CaA' , 2aq flat prisms, $\frac{1}{2}$ sol water — AgA' bunches of slender silky needles (from hot water)

Ethyl ether of the ethyl derivative $C_4H_8O_4$ \cdot $\frac{1}{2}$ $CH_3CH(OEt)CO_2Et$ (155° \cdot V) (Schreiner, *B* 12, 179) SG $\frac{1}{2}$ 9498 VD 505 (calc 506) Formed by the action of $NaOEt$ on a chloro propionic ether (Wurtz, *A Ch* [3] 59, 169) Formed also by treating lactic ether with sodium and EtI (Wurtz & Friedel, *A Ch* [3] 63, 103) Also from silver ethyl lactate and EtI Colourless liquid, nearly insol water, sol alcohol and ether Alkalis convert it into alcohol and ethyl lactic acid

Amide of the ethyl derivative $CH_3CH(OEt)CONH_2$ **Lactamethane** [63°] (219°) Formed by allowing $CH_3CH(OEt)CO_2Et$ to stand a few days with aqueous NH_3 (Wurtz, *A Ch* [3] 59, 174) Broad plates, sol water, alcohol, and ether Decomposed by distillation with potash into NH_3 and lactic acid

Phenyl derivative $CH_3CH(OPh)CO_2H$ **Phenoxy propionic acid** [113°] Formed from a chloro propionic acid (25 g), strong caustic soda (to neutralisation), and sodic phenylate (24 g) The liquid is evaporated till it becomes thick, dissolved in water, and treated with HCl (L Saarbach, *J pr* 129, 152) Glassy needles (from water) Sol cold water, volatile with steam, $\frac{1}{2}$ sol hot water, alcohol, and ether. Aqueous solutions give a yellow pp with $FeCl_3$. Its salts are soluble in water — NaA' Exhibits rotatory action while dissolving in water Deliquescent — KA' 14aq at 180° — CaA' , 2aq — AgA' sharp needles, blackened by light

Ethyl ether of the phenyl derivative EtA' (244°) SG $\frac{1}{2}$ 1360 Formed when an alcoholic solution of the acid is allowed to stand

More rapidly by passing HCl into such a solution

Amide of the phenyl derivative
 $\text{CH}_3\text{CH}(\text{OPh})\text{CONH}_2$ [180°] Formed from the ether by aqueous ammonia. Crystallises from hot water in long needles. V sol alcohol and ether. Dissolves in hot HCl, on cooling crystals of the hydrochloride of the amide separate.

Bromo-phenyl derivative
 $\text{CH}_3\text{CH}(\text{OC}_6\text{H}_4\text{Br})\text{CO}_2\text{H}$ *Bromo phenoxy propionic acid* [106°] Formed by the action of bromine water on a solution of phenylated lactic acid. Crystallised from alcohol. V sol alcohol and ether, sl so water. Boiling aqueous NaOH cannot turn out the bromine, hence Br is in the benzene nucleus (Saarbach, *J pr* [2] 21, 157) — NaA. Deliquescent needles.

Thymyl derivative
 $\text{C}_6\text{H}_5\text{PrMe O CHMe CO}_2\text{H}$ Formed by heating thymol with α -chloro propionic acid in presence of a 50 p.c. solution of KOH, acidifying with HCl, and adding ammonium carbonate (Seichlone, *G* 12, 48). The acid from synthetical thymol crystallises in prisms [74°], v sol alcohol, ether, and chloroform. The acid from natural thymol crystallises in needles [48°], and forms very soluble and amorphous Ba and Ag salts.

p-Benzyl-phenyl derivative
 $\text{C}_6\text{H}_5\text{CH}_2\text{C}_6\text{H}_4\text{O CHMe CO}_2\text{H}$ [102°] From p-benzyl phenol, KOH, and α -chloro propionic acid (Mazzara, *G* 12, 264).

Benzyl p-tolyl derivative
 $\text{C}_6\text{H}_5\text{CH}_2\text{C}_6\text{H}_4\text{Me O CHMe CO}_2\text{H}$ [115°] Formed in like manner (Mazzara). Small crystals, sl sol water, v sol alcohol and ether. Its solutions give a yellow turbidity with ferric salts and crystalline pps with lead acetate and AgNO_3 .

Allophanyl derivative
 $\text{NH}_2\text{CO NH CO O CHMe CO}_2\text{H}$ [190°] Formed by passing gaseous cyanic acid into an ethereal solution of lactic ether and saponifying the resulting allophanyl lactic ether with conc HClAq at 100° (Traube, *B* 22, 1572). Minute colourless needles, m sol cold, v e sol boiling alcohol or water. When heated above 190° it splits up into lactic and cyanic acids — AgA' white powder, decomposed by boiling water — PbA', crystalline pp.

Ethyl ether of the allophanyl derivative Et' [170°] Colourless needles, v sol hot alcohol and hot water, v sl sol ether. Decomposed by alkalis into CO_2 , ammonia, alcohol, and lactic acid. Ammonia yields biuret.

Isoamyl ether $\text{C}_6\text{H}_{11}\text{A}$ [131°].

Monobasic anhydride

$\text{C}_6\text{H}_{10}\text{O}_5$ i.e. $\text{CH}_3\text{CH}(\text{OH})\text{CO O CHMe CO}_2\text{H}$ *Dilactic acid* Formed when aqueous lactic acid is left for several months over sulphuric acid *in vacuo* (Wislizenus, *A* 164, 181). Requires 1 mol of KOH for each mol of $\text{C}_6\text{H}_{10}\text{O}_5$ to neutralise it, but the neutral solution gradually becomes acid from liberation of free lactic acid, potassium lactate being also formed. By heating lactic acid at 140° this anhydride is formed together with lactide. It may also be formed from α -bromo propionic acid and potassium lactate (Brüggen, *Z* 1869, 338). It forms

amorphous Mg and Ca salts. The ethyl ether $\text{CH}_3\text{CH}(\text{OH})\text{CO O CHMe CO}_2\text{Et}$ (c 235°) S G 2 1 134 is formed by the action of chloro-propionic ether on potassium lactate (Wurtz & Friedel, *A Ch* [3] 63, 112). It is decomposed when heated with water into lactic acid and alcohol.

Di-ethyl ether of the di-basic anhydride $\text{C}_6\text{H}_{10}\text{O}_5$ i.e. $(\text{CH}_3\text{CH}(\text{CO}_2\text{Et}))_2\text{O}$ or $\text{CH}_3\text{CH}(\text{OEt})\text{CO O CHMe CO}_2\text{Et}$ (190° *in vacuo*) Formed by treating α -chloro propionic ether with $\text{CH}_3\text{CH}(\text{ONa})\text{CO}_2\text{Et}$ at 115° (Brüggen, *A* 148, 224). Scarcely attacked by conc KOHAq. Alcoholic KOH gives lactic acid and ethyl lactic acid. NH_3 in ether forms an oily amide $\text{C}_6\text{H}_{10}\text{NO}_5$, decomposed by KOH giving lactic and ethyl-lactic acids.

Neutral anhydride $\text{C}_6\text{H}_{10}\text{O}_5$ i.e.

$\text{CH}_3\text{CH} < \begin{smallmatrix} \text{CO} \\ \text{O} \end{smallmatrix} > \text{CH CH}_3$ *Lactide* [124 5°]

(255°) V D 4 81 (calc 4 96) (Henry, *B* 7, 753). Formed by the dry distillation of lactic acid (Gay Lussac & Pelouze, *A* 7, 43, Pelouze, *A* 53, 116, Engelhardt, *A* 70, 243, Wurtz & Friedel, *A Ch* [3] 63, 101). Prepared by passing dry air through lactic acid at 150° (Wislizenus, *A* 167, 318). Monoclinic tables (from alcohol). May be sublimed. V sl sol. hot water, but gradually converted thereby into lactic acid. Bases quickly convert it into lactic acid. Ammonia gives lactamide. Ethylamine gives the ethylamide of lactic acid. $\text{CH}_3\text{CH}(\text{OEt})\text{CO}_2\text{Et}$ forms a compound $\text{C}_{11}\text{H}_{20}\text{O}_5$ (270°) decomposed by potash into alcohol and lactic acid.

Nitrile $\text{C}_6\text{H}_9\text{NO}$ i.e. $\text{CH}_3\text{CH}(\text{OH})\text{CN}$ *Aldehyde cyanhydrin* (183°) Formed in the cold by allowing a mixture of aldehyde (1 mol.) and anhydrous HCN to stand for 9 days (Maxwell Simpson & Gautier, *Bl* [2] 8, 277). Liquid, not solidified at -21°. Partially decomposed into its components by distillation. Miscible with water, alcohol, and ether. Aqueous KOH forms KCy and aldehyde (or aldehyde resin). Conc HClAq acts violently upon it, at 0° the products are lactic acid and NH_4Cl .

Sarcolactic acid $\text{C}_6\text{H}_8\text{O}_5$ i.e.

$\text{CH}_3\text{CH}(\text{OH})\text{CO H}$ *Paralactic acid* (Heintz, *P* 75, 391) $[\alpha]_D = 3.5^\circ$. Occurs in muscular tissue, thymus and thyroid gland, urine after much exercise, spleen, lymphatic glands, and in pig's bile (Liebig, *A* 62, 278, 326, Wislizenus, *A* 167, 302, Strecker, *A* 123, 354, Colasanti & Moscatelli, *H* 12, 416, *G* 17, 548, 18, 548, Marcuse, *BC* 1887, 92, Nebelhaus, *Zent Biol* 25, 123, Hirschler, *H* 11, 41, Gleiss, *Pf* 41, 69). Sarcolactic acid is absent from living blood, but occurs in blood after death (Salomon, Virchow's *Archiv*, 113, 356, cf. Berlinerblau, *C C* 1888, 757, Vissokovitch, *C C* 1888, 117). Sarcolactic acid may occur sometimes along with ordinary lactic acid as a product of fermentation (Maly, *B* 7, 1567). Its presence is then due to *micrococcus acidiparalactici*, which can convert glucose into sarcolactic acid (Nenoki & Sieber, *M* 10, 532). Sarcolactic acid is formed in the fermentation of moists by cheese (Helger, *A* 160, 336). It is also formed by the action of nitrous acid on the amido propionamide present in urine. A dextro-

rotatory lactic acid is formed by the action of *Penicillium glaucum* on ordinary ammonium lactate (Lewkowitzh, *B* 16, 2720)

Preparation.—1 Chopped flesh is exhausted with cold water, the extract mixed with baryta water, the albumen coagulated by boiling and removed by filtration, and the clear liquid concentrated by evaporation. Sulphuric acid is added to the syrupy residue, and it is shaken with ether, which leaves sarcosolactic acid when evaporated—2 Extract of meat (1 pt) is dissolved in warm water (4 pts) and added with 90 p.c. alcohol. The filtrate is evaporated to a syrup, mixed with 4 pts of alcohol, filtered, evaporated, acidified by H_2SO_4 and extracted with ether (Khimenko, *J R* 12, 17, *Bl* [2] 34, 321)

Properties.—Dextrorotatory syrup, forming levorotatory salts. Resembles ordinary lactic acid in its reactions. In a dry atmosphere it changes in the cold to a levorotatory anhydride $[\alpha]_D = -86^\circ$. At 140° it forms ordinary lactide, whence water produces ordinary inactive lactic acid (Strecker, *A* 105, 313)

Reactions.—1 Hot dilute sulphuric acid splits it up into formic acid and aldehyde—2 Chromic acid mixture gives CO_2 and acetic acid

Salts.— CaA' , 4aq (Engelhardt, *A* 65, 359) S 8 in the cold. V sol boiling water and alcohol— CaA' , 5aq changes into CaA' , 5aq on recrystallisation— MgA' , 4aq. More soluble in water and alcohol than ordinary magnesium lactate— NiA' , 3aq— ZnA' , 2aq. Forms more distinct crystals than ordinary zinc lactate. S (of ZnA' , 2aq) 5.7 at 14.5° , the solubility of ordinary zinc lactate being 1.7 (Wislicenus) S (98 p.c. alcohol) 10.4— ZnA' , 3aq. Ppd by adding alcohol to an aqueous solution of the zinc salt— AgA' , 1/2aq flat needles

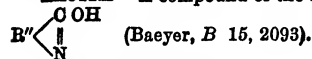
Ethyl ether EtA' $[\alpha]_D = -14.2^\circ$. From the silver salt and EtI (Khimenko)

Chloro lactic acid *v* **CHLORO OXY-PROPIONIC ACID**

Chloride of lactic acid *v* **Chloride of α -CHLORO-PROPIONIC ACID**

LACTIDE *v* **Neutral anhydride of LACTIC ACID**

LACTIM A compound of the form



LACTIMIDE C_3H_5NO *s*

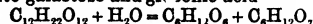
$CH_3CH \begin{array}{c} \diagup \text{CO} \\ | \\ \text{NH} \end{array}$ $[275^\circ]$ Produced together with ethylamine and CO_2 by heating alanine in dry HCl at 190° as long as water comes off (Preu, *A* 134, 872). The brown amorphous product is treated with lead hydroxide and H_2S and recrystallised from alcohol. Colourless needles, may be sublimed. V sol water and alcohol. Has a bitter taste. Its solution does not dissolve Ag_2O or give a pp with $ZnCl_2$ or $AgNO_3$.

LACTO-ALBUMEN *v* **MILK**

LACTO-ALBUMOSE *v* **MILK**

LACTO-ZONIC ACID $C_{12}H_{21}O_{17}$. Formed by treating a solution of milk sugar (1 pt) in water (7 pts) with bromine (1 pt) at ordinary temperatures for some days, the bromine is next removed by a stream of air and H_2S , the HBr is

removed with white lead followed by Ag_2O and the metals with H_2S , on treating the syrup so obtained with glacial acetic acid the lacto bionic acid is left. The product is purified by conversion into lead salt, with specially prepared basic lead acetate, and decomposition of the same with H_2S (Emil Fischer & Jacob Meyer, *B* 22, 362). Colourless syrup of strong acid reaction, v sol water, v sl sol alcohol, insol ether. Shows no tendency to crystallise. Does not reduce alkaline copper solutions even on boiling. On warming a short time with dilute mineral acids it splits up into galactose and glycolic acid

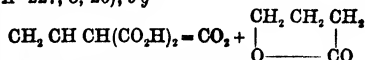


LACTOCYANAMIDE *v* **LACTIC ACID**, p 112

LACTONES Anhydrides of oxy acids formed by elimination of water between the hydroxyl and carboxyl groups, both being in the same carbon chain. The name is derived from *lactide* which, until its vapour density had been determined, was written $CH_3CH \begin{array}{c} \diagup \text{O} \\ | \\ \text{CO} \end{array}$. Lactones are usually derived from γ - or from δ -oxy acids. The formula $R''R'C \begin{array}{c} \diagup \text{CH}_2\text{CH}_2 \\ | \\ \text{O CO} \end{array}$ represents a γ lactone, while

$R''R'C \begin{array}{c} \diagup \text{CH}_2\text{CH}_2 \\ | \\ \text{O CO} \end{array} > CH_2$ is a δ lactone (Fittig, *A* 200, 21, 203, 67, 216, 52)

Formation.—1 γ Oxy acids split up in the cold, and immediately on heating their aqueous solution into water and a γ -lactone—2 Formed by boiling their carboxylic acids with dilute H_2SO_4 (Erdmann, *A* 228, 176)—3 From esters of γ oxy acids on distillation, alcohol being split off, *e.g.* oxy isocaproic ether (Bredt), oxy valeric ether (Kissling)—4 From unsaturated acids (*v* *A* 227, 8, 26), *e.g.*



Properties.—Volatile liquids, neutral to litmus. Volatile with steam.

Reactions.—1 γ Lactones do not take up water to form an oxy acid by mere boiling. On the other hand most δ lactones in presence of water, cold or hot, are partially converted into acid, and tend to assume equilibrium with 65 p.c. of lactone to 35 p.c. of acid (Fittig & Wolff, *A* 216, 187, Hantzsch, *A* 222, 28)—2 All lactones are converted into salts of the corresponding oxy acids by boiling with aqueous solutions of alkalis, alkaline earths, and sometimes even with $CaCO_3$ (Fittig, *A* 208, 116). 3 All lactones form compounds with NH_3 , which easily split up into their components (Wolff, *A* 229, 278). These compounds are probably acid amides—4 No lactone at present known reacts with hydroxylamine, but a few aromatic lactones readily react with phenylhydrazine (*e.g.* phthalide). Hence the fact of an oxygen compound reacting with phenylhydrazine cannot be taken (as previously assumed) as a proof of its aldehydic or ketonic nature. This can only be proved by its behaviour towards hydroxylamine (V Meyer & Münchmeyer, *B* 19, 1706). The compound formed with phthalide appears to be $HOCH_2C_6H_4CO \cdot N_2H_5$, ph formed by direct addition (Wislicenus, *B* 20, 401)—5 Lactones (1 mol) appear to react with oxalic ether (1 mol.) in presence of $NaOEt$

(1 mol) Phthalide gives $C_{12}H_{10}O_5$ [122°] and valerolactone also gives a crystalline compound (Wilschusen, *B* 20, 2061)

Lactones are for the most part described in this Dictionary under the oxy acids of which they are the anhydrides

LACTONIC ACID v GALACTONIC ACID

Lactonic acids Acids which are at the same time lactones When warmed with alkalis they give rise by assimilation of water to acids of higher basicity

LACTONITRILE v Nitrile of LACTIC ACID

LACTO-PROTEIN v MILK

LACTOSE or Milk sugar v SUGAR and MILK

LACTOSIN $C_{12}H_{22}O_{11}$ Occurs in the roots of all the Caryophyllaceae, being most conveniently obtained from *Silene vulgaris* (A Meyer, *B* 17, 685) White amorphous powder (anhydr), $[a]_D = +168$, or small glistening crystals ($+H_2O$), $[a]_D = +211.7$ It forms gummy solutions with water, but is sparingly soluble in alcohol The aqueous solution of the crystallised lactosin gives no pps with neutral or basic lead acetate, although a pp is produced by lead acetate and NH_4 Fehling's solution is not reduced on short boiling By boiling with dilute H_2SO_4 it is inverted into lactose and a new sugar

LACTUCARIUM A brownish viscid substance obtained by evaporating juice which exudes from incisions in the leaves and stem of certain species of lettuce, especially *Lactuca virosa* It has an odour resembling opium and acts as a narcotic According to Ludwig (*Ar Ph* [2] 7, 129) it contains (50 p c of) 'lactucone' $C_{14}H_{26}O_8$, a crystalline substance [150°–160°] soluble in boiling alcohol, 'lactucic acid,' lactucin, a wax, and oxalic acid According to Hesse (*A* 234, 243) lactucarium contains the acetyl derivatives of (a) and (b) lactucerin

LACTUCERIN $C_{14}H_{26}O_8$ (L), or $C_{12}H_{24}O_8$ (K), or $C_{10}H_{20}O_8$ (H) *Lactucone* [210°] Obtained by washing lactucarium with benzene and extracting the residue with boiling alcohol, the crystals obtained may be purified by shaking their ethereal solution with aqueous KOH, and ppg by the addition of alcohol and water (Lenoir, *A* 60, 83, Kassner, *A* 238, 220) Lactucerin so prepared forms minute white needles [200°], but after sublimation in CO_2 it melts at 210° On fusion with KOH it gives lactucol $C_{14}H_{26}O_8$, hydrogen, and acetic acid It is dextrorotatory According to Hesse, lactucerin consists of the mono-acetyl derivatives of (a) and (b) lactucerinol From the milk juice of *Lactuca canadensis*, by crystallising from hot alcohol Flowers (*Ph* [3] 10, 44) obtained a 'lactucern' [89°] crystallising in colourless needles

Lactucone $C_{14}H_{26}O_8$ (?) [296°] Microscopic needles Insoluble in water, difficultly soluble in alcohol Occurs in French lactucarium from *Lactuca altissima*, from which it is extracted with 90 p c alcohol Acetic anhydride does not act upon it even at 200° By distillation with P_2S_5 it gave a hydrocarbon of the constitution $C_{14}H_{22}$, boiling between 247°–252° (Franchimont, *B* 12, 10)

Lactucol $C_{14}H_{26}O_8$ [162°] Obtained by fusing lactucerin with KOH (Kassner, *A* 233, 224) Needles Dextrorotatory, $[a]_D = 46^\circ$

Acetyl derivative $C_{16}H_{28}AcO$ [200°]

Like lactucol and lactucerin its solutions in ether, chloroform, and CS_2 are dextrorotatory, $[a]_D = 68^\circ$

(a)-**LACTUCEROL** $C_{14}H_{26}O_8$ [166°–181°] Obtained by exhausting lactucarium with ligroin and extracting the residue with alcohol The crystals so obtained are saponified by alcoholic potash, and the product ppg with water The pp is recrystallised from alcohol, from which (a) lactucerinol separates first (Hesse, *A* 234, 245, 244, 268)

Properties—Crystallises from 90 p c alcohol in silky needles (containing 2aq) From chloroform or ether it separates in anhydrous crystals. Insol water and alkalis It melts at 162°, but after purification by conversion into its di-acetyl derivative and saponification of the product its melting point is higher Conc H_2SO_4 colours its solution in chloroform red It absorbs Br with evolution of HBr It is dextrorotatory, in a 2.3 p c solution in chloroform $[a]_D = 76.2$ at 15° It may be distilled in a current of CO_2

Acetyl derivative $C_{16}H_{28}AcO$ [202°–207°] Occurs in lactucarium, and is formed by heating (a) lactucerinol for a short time with Ac_2O at 80° Small plates, m sol cold alcohol, v sol ether Dextrorotatory

Di-acetyl derivative $C_{16}H_{28}Ac_2O$ [198°–200°] From (a) lactucerinol and Ac_2O by boiling for 2 hours Satiny plates In a 1 p c solution in chloroform $[a]_D = 63.6$ at 15°

Di-propionyl derivative $C_{16}H_{28}(C_2H_5O)_2$ [152°] Minute needles (from alcohol), v e sol $CHCl_3$, and ether

Di-benzoyl derivative $C_{16}H_{28}Bz_2O_2$ [156°] White crystals, insol water, v sol ether and chloroform, sl sol alcohol Saponified by alcoholic potash

(b)-**Lactucerinol** $C_{12}H_{24}O_8$ $[a]_D = 38$ at 15° in a 4 p c solution in chloroform Occurs as a mono acetyl derivative in lactucarium, and separated from its (a)-isomeride by crystallisation from alcohol Long silvery needles (from ether or chloroform) Separates from alcohol as a gelatinous mass (containing 2aq) More soluble in alcohol, and less dextrorotatory than its isomeride

Acetyl derivative $C_{14}H_{26}Ac_2O$ [230°] Plates (from alcohol) Less sol alcohol and ligroin than its isomeride

LACTUCIC ACID Obtained from the juice of *Lactuca canadensis* after separating 'lactucern,' adding water, ppg with lead acetate, decomposing the lead salt with H_2S and evaporating (Flowers, *Ph* [3] 10, 44) Brownish green amorphous substance, with acid, bitter taste. Sol alcohol, insol petroleum spirit, ether, and chloroform Ludwig (*J* 1847, 824) and Wals (*N Jahr Pharm* 15, 118) obtained a substance ($C_{10}H_{20}O_8$, ?) called lactucic acid by triturating lactucarium (1 pt) with dilute H_2SO_4 (1 pt), adding alcohol (5 pts of 84 p c), filtering, shaking the filtrate with slaked lime, decolourising with animal charcoal, evaporating, and crystallising the residue from boiling water Light yellow amorphous mass, gradually becoming crystalline Its solutions are coloured wine-red by alkalis, and reduce boiling Fehling's solution.

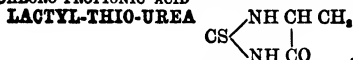
LACTUCIN $C_{14}H_{26}O_8$, or $C_{12}H_{24}O_8$, or $C_{10}H_{20}O_8$ S 125 in the cold Occurs in the juice of common lettuce (*Lactuca sativa*) and of *Lactuca*

althusma (Aubergier, *B J* 24, 522, *A* 44, 299, Wals, *A* 82, 85, *N Jahr Pharm* 15, 118, Ludwig & Kromayer, *Ar Ph* [2] 111, 1, Kromayer, *Ar Ph* [2] 105, 3, Buchner, *Rep Pharm* 43, 1, Flowers, *Ph* [3] 10, 44) Obtained by macerating the dried juice with hot water ($\frac{1}{2}$ pt) for four days, pressing the mass, boiling the residue with water, and ppg the filtrate with lead subacetate, removing excess of lead from the filtrate by H_2S , and evaporating Pearly scales (from alcohol). M sol alcohol and HOAc, sl sol ether Conc HNO_3 turns it brown Its solutions are not ppg by ordinary reagents

LACTUCOPICRIN $C_{14}H_{16}O_{11}$. An amorphous very bitter substance which remains in the mother-liquor in the preparation of lactucin (Kromayer, *Die Bitterstoffe*, 1861, Flowers, *Ph* [3] 10, 222) Sol water and alcohol, its solutions not being ppg by lead salts

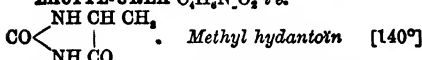
LACTURAMIC ACID ν **URAMIDO PROPIONIC ACID**

LACTYL CHLORIDE is the chloride of α -CHLORO PROPIONIC ACID



From this urea and a chloro propionic ether at 100° (Freytag, *J pr* [2] 20, 380) Crystalline.

LACTYL-UREA $C_4H_6N_2O_4$: a.



Methyl hydantoin [140°]

(H), [145°] (Urech, *B* 6, 1113) Formed by treating aldehyde ammonia with crude KCy and hydrochloric acid $CH_3CH(OH)NH_2 + HCN + HClNO = NH_4 + C_2H_5N_2O_2$ (Heintz, *A* 169, 120) Formed also by heating uramido propionic acid Prisms, or cauliflower-like tufts (containing aq), v sol water and alcohol, nearly insol ether May be sublimed Tastes bitter Neutral in reaction Boiling baryta water converts it into uramido-propionic acid Heating with barium hydrate at 100° – 140° gives alanine Pure HNO_3 gives a quantitative yield of a nitro derivative, but no gas is evolved (Franchimont, *R T C* 6, 217) — $AgC_2H_5N_2O_2$ formed by treating lactyl urea with moist Ag_2O Insol water, sol NH_4Aq Ppd as a white powder by adding HNO_3 to its ammoniacal solution

LEVULAN $C_7H_{10}O_6$ [250°] $[\alpha]_D = -221^\circ$ Occurs in an impure condition in the residues obtained from desugarising molasses by Steffen's process (Lippmann, *B* 14, 1509) White amorphous powder Sol hot water, the solution gelatinises on cooling By long boiling it becomes much more soluble and loses its power of gelatinising Insol alcohol It does not reduce Fehling's solution but produces a blue pp By HNO_3 it is oxidised to mucic acid Dilute H_2SO_4 at 120° converts it into levulose

LEVULIN $C_5H_8O_4$ (dried at 110°) A substance resembling dextrin, contained in the juice of the tubers of the Jerusalem artichoke (*Helianthus tuberosus*) and of *Dahlia variabilis*, and in oak bark (Ville & Joulie, *Bl* [2] 7, 262, Popp, *A* 156, 181, Dieck & Tollens, *A* 198, 228, *B G* 1879, 275, Etti, *B* 14, 1826, Lefranc, *J Ph* [5] 2, 216, Reidemeister, *J Th* 1881, 68) Prepared by ppg the juice of artichokes with lead acetate, filtering, removing excess of lead by

H_2S , neutralising with $MgCO_3$, filtering and evaporating The residue is extracted with alcohol, until it is optically inactive It is then extracted with absolute alcohol, which leaves inulin undissolved, and ppg with ether

Properties —Amorphous deliquescent mass, optically inactive, but becomes levorotatory on boiling with dilute HCl, being split up into levulose and glucose V sol water and dilute alcohol, sl sol absolute alcohol, insol ether Has an insipid taste Turns brown at 140° forming caramel Not ppg by lead subacetate Reduces Fehling's solution after long boiling Dilute HNO_3 gives oxalic acid and saccharic acid Alkalis do not turn it brown First hydrolysed by yeast and then undergoes alcoholic fermentation Levulin prevents the ppg of ferric and cupric salts by alkalis $AgNO_3$ gives a white pp blackened on heating — $K_2C_2H_3O_6$ ppg by adding alcoholic KOH to its alcoholic solution — $Ba_2C_2H_3O_6$ aq amorphous — $Pb_2C_2H_3O_6$ aq amorphous, ppg by adding an alcoholic solution of lead subacetate to an alcoholic solution of levulin

LEVULINIC ACID ν β **ACETYL PROPIONIC ACID**

LEVULOSE ν **SUGAR.**

LANTHANUM La At w 138.2 Mol w. unknown SG 6.163 Melts between M P of Sb (450°) and that of Ag (950°) (Hillebrand & Norton, *P* 156, 466) SH 04485 (H & A, *N P* 158, 71) SVS 22.4 Chief lines in emission spectrum 5183, 4921, 4920 in the green, 4824, 4655, 4558 in the blue, 4522, 4330, 4268 in the indigo, 4238, 4196, 4086, 4077 in the violet (Thalén) In 1803 Klaproth separated a new earth from a Swedish mineral, the earth was examined by Hisinger and Berzelius and called by them *ceria* (from the planet *Ceres* then recently discovered) In 1839 Mosander showed that *ceria* contained two oxides (*P* 46, 648) the name *ceria* was retained for one, and the other was given the name *lanthana* ($\lambda\alpha\nu\theta\acute{\alpha}\nu\epsilon\iota\nu$ = to be concealed) In 1841 Mosander discovered that *lanthana* was a mixture of two oxides (ν *P* 60, 297), one of these he called *lanthana* and the other *didymia* ($\delta\acute{\iota}\delta\upsilon\mu\omicron\varsigma$ = twofold)

Occurrence —As silicate, with silicates of Ce and Di, in *cerite*, *gadolinite*, *orthite*, &c (Rammelsberg, *P* 107, 631) *Lanthanite*, from Bethlehem in Pennsylvania, U.S., is a carbonate of La and Di (Lawrence Smith, *Am S* [2] 18, 378) La containing minerals occur in small quantities in a few localities

Preparation —The mixed oxides of Ce, La, and Di are separated from *cerite* by treatment with H_2SO_4 , &c, as described under *Cerium* (vol 1 p 723), the oxides are dissolved in HNO_3Aq , the solution is evaporated to dryness the residue is heated to full redness until pale yellow, and then treated with boiling dilute HNO_3Aq in which nitrates of La and Di dissolve while basic Ce nitrate remains insoluble There are various ways of separating La from Di in the nitric acid solution The hydrated oxides may be obtained by ppg with NH_4Aq , the pp is dissolved in H_2SO_4Aq , and the mixed sulphates are obtained by crystallisation, the sulphates are dried and powdered, 1 part is dissolved in small successive portions in 6 parts

water at 2°-3°, the solution is then heated to ca. 40° when $\text{La}_2(\text{SO}_4)_3$ separates nearly free from $\text{D}_1(\text{SO}_4)_3$. The $\text{La}_2(\text{SO}_4)_3$ may be purified by re solution in ice cold water and heating to 40°, the operations being repeated until the sulphate is perfectly white and shows no D₁ lines in the spectrum. Or the solution of $\text{La}_2(\text{SO}_4)_3$ may be pptd by oxalic acid, the pp strongly heated, dissolved in HNO_3 , and fractionally pptd by NH_4Aq , the ppn must be effected from very dilute solutions by means of very dilute NH_4Aq , the first third of the pp contains most of the D₁ oxide, ppn is continued so long as the pp shows D₁ lines in the spectrum. This method gives good results when comparatively much La is present (Mosander, *l.c.*, Bunsen a. Jøgel, *P* 155, 377), but it is slow and laborious. Marignac (*J pr* 48, 406) adds excess of HNO_3 to the solution of the nitrates of La and D₁, heats, and adds oxalic acid, the pp dissolves, when crystallisation begins the solution is allowed to cool, and is then poured off from the pp which is rich in D₁ oxalate, these operations are repeated, a very acid solution of La salt is finally obtained from which NH_4Aq pptts $\text{La}_2\text{O}_3 \cdot x\text{H}_2\text{O}$. This method is recommended when much D₁ is present with little La.

The method of Von Welsbach (*Sitz W* 92 [2nd part], 317) was found very good by Robinson (*priv comm*). A large quantity of the mixed nitrates of Ce and La obtained from *cerite* after separating basic Ce nitrate (*v* vol 1 p 723) is mixed with the necessary quantity of NH_4NO_3 , about $\frac{1}{10}$ th part conc HNO_3Aq is added, and the liquid is evaporated until small crystals appear on the surface, a little water is then added, and crystallisation is allowed to proceed for about 24 hours, the crystals are drained and washed with a little HNO_3Aq which is added to the mother liquor, the mother liquor is evaporated and crystallised, the liquor from this is again evaporated, and so on until 6-8 fractions are obtained. The first fraction contains almost all the La. By fractionating the middle fraction, placing the first fractional pp in the former first fraction, and repeating this process a few times, pure $\text{La}(\text{NO}_3)_3$ is obtained. For other methods of separating La salts from salts of Ce and D₁ v Hermann, *J pr* 82, 385, Erk, *Z* [2] 7, 100, Cleve, *Bl* [2] 21, 196, 246.

La_2Cl_6 is obtained by dissolving the pp of $\text{La}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ (obtained as described above) in HClAq , adding NH_4Cl , evaporating to dryness, and heating strongly in a covered crucible. By reducing La_2Cl_6 by heating with K and washing out KCl in alcohol, metallic La is obtained. Hillebrand a. Norton obtained approximately pure La by electrolysis La_2Cl_6 covered with a fused mixture of KCl and NaCl, using a thick iron wire as negative electrode and a battery of 4 Bunsen's cells (*P* 156, 466, cf Bunsen, *P* 155, 633).

Properties and Reactions—White metal, fairly malleable and ductile, rather harder than Ce. Oxidises rapidly in dry air, but burns only at temperature considerably higher than that at which Ce burns. Decomposes cold water slowly, hot water rapidly. Easily dissolved by acids, including cold conc HNO_3 , which scarcely acts on Ce.

The atomic weight of La has been deter-

mined (1) by converting the oxide into the sulphate and *vice versa* (Rammelsberg, *P* 55, 65, Marignac, *A Ch* [8] 27, 228, [4] 30, 67, Holzmänn, *J pr* 75, 348, Czudnowicz, *J pr* 80, 33, Hermann, *J pr* 82, 395, Zschiesche, *J pr* 104, 174, Erk, *Z* [2] 7, 106, Cleve, *Bl* [2] 81, 196, Brauner, *O J* 41, 75, Crookes, *Pr* 38, 414, (2) by analysing La iodate (Holzmänn, *J pr* 75, 349), (3) by estimating Cl in La chloride (Hermann, *J pr* 82, 395), (4) by converting La carbonate into oxide (Hermann, *l.c.*), (5) by determining S H of La (Hillebrand a. Norton, *P* 158, 71), (6) by considering the chemical relations of La with other elements in the light of the periodic law.

The at w of La was taken for many years as c 92.2, the oxide was formulated LaO and the chloride LaCl . Mendeleeff (*v C N* 41, 49) proposed to multiply the usually accepted at w by 2, and to regard the oxide as LaO_2 , by doing this he placed La in Group IV along with Ce. As the properties of La salts were not much known when Mendeleeff's memoir was published, he did not strongly press the arguments in favour of the position assigned by him to La. Fuller investigation showed that the usually accepted at w of La should be increased by one half, that the oxide should be regarded as similar to the oxides of the earth metals (M_2O_3), and that La should be placed in Group III along with Al, Ga, Sc, &c (*v* **EARTHS, METALS OF THE**, vol II p 424).

La is distinctly metallic in its chemical relations, it forms the oxide La_2O_3 , and there are indications of the existence of a higher oxide, the chloride is La_2Cl_6 or LaCl_3 . La forms several

salts of the form LaX_3 , $\text{X} = \text{NO}_2, \frac{\text{SO}_4}{2}, \frac{\text{PO}_4}{3}$

&c, a few double salts and one or two basic salts are known. The investigation of the La compounds shows that this metal is to be placed with the metals of the earths (Group III), it is more closely related to the even series members of the group (Sc, Y, and Yb) than to the odd series members (Al, Ga, In, Tl). The strongly basic character of La_2O_3 marks the connection of La with the alkaline earths and alkali metals. The examination of La compounds is yet far from complete (*cf* **METALS, RARE**).

Detection and Estimation—Most of the La salts are colourless, the soluble salts have an astringent, sweetish taste, solutions of La salts do not show any absorption-bands. Alkalis, NH_4HS , and KCN, form gelatinous pps insoluble in excess of pptant. Alkali carbonates ppt $\text{La}_2(\text{CO}_3)_3$, BaCO_3 forms a pp without warming, $\text{H}_2\text{C}_2\text{O}_4$ forms a white pp at first curdy, then crystalline, more soluble in acids than the oxalates of Ce and D₁, $\text{Na}_2\text{S}_2\text{O}_3$ does not produce any pp. La salts do not colour beads of borax or microcosmic salt.

La may be estimated as La_2O_3 or $\text{La}_2(\text{SO}_4)_3$. La_2O_3 is obtained (1) by ppg with NH_4Aq , washing as rapidly as possible with water containing NH_3 (to prevent formation of La_2CO_3 and partial solution of La), solution in HNO_3Aq , re ppn by NH_4Aq , washing with NH_4Aq , and strongly heating, (2) by ppg $\text{La}_2(\text{C}_2\text{O}_4)_3$ by addition of $\text{H}_2\text{C}_2\text{O}_4$ and standing, washing, and heating to white heat. $\text{La}_2(\text{SO}_4)_3$ is obtained by ppg with

NH_4Aq , as above, dissolving in warm dilute $\text{H}_2\text{SO}_4\text{Aq}$, evaporating to dryness at 100° , and gradually heating to redness

Lanthanum, arsenate of $\text{La}_2(\text{HASO}_4)_3$, and arsenite of $\text{La}_2(\text{HASO}_3)_3$, v Smith, *A* 191, 331 Lanthanum, borate of $(2\text{La}_2\text{O}_3 \cdot \text{B}_2\text{O}_3)_n$, obtained, along with crystals of La_2O_3 , by dissolving La_2O_3 in molten borax (Nordenskjöld, *J* pr 85, 431)

Lanthanum, bromide of $\text{La}_2\text{Br}_6 \cdot 14\text{H}_2\text{O}$, or $\text{LaBr}_3 \cdot 7\text{H}_2\text{O}$. Colourless crystals, e sol water or alcohol, obtained by dissolving $\text{La}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ in HBrAq and evaporating (Cleve, *Bl* [2] 89, 151, 43, 56) By the action of Br in vapour on La_2O_3 , the oxybromide LaOBr is obtained (Frerichs a Smith, *A* 191, 331) Double salts are described by Cleve (*l c*) — $\text{La}_2\text{Br}_6 \cdot 2\text{AuBr}_3 \cdot 18\text{H}_2\text{O}$, $\text{La}_2\text{Br}_6 \cdot 3\text{ZnI}_2 \cdot 27\text{H}_2\text{O}$, also by F a S (*l c*), $\text{La}_2\text{Br}_6 \cdot 3\text{NiBr}_2 \cdot 18\text{H}_2\text{O}$, and $\text{La}_2\text{Br}_6 \cdot 3\text{ZnBr}_2 \cdot 39\text{H}_2\text{O}$

Lanthanum, carbide of A carbide of La is said to be produced by heating the oxalate or formate in absence of air, it is described as similar to, but more easily acted on by acids than, Ce carbide (Delafontaine, *C N* 11, 253)

Lanthanum, chloride of La_2Cl_6 or LaCl_3 A white crystalline mass, e sol water or alcohol Obtained by adding NH_4Cl to a solution of $\text{La}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ in HClAq , evaporating to dryness, and heating in a closed crucible until all NH_4Cl is volatilised (Hermann, *J* pr 82, 406, Hillebrand a Norton, *P* 158, 71) Mosander (*P M* 23, 241) says that LaCl_3 can be obtained by evaporating $\text{La}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ in HClAq to dryness, and heating the residue in a stream of HCl gas

By slowly evaporating $\text{La}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ in HClAq , large colourless triclinic crystals of $\text{LaCl}_3 \cdot 7\text{H}_2\text{O}$ are obtained Double salts are described by Smith (*A* 191, 331), and Cleve (*Bl* [2] 21, 196, 39, 151) — $\text{M} 9\text{H}_2\text{Cl}_2 \cdot 24\text{H}_2\text{O}$,

$\text{M} 3\text{PtCl}_2 \cdot 24\text{H}_2\text{O}$, $\text{M} 2\text{PtCl}_2 \cdot 26\text{H}_2\text{O}$, $\text{M} 8\text{AuCl}_3 \cdot 21\text{H}_2\text{O}$, $\text{M} 2\text{AuCl}_3 \cdot 20\text{H}_2\text{O}$ ($\text{M} = \text{La}_2\text{Cl}_6$) Oxychlorides are obtained by heating the hydrated chloride in air, and by the action of Cl on La_2O_3 (v LANTHANUM, OXYCHLORIDES OF)

Lanthanum, chromate of $\text{La}_2(\text{CrO}_4)_3 \cdot 8\text{H}_2\text{O}$, v. Smith, *A* 191, 355

Lanthanum, cyanide of LaCy_3 , v. vol u p. 341

Lanthanum, fluoride of $\text{La}_2\text{F}_6 \cdot \text{H}_2\text{O}$ A gelatinous pp by adding HFAq to solution of La acetate (Cleve, *Bl* [2] 89, 151, 43, 56) From solution of La sulphate, Smith obtained a pp of $\text{La}_2\text{F}_6 \cdot 3\text{HF}$ (*A* 191, 331) According to Marignac, H_2SiF_6 pptts La_2F_6 from La salts (*J* pr 48, 406)

Lanthanum, hydroxide of $\text{La}_2\text{O}_3 \cdot \text{H}_2\text{O}$, or La_2O_3 , may also be regarded as *hydrated oxide* $\text{La}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$ A white gelatinous pp by adding KOH Aq or NaOH Aq to solution of a La salt (NH_4Aq pptts basic compounds) Also produced by action of warm water on La_2O_3 $\text{La}_2\text{O}_3 \cdot \text{H}_2\text{O}$ is a strongly basic hydroxide, it turns red litmus blue, decomposes NH_4ClAq on warming with evolution of NH_3 , absorbs CO_2 from the air, reacts with acids to form salts

LaX_3 ($\text{X} = \text{NO}_2, \frac{\text{SO}_4}{2}, \frac{\text{PO}_4}{3}$, &c) Thomsen gives

the heat of neutralisation with $\text{H}_2\text{SO}_4\text{Aq}$ and HClAq as $[\text{La}^\circ\text{O}^\circ\text{H}^\circ \cdot 3\text{H}^\circ\text{SO}^\circ\text{Aq}] = 82,820$; $[\text{La}^\circ\text{O}^\circ\text{H}^\circ \cdot 6\text{HClAq}] = 74,970$ (*Th.* 1, 375).

Lanthanum, haloid compounds of. LaF_3 , LaCl_3 , and LaBr_3 have been isolated, the formulae may be written La_2F_6 , &c, as none of these compounds has been gasified All form hydrates, and all combine with haloid compounds of Au and some other heavy metals to form double salts Oxychlorides and oxybromides, LaOX , are known

Lanthanum, iodide of LaI_3 has not been isolated, but the double salt $2\text{LaI}_3 \cdot 3\text{ZnI}_2 \cdot 27\text{H}_2\text{O}$ is described by Frerichs a Smith (*A* 191, 355)

Lanthanum, oxides of Only one oxide, La_2O_3 , is known with certainty, there are indications of the existence of an oxide containing more O

LANTHANUM SESQUIOXIDE, La_2O_3 Obtained by strongly heating $\text{La}_2\text{O}_3 \cdot x\text{H}_2\text{O}$, or the oxalate, or any La salt the acid of which is volatilisable Forms a white, amorphous, infusible powder, SG 6.48–6.53 (Cleve, *l c*, Nilsson a Pettersson, *B* 13, 1464) SH 0749 (N a P, *l c*, v also *Pr* 31, 46) Diamagnetic (N a P, *l c*) Combines with water to form $\text{La}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$ (v LANTHANUM, HYDROXIDE OF) Easily sol in acids Nordenskjöld (*J* pr 85, 431) obtained La_2O_3 in lustrous rhombic crystals, $abc = 56581 \cdot 6863$, SG 5.296, by dissolving the amorphous oxide in borax in a porcelain oven, the crystals did not directly combine with water, but were easily sol in acids The great infusibility of La_2O_3 , and its power of emitting white light when strongly heated, render it useful as a light giver, Von Welsbach has patented an arrangement whereby La_2O_3 is heated in the flame of a Bunsen lamp and emits a clear white light (English Patent, 15,286, July 1886).

LANTHANUM PEROXIDE According to Mosander (*P M* 23, 241) a peroxide of La is ppd by adding BaO_2 to a neutral La salt, on drying it loses O Hermann (*J* pr 82, 397) says that a peroxide is obtained by heating oxalate, nitrate, or carbonate of La in the air, it dissolves in HClAq with evolution of Cl, heated in H it yields La_2O_3 Zschiesche (*J* pr 104, 74) was unable to confirm Hermann's observations Cleve (*Bl* [2] 43, 56) by ppg La salts by alkali and $\text{H}_2\text{O}_2\text{Aq}$ obtained an oxide to which he assigned the composition La_2O_5

Lanthanum, oxybromide of LaOBr Obtained by the action of Br vapour on heated La_2O_3 (Frerichs a Smith, *A* 191, 331)

Lanthanum, oxychlorides of LaOCl and $\text{La}_2\text{O}_2\text{Cl}_2$ ($= 3\text{La}_2\text{O}_3 \cdot \text{La}_2\text{Cl}_6$) The former is a greyish mass, unchanged by water, obtained by heating La_2O_3 in Cl to 200° (Frerichs a Smith, *A* 191, 331, Cleve, *Bl* [2] 89, 151, 43, 56) The latter is obtained by heating $\text{LaCl}_3 \cdot 7\text{H}_2\text{O}$ and washing the residue with water (Hermann, *J* pr. 82, 385)

Lanthanum, salts of. La forms one series of salts, LaX , where $\text{X} = \text{NO}_2, \frac{\text{SO}_4}{2}, \frac{\text{PO}_4}{3}$, &c Most

of the La salts are colourless, those which are soluble have a sweetish astringent taste The salts of volatilisable acids yield La_2O_3 when strongly heated La_2SSO_4 combines with the alkali sulphates, but the products are not alums A good many double La salts are known, a few basic salts have been isolated. The following are the chief La salts. *arsenate, arsenite, borate,*

bromate, carbonate, chlorate, chromate, hypochlorite, iodate, molybdate, nitrate, perchlorate, periodate, phosphates, phosphite, selenate, selenite, silicate, sulphates, sulphites, thiosulphate, tungstate, v CARBONATES, NITRATES, SULPHATES, &c

Lanthanum, sulphide of La_2S_3 , Red-yellow, microscopic crystals, obtained by the action of 8 parts Na polysulphide with 1 part La_2O_3 , and washing with water (Beringer, *A* 42, 134) Mosander obtained a greyish powder by heating La_2O_3 in CO_2 laden with CS_2 (*P* 60, 297), and by a similar reaction Frichs & Smith (*A* 191, 355) obtained La_2S_3 as a brownish grey powder, soluble in acid with evolution of H_2S , decomposed by water to LaO_2H_3 and H_2S Didier (*C R* 100, 1461) obtained a similar body by heating La_2O_3 in H_2S gas M M P M

LAN-HOPINE $\text{C}_{22}\text{H}_{23}\text{NO}$, [α 200°] A base homologous with papaverine, occurring in opium (Hesse, *A* 153, 57, *Suppl* 8, 271, *C C* 1870, 168)

Preparation—The aqueous extract of opium is pptd by Na CO_3 , the pp dissolved in ether, the ethereal solution shaken with dilute HOAc, and the acid solution poured into aqueous NaOH After 24 hours the ppt thebaine and papaverine are filtered off, the filtrate neutralised with HCl, pptd by NH_3 , and shaken with chloroform The chloroform takes up codeine, lanthopine, and meconidine, and is then shaken with dilute HOAc, and the solution exactly neutralised by NaOH, when lanthopine is pptd

Properties—White powder composed of minute prisms (from CHCl_3) Insol water, nearly insol alcohol, v sl sol ether and benzene, m sol chloroform When pptd from solutions by KOH or lime it dissolves in excess of the precipitant NH_3 gives a pp insol excess It does not give a blue colour with FeCl_3 Conc HNO_3 gives an orange red colour H_2SO_4 gives no colour in the cold, but a brownish yellow colour at 150°

Salts—The sulphate forms extremely thin needles— $\text{B}^+\text{HCl}6\text{aq}$ extremely thin needles, appearing like a jelly when in mass, v sl boiling water— $\text{B}^+\text{H}_2\text{PtCl}_62\text{aq}$ lemon yellow crystalline powder, insol water, alcohol, and HCl aq

LANUGINIC ACID $\text{C}_{41}\text{H}_{64}\text{O}_6$, H 73 p.c., N 163 p.c., S 34 p.c., O 314 p.c. An acid produced by the action of boiling baryta water upon wool (Champion, *C R* 72, 330, Knecht & Appleyard, *J* 22, 1120) The excess of baryta is removed by CO_2 , the acid pptd by lead acetate, and the pp decomposed by H_2S Yellowish porous mass, v sol hot water, sl sol alcohol, insol ether Its aqueous solution ppts colouring matters as lakes It also ppts tannin and most metallic oxides from their acetates It reacts like a proteid with Millon's reagent and with phosphotungstic acid

LAPACHIC ACID $\text{C}_{11}\text{H}_{14}\text{O}_5$, α $\text{C}_{11}\text{H}_{13}\text{O}_5(\text{OH})\text{CH}^+\text{Pr}$ *Oxy-amenyl naphthogumone* *Targuine* [138°]

Occurrence—A yellow colouring matter present in the 'lapacho' wood of a genus of the Bignonaceae, several species of which are indigenous to the Argentine Republic and other parts of South America (Siewert, *Report of Argentine Republic*, cap 15, *Philadelphia*). Occurs

also in Greenheart from Surinam (Stein, *J pr* 99, 8), and in Bethabarra wood (Greene & Hooker, *Am* 11, 267)

Preparation—The wood (10 kilos), in the form of saw dust, is boiled with a moderately-concentrated solution of sodium carbonate (500 grms crystallised salt in 80 litres water), the extraction is repeated several times The solution is of a blood red colour, and the addition of hydrochloric acid precipitates the crude acid together with a red brown resin, from which it is best purified by frequent solution and recrystallisations from benzene (Paternò, *G* 9, 505, 12, 337, Arnaudon, *C R* 46, 1152) Yield 5 p.c. of pure material

Properties—Monoclinic prisms, $\alpha b c = 7206 \cdot 1 \quad 6492$, $n = 97 \cdot 9$ (Panebianco, *G* 10, 80), v sol boiling alcohol, benzene, and ether It dissolves readily in solutions of the alkalis and alkaline earths, forming red solutions containing salts of the acid It decomposes carbonates on boiling

Reactions—1 Completely oxidised by *chromic acid*—2 Alkaline KMnO_4 gives oxalic acid—3 Boiling *nitric acid* (*S G* 138) yields phthalic acid—4 Distillation with *zinc-dust* yields isobutylene, naphthalene, and possibly a homologue of naphthalene—5 Boiling with HIAq and phosphorus forms amyl naphthalene (305°) and a little di (β)-naphthyl, both being perhaps derived from an intermediate naphthyl amylene 6 Cold conc HNO_3 (*S G* 149) or H_2SO_4 gives lapachone—7 *Zinc-dust* and *potash* reduce it to an unstable crystalline hydride, re-oxidised by air to lapachic acid

Salts— $\text{NaA}'5\text{aq}$ scarlet radio-crystalline mass S (of NaA') 151 at 24° Sol alcohol— $\text{KA}'\text{S}$ 333 at 24°— $\text{NH}_4\text{A}'\text{aq}$ large brick red crystals— $\text{CaA}'_2\text{aq}$ amorphous red pp S 224 at 24°— $\text{BaA}'_2\text{aq}$ long slender needles (from boiling water) S 23 at 26°— $\text{SrA}'_2\text{aq}$ — PbA'_2 , amorphous red pp, insol water— AgA' scarlet powder—Aniline salt $\text{C}_{11}\text{H}_9\text{NH}_2\text{A}'$ [123°], orange prisms (from alcohol)— α -Toluidine salt $\text{C}_{11}\text{H}_9\text{MeNH}_2\text{A}'$ [135°], yellow laminae— p -Toluidine salt [130°], orange yellow laminae

Acetyl derivative $\text{C}_{11}\text{H}_{12}\text{AcO}_5$ [83°] Formed, together with a compound $\text{C}_{11}\text{H}_{11}\text{O}_5$ (?), by heating lapachic acid with NaOAc and Ac_2O (Paternò) Sulphur yellow prisms, insol water, sol alcohol, ether, and benzene Readily converted by potash into lapachic acid Br in HOAc converts it into bromo lapachone HNO_3 (*S G* 148) at 0° forms a nitro-compound $\text{C}_{11}\text{H}_{12}(\text{NO}_2)\text{AcO}_5$ [170°], crystallising in reddish plates The compound $\text{C}_{11}\text{H}_{11}\text{O}_5$ forms small white acicular prisms [132°], v sl sol cold alcohol and ether Boiling alcoholic KOH does not saponify it, but converts it into a compound $\text{C}_{11}\text{H}_{12}\text{O}_5$, which crystallises (from alcohol) in orange needles [141°] Br does not convert the compound $\text{C}_{11}\text{H}_{11}\text{O}_5$ into bromo lapachone

Bromo-lapachone $\text{C}_{11}\text{H}_{11}\text{BrO}_5$ α

$\text{C}_{11}\text{H}_{12}\text{O}_5 < \frac{\text{CHBr}}{\text{O}} > \text{CHPr}$ (?) [140°] Formed by

warming lapachic acid with Br in HOAc (Paternò, *G* 12, 353) Orange laminae, v sol Lot alcohol, sol HOAc, and benzene, sl sol ether, insol cold aqueous alkalis HNO_3 oxidises it to phthalic acid.

Lapachone $C_{12}H_{10}O_4$, *see*

$C_{12}H_{10}O_4 < \begin{smallmatrix} OH \\ | \end{smallmatrix} > CHPr (?)$ [156°] Formed by the action of H_2SO_4 or cold conc HNO_3 (SG 149) on lapachic acid (Paternò, *G* 12, 837, Hooker & Greene, *B* 22, 1723) Silky orange red needles (from alcohol), insol water, v sol hot alcohol and benzene HNO_3 oxidises it to phthalic acid Distillation over red hot zinc dust gives naphthalene and isobutylene Ac_2O has no action Boiling with $NaOAc$ and Ac_2O forms a compound $C_{22}H_{22}O_4$, crystallising in brown tables with blue reflex, nearly insol Ac_2O $NaHSO_4$ forms a white crystalline compound Combines readily with hydroxylamine and with NH_3 Its molecular weight, found by Raoult's method, agrees with the formula $C_{12}H_{10}O_4$ Lapachone is converted by heating with alkalis into $C_{10}H_8O_4(OH)CH_2CH(OH)Pr$, which separates from cold alcohol or $HOAc$ in large crystals [125°], v sol most solvents, readily converted into lapachone by dilute HCl , and forming the salts $BaA''aq$ and $AgA''aq$, both crystallising in red needles

LARCH FUNGUS This fungus extracted with 95 p c alcohol yields a mixture of at least four different resins (Masing, *Ar Ph* [3] 6, 111)

(a) $C_{12}H_{12}O_4$ [125°] S (95 p c alcohol) 3 at 14° Insol chloroform

(b) $C_{12}H_{10}O$ [90°] S (95 p c alcohol) 7 Sol chloroform

(c) $C_{12}H_{10}O_4$ Agaric acid. [272°] Needles, v sl sol boiling alcohol, insol chloroform (Jahns, *Ar Ph* 221, 269)

(d) A portion easily soluble in alcohol, and partly ppd by water, leaving in solution a resin [C 61.9 p c, H 8.1 p c, O 30.0 p c] S 17, S (alcohol) 17 The resinous mixture is altered by boiling with milk of lime, the pp then thrown down by HCl being separated by chloroform into two substances $C_{12}H_{10}O_4$, insol chloroform, S (alcohol) 6, and $C_{12}H_{12}O_4$, sol chloroform, S (alcohol) 21

LARDACEIN v PROTEINS

LARD OIL A nearly colourless fatty oil, obtained by pressing hog's lard SG 20° 9122, d_4^{20} 1.4686 (Long, *Am* 10, 392)

LARICIC ACID $C_{16}H_{16}O_4$ [153°] S 1.14 at 60° Obtained from the small branches of larch trees (*Pinus larix*) from 20 to 30 years old (Stenhouse, *Pr* 11, 405) The bark is exhausted with water at 80°, and the extract evaporated and distilled The distillate is evaporated at 80° and the resulting crystals purified by sublimation Long monoclinic crystals resembling benzoic acid, usually in twins Sublimes at 93° Volatile with steam Has a bitter astringent taste and powerful odour Reddens litmus slightly M sol boiling water and alcohol, insol ether Nitric acid oxidises it to oxalic acid Its aqueous solution is ppd by baryta water, but not by lead or silver salts Ferric chloride gives a purple-red colour The K salt forms flat reddish brown crystals, decomposed by CO_2

LARREA RESIN Occurs on branches of *Larrea mexicana* (Stillmann, *B* 13, 756) 61.7 p c of it dissolves in alcohol, 26.3 in aqueous potash, and 1.4 p c consists of a colouring matter soluble in water

LASERPITIN $C_{22}H_{22}O_4$ (Feldmann, *A* 135, 886; *B* 1866, 1.457) or $C_{12}H_{22}O_4$ (Kütz, *Ar Ph*

[3] 21, 161) [114°] (F); [118°] (K) A bitter principle contained in the root of *Laserpitium latifolium* Extracted by 80 p c alcohol (F) or by light petroleum (K) Prisms Insol water, sol benzene, chloroform, ether, and CS_2 May be sublimed Its alcoholic solution is ppd by water, but not by alcoholic lead acetate (F) Insol dilute acids and alkalis Conc H_2SO_4 and conc $HClAq$ form deep red solutions (K) When heated with conc alcoholic KOH it is resolved into angelic acid and laserin $C_{11}H_{12}O_4$ (F) or $C_{22}H_{22}O_4$ (K), a crystalline resin, insol acids, sol ether, alcohol chloroform, and $HOAc$ Potash fusion gives methyl crotonic acid and laserin (K)

Laserpitin forms an acetate $C_{13}H_{22}O_4HOAc$ crystallising from acetic acid in silky needles (K)

Acetyl derivative $C_{13}H_{21}AcO_4$ [113°] From laserpitin, Ac_2O , and dry $NaOAc$ (K) Colourless needles, insol water, sol $HOAc$, alcohol, ether, and $CHCl_3$

Bromo derivative $C_{20}H_{20}Br_2O_4$ [90°] Formed by the action of Br on a solution of laserpitin in $CHCl_3$ (K) Needles, sol alcohol, ether, $CHCl_3$, and $HOAc$

Di-nitro- derivative $C_{13}H_{20}(NO_2)_2O_4aq$ [115°] From laserpitin and HNO_3 (K) Amorphous mass, insol water, sol alcohol, ether, $CHCl_3$, and $HOAc$

LAUDANINE $C_{20}H_{27}NO_3$ (Hesse, *A* 153, 57, 176, 201, *Suppl* 8, 272) [165°] SG 1.256 (Schröder, *B* 13, 1075) S (ether) 155 at 18° [α] = -13.5 in a 2 p c chloroform solution at 22.5°

Preparation—An aqueous extract of opium is ppd by lime or Na_2CO_3 , the filtrate is shaken with ether, the ethereal solution is shaken with dilute acetic acid, the acetic acid solution is neutralised exactly with NH_3 , the ppd lanthopine is removed by filtration, the filtrate is ppd by excess of ammonia, and the pp crystalised from ether Laudanine separates first from the ethereal solution, and afterwards codamine. The laudanine is dissolved in acetic acid solution, and the solution mixed with excess of $NaOH$, which ppt's cryptopine The alkaline filtrate is ppd by ammonium chloride, and the pp dissolved in aqueous $HOAc$, on adding KI laudanine hydro iodide is ppd, and this is decomposed by ammonia and the free base crystallised from ammonia

Properties—Stellate groups of small six-sided prisms (from alcohol) Cannot be sublimed Tasteless Lævorotatory In the crystalline state it is v sol benzene, $CHCl_3$, and boiling alcohol, sl sol cold alcohol, v sl sol ether In the amorphous state it is much more soluble Its salts have a bitter taste From their solutions caustic potash and ammonia ppt the base in white amorphous flocks, which soon become crystalline, and dissolve in excess of the precipitant Chloroform extracts the base from the ammoniacal but not from the potash solution Conc H_2SO_4 (containing ferric salt) gives an intense rose red solution, which at 150° changes to dark violet Conc HNO_3 gives an orange-red solution $FeCl_3$ gives an emerald-green colour The base is poisonous, its hydrochloride acting physiologically like strychnine.

Salts— B, H_2SO_4, aq concentric groups of

needles, v e sol water, almost insol dilute H_2SO_4 .—B'HC1 6aq prisms, v sol water and alcohol, v sl sol NaClAq Inactive to light—B'HB'r 2aq nodules S 35 at 20°—B'HI aq crystalline powder S 2 at 15° v sol boiling water, insol KIAq—B' H_2PtCl_6 2aq yellow amorphous pp, v sol boiling water—B' $H_2C_2O_4$ 6aq concentric groups of delicate needles [110°] S 22 at 10°—B' $C_4H_9O_3$ 3aq [100°] S 49 at 15°

LAUDANOSINE $C_{21}H_{33}NO_4$ [89°] S (ether) 52 at 16° $[\alpha]_D^{20} = 105$ in a 2 p c alcoholic solution at 22 5° $n_D^{20} = 56$ in a 2 p c chloroform solution at 22 5°

Preparation—Obtained from opium by ppg the aqueous extract with Na_2CO_3 , dissolving the pp in ether, shaking the ethereal solution with dilute HOAc, and pouring the acid solution into aqueous NaOH The pp contains a large number of bases, and is warmed with alcohol and dilute HOAc, and partially evaporated at 50° Papaverine and narcotine are then ppd, and thebaine and tartrate is ppd by adding tartaric acid to the filtrate The filtrate is exactly neutralised with NH_3 , and $NaHCO_3$ added After a week the pp is collected and extracted with benzene Cryptopine and protopine separate first from the benzene, and when the filtrate is shaken with $NaHCO_3$, laudanosome separates (Hesse, *A Suppl* 8, 321) It is purified by dissolving in acetic acid, ppg with KI, decomposing the resulting salt with NH_3 , and recrystallising from benzene

Properties—Needles More soluble in ether than thebaine and cryptopine, extremely sol alcohol and chloroform, v sol boiling benzene and ligroin, insol water and alkalis Conc H_2SO_4 (containing ferric salt) gives a brownish red solution, which at 150° becomes green, and finally dark greenish violet $FeCl_3$ gives no colour Tastes slightly bitter, its salts have an extremely bitter taste Its alcoholic solution exhibits a strong alkaline reaction Dextro rotatory Its hydrochloride is also dextro rotatory, $[\alpha]_D^{20} = 108$ in a 2 p c solution at 22 5°

Salts—B'HI aq small prisms, v sl sol cold water, v sol alcohol—B' H_2PtCl_6 3aq yellow amorphous pp, insol cold water—B' $H_2C_2O_4$ 3aq prisms, v e sol water

LAUREL OILS The essential oil from the leaves of the common cherry laurel (*Cerasus laurocerasus*) consists of benzoic aldehyde, HCy, a volatile oil (possibly benzyl alcohol) convertible by oxidation into benzoic acid, and minute quantities of an odorous resin (Elden, *Ph* [3] 5, 761, cf Leger, *Ph* [3] 3, 971) The essential oil from the fruits of *Laurus nobilis* contains a levorotatory terpene $C_{15}H_{26}$ (164°), SG 15 908, a levorotatory sesquiterpene $C_{15}H_{22}$ (250°), SG 15 925, and lauric acid $C_{12}H_{24}O_2$ (Blas, A 134, 1, Gladstone, *C J* 17, 1) The essential oil from the leaves of the Californian laurel (*Oreodaphne californica*) contains terpineol (168°) and umbellol $C_8H_{10}O$ (216°) (Stillmann, *B* 13, 629)

Laurel-nut oil is a fatty oil, SG 932, derived from *Calophyllum inophyllum* growing in the East (Hooper, *Ph* [3] 19, 525)

LAURENE v DI METHYL-ETHYL-BENZENE

LAURIC ACID $C_{12}H_{24}O_2$ Dodecic acid Mol. w. 200 [43 6°] (Heintz) (225°) at 100 mm.

(Krafft, *B* 13, 1415) SG 22 883 (Görgey). H C 1759730 (Lougumme, *A Ch* [6] 11, 222) Occurs as glyceryl ether in the berries of the bay tree (*Laurus nobilis*) (Marsson, *A Ch* 41, 33, Blas, A 134, 1), in the fat of pichurum beans (Schamer, *A* 53, 393), in the volatile oil of these beans (Müller, *J pr* 58, 469), in small quantity in spermaceti (Heintz, *A* 92, 394), in croton oil (Schlippe, *A* 105, 14), in the fruit of *Cylco daphne sebifera* (Gorkom, *Tydschrift af neerl Indie*, 81, 410), in the so called Dika bread, the fruit of *Mangifera gabonensis* (Oudemans, *J pr* 81, 356), in the age of the Mexicans, a fat obtained from *Coccus Azza* (Hoppe, *J pr* 80, 102), and in cocoa nut oil (Görgey, *A* 66, 303)

According to Scherring (*A* 96, 236) it is among the products of the distillation of cetyl alcohol with potash-lime, but Heintz (*A* 97, 271) denies this A dodecic acid [35°], isomeric with lauric acid, may be obtained by treating the amide of myristic acid with Br and NaOH, and converting the resulting dodecylamine into the nitrile and thence into the amide [97°] of dodecic acid, and boiling the amide with conc HClAq (Lutz, *B* 19, 1433)

Preparation—Lauric acid may be obtained from fats containing it by saponification followed by fractional precipitation of the acids by barium acetate (Heintz, *A* 92, 294, *P* 92, 429, 583, *J pr* 66, 1) It may also be obtained from the fat of bay berries by saponifying, distilling the solid fatty acids under reduced pressure, and rectifying *in vacuo* (Krafft, *B* 12, 1665)

Properties—Silky needles united in tufts (from alcohol) or scaly crystalline mass (after fusion) Its alcoholic solution has a slight acid reaction It is slightly volatile with steam Insol water, v sol alcohol and ether. Gives diennyl ketone ($C_{11}H_{22}O$) when its calcium salt is distilled

Salts—KA' Amorphous S (alcohol) 45 at 15°, 38 at 78° (Oudemans, *C C* 1863, 737)—KHA' Crystalline S (alcohol) 15 at 15°; 400 at 78°—NaA' White powder S (alcohol) 25 at 15°, 145 at 78°—NaHA' S (alcohol) 2 at 15°—NH₄HA' S (alcohol) 6 at 15°—BaA' crystalline spangles (from alcohol) S 07 at 100°, 0054 at 15° (Oudemans), 009 at 17°, 50 at 100° (Görgey) S (alcohol) 0187 at 15°, 1009 at 78° (O), 07 in the cold, 5 at 78° (G)—CaA' aq S 0039 at 15°, 0547 at 100° S (alcohol) 0719 at 15°, 220 at 78° (O)—SrA' aq S 0272 at 15°, 036 at 100° S (alcohol) 96 at 15°, 36 at 78°—MgA' 3aq S 023 at 15°—PbA' [110°–120°] (Heintz) S 0011 at 100° S (alcohol) 0047 at 15°, 235 at 100°—CoA' aq—NiA' aq—NiA' 3aq—CuA' S 0023 at 15°, 0029 at 100°—AgA' white powder, consisting of minute slender needles S 0001 at 15°, 0405 at 100° S (alcohol) 0323 at 15°, 0824 at 78°

Ethyl ether EtA' [–10°] (269°) at 750 mm (Delfs, *A* 92, 278) SG 22 86 (Görgey) V D 84 (calc 79) From the acid, alcohol, and HCl

Glyceryl ether $C_3H_7(C_{12}H_{25}O_2)_3$ (Schiff, *B* 7, 781) *Trilaurin Lauratearin* [45°] H. C. 5707420 (Lougumme, *A Ch* [6] 11, 222). Obtained from bay berries by extracting with alco-

hol Silky needles, sl sol cold alcohol, v sol ether

Phenyl ether $\text{A}'\text{C}_6\text{H}_5$, [24½°], (210° at 15 mm), pearly plates

p-Tolyl ether $\text{A}'\text{C}_6\text{H}_4$, [28°], (220° at 15 mm) (Krafft a Burger, B 17, 1878)

Amide $\text{C}_{11}\text{H}_{23}\text{CONH}_2$, [102°] (Krafft a Stauffer, B 15, 1729) From the chloride and NH_3

Chloride $\text{C}_{11}\text{H}_{23}\text{COCl}$ [-17°] (142½° at 15 mm) Colourless liquid (Krafft, B 17, 1878)

Nitrile $\text{C}_{11}\text{H}_{23}\text{CN}$ [4°] (198° at 100 mm) S G $\frac{1}{4} = 835$, $\frac{1}{2} = 827$, $\frac{100}{100} = 767$ Formed by distilling lauramide with P_2O_5 (Krafft a Stauffer, B 15, 1729) Colourless liquid of peculiar odour

LAURIC ALDEHYDE $\text{C}_{11}\text{H}_{23}\text{CHO}$ [45°] (143° at 22 mm) Prepared by the dry distillation of a mixture of calcium laurate and formate (Krafft, B 18, 1414) White crystalline odourless solid

LAURIN $\text{C}_{12}\text{H}_{24}\text{O}_2$ Bay berry camphor A substance discovered by Bonastre (*J Ph* 10, 32) in the berries of the bay tree, and further examined by Marsson (A 41, 329) and Delffs (A 88, 354) It is prepared by boiling the skinned and pounded berries with alcohol of 85 to 90 p.c., filtering at the boiling heat, and leaving the liquid to itself for several days Lauric stearin is then first deposited, and on filtering again and leaving the liquid to evaporate, laurin is deposited in crystals contaminated with a viscid oil, from which they may be freed by pressure between paper and recrystallisation Dimetric crystals, destitute of taste and smell, insol water, v sol alcohol, even in the cold, sol ether The solutions are neutral to test papers Laurin does not dissolve in alkalis Laurin can not be distilled without decomposition Its alcoholic solution is not precipitated by acetate of lead or nitrate of silver

LAUROCERASIN A name given by Lehmann (*N R P* 23, 440) to amygdalin when extracted from the leaves of the cherry laurel (*Cerasus laurocerasus*) or the bark of the berry-bearing alder (*Rhamnus Frangula*), v AMYGDALIN

LAURONE $\text{C}_{23}\text{H}_{46}\text{O}$ i.e. $(\text{C}_{11}\text{H}_{23})_2\text{CO}$ Di-*enyl ketone* [69°] S G, $\frac{1}{4} = 802$, $\frac{100}{100} = 788$ Formed by distilling barium laurate with lime Shining plates (from alcohol) On reduction it gives tricosane (Overbeck, P 86, 591, A 84, 289, Krafft, B 15, 1712)

LAURONOLIC ACID $\text{C}_6\text{H}_{11}\text{O}_2$ i.e. $\text{C}_6\text{H}_9\text{CO}_2\text{H}$ Formed by distilling camphanic acid, or by heating its barium salt with water at 200°, campholactone is formed at the same time (Woringer, A 227, 7)

Properties—Oil Sol water and ether Volatile with steam In presence of HCl some of it changes into the isomeric campholactone When distilled with water, a great part changes to the campholactone

Salts— CaA' , 3aq Forms dendritic crystals on the surface of an evaporating solution— AgA'

LAUROSTEARIN v *Glyceryl ether of LAURIC ACID*

LAUROXYLIC ACID v. DI METHYL BENZOIC ACID.

LAUTH'S VIOLET v AMIDO IMIDO IMIDO DI PHENYL SULPHIDE

LAVENDER OIL A volatile oil obtained by distilling the flowers of *Lavendula officinalis* with steam S G 12 875 It contains a levorotatory terpene $\text{C}_{10}\text{H}_{18}$ (162°) which forms a crystalline hydrochloride and various oxygenated bodies which yield camphor on further oxidation (Dumas, A Ch 13, 775, Lallemand, A 114, 198, Barth, Z 1867, 509, Bruylants, J Ph [4] 30, 39) English oil of lavender submitted to fractional distillation yields 30 p.c. of the terpene (176°-186°), and 30 p.c. of oxygenated products (200°-207°) (Shenstone, Ph [2] 13, 207)

LEAD Pb At w 206.4 Mc° w not known with certainty, but probably same as w (v. p 124) [330°-335°] (Brigel, B 6, 191, Pictet, P M [5] 7, 446), [326°] (Riemsdyk, C N 20, 32, Person, J 1849) (Between 140° and 1600°) (Carnelley a Williams, C J 35, 565) S G 11 335 at 0° (Quincke, P 97, 396), 11 37 at 0° (Reich, J pr 78, 328), 11 345 to 11 358 at 4°, 11 352 to 11 366 at 23° (Schweitzer, Am Ch 7, 174) S G molten 10 37-10 65 (Roberts a Wrightson, A Ch [5] 30, 181) For other values for S G v Clarke's *Table of Spec Gravities* [1888], pp 5-6 S H -78° to 11° 03065 (Regnault, A Ch [3] 26, 286), S H 19° to 48° 0315 (Kopp, Tr 1865 71), S H molten 340° to 450° 0402 (Person, A Ch [3] 24, 129) C E 0° to 100° 00002799 (Matthiessen, Pr 15, 220) T C (Ag=100) 8.5 (Wiedemann a Franz, P M [4] 7, 33) Heat of fusion 5858 (Rudberg, P 19, 125), 5369 (Person, A Ch [3] 24, 129) E C (Hg at 0°=1) 4.8 at 0°, 3.363 at 100° (Lorenz, W 13, 422, 582) Crystallises in regular octahedra For emission spectrum of lead v Werther, J pr 88, 180, Cornu, C R 73, 332, L de Boisbaudran, C R 77, 1152, Hartley a Adeney, Tr 1884 63

Occurrence—Lead occurs native in small quantities (e.g. v Chapman, P M [4] 31, 176, Koksharov, J M 1875 873, Igelstrom, J M 1889 (ii) 32) *Galena* (PbS) is very widely distributed, the other most important ores of lead are *cerussite* (carbonate), *anglesite* (sulphate), *pyromorphite* (phosphate), and *mimetesite* (arsenate) Small quantities of oxychloride, chromate, molybdate, tungstate, vanadate, &c., of Pb also occur Lead has been known and used from very early times

Formation—1 PbS is roasted in a reverberatory furnace until a portion is oxidised partly to PbO and partly to PbSO₄, the doors are then closed, and the PbS, PbO, and PbSO₄ react to produce SO₂ and Pb, PbS + 2PbO = 3Pb + SO₂, PbS + PbSO₄ = 2Pb + 2SO₂, -2 PbS is roasted in a reverberatory furnace, the temperature being gradually increased until a considerable quantity of PbSO₄ and a little PbO are produced, the doors are then closed, and the temperature is raised until the mass softens but does not melt When the PbS and PbSO₄ react to produce Pb and SO₂, the residue is again roasted, whereby more PbSO₄ is formed, and on raising the temperature with closed doors the PbSO₄ and PbS react as before After a time the quantity of PbSO₄ relatively to PbS becomes so great that the chief product of their reaction is PbO (PbS + 3PbSO₄ = 4PbO + 4SO₂), coal and wood are then thrown into the furnace, and heating is

continued, when the PbO is reduced to $\text{Pb}(\text{PbO} + \text{C} = \text{Pb} + \text{CO})$, at the same time some of the PbSO_4 is partly reduced to PbS , which, reacting with the remaining PbSO_4 , produces Pb and SO_2 ($2\text{PbSO}_4 + 2\text{C} = \text{PbSO}_4 + \text{PbS} + 2\text{CO}_2$) — 3 PbS is melted with scrap Fe , $\text{PbS} + \text{Fe} = \text{FeS} + \text{Pb}$ — 4 PbCO_3 is strongly heated with C , $\text{PbCO}_3 + 2\text{C} = \text{Pb} + 3\text{CO}$ — 5 PbO is reduced by heating in H or CO , or with C , KCN , or Na — 6 Zinc is suspended in an acidulated solution of a Pb salt, when Pb is ppt on the zinc — 7 PbCl_2 is mixed with Na_2CO_3 , and heated with KCN or C

Preparation — 1 By melting Pb prepared by one of the foregoing methods, exposing the molten mass to a gentle current of air, blowing water vapour through the mass, and running off the metal from beneath the film of oxides of Cu , Sb , Fe , &c., nearly pure Pb is obtained — 2 Pb oxalate is strongly heated in a carbon crucible, or is mixed with powdered C and heated — 3 Stas prepared pure lead by the following process (*Chem. Propriétés* 324) Commercial Pb acetate was dissolved in water, and digested at 40° to 50° in a leaden vessel with sheets of Pb until all Cu and Ag were ppt , the filtered liquid was run into almost boiling water, strongly acidulated with H_2SO_4 , the ppt PbSO_4 was very thoroughly washed, and then suspended in a solution of NH_4 sesquicarbonate and NH_3 until transformed into PbCO_3 , the PbCO_3 was thoroughly washed, a part of it was decomposed to PbO by heating in a Pt dish, and the rest was almost, but not quite, entirely dissolved in dilute HNO_3Aq , the solution of $\text{Pb}(\text{NO}_3)_2$ was heated to boiling, and the PbO was added little by little, the last traces of Fe were thus ppt as oxide, the boiling liquid was filtered and poured into solution of NH_4 sesquicarbonate. In this way pure PbCO_3 was obtained. The PbCO_3 was dried, and added little by little to pure molten KCN contained in an unglazed porcelain crucible placed within a larger crucible, the space between being filled with powdered Al_2O_3 , previously heated and mixed with 5 p.c. melted and powdered borax. The Pb thus obtained was again placed in pure molten KCN , and kept there until the upper surface of the Pb appeared convex and lustrous like pure Hg , after partial cooling the Pb was run off into a mould of polished steel. If a trace of PbO or PbS is present in the molten Pb the surface does not become convex.

Properties — Almost white, lustrous metal. Ordinary lead is blue gray, it contains traces of Ag , Cu , Sb , and sometimes Fe and Mn . Very malleable and ductile, but the properties are greatly affected by small quantities of impurities, notably by PbO , which is somewhat soluble in molten Pb . Tenacity very low. Somewhat sonorous. Leaves a mark on paper. Lead is so soft that it can be cut by a knife or scratched by the nail, traces of foreign metals, or of As or S , increase the hardness of Pb . By slowly cooling molten Pb , piercing a hole in the crust by an iron rod, and pouring off the still molten portion, the metal is obtained in regular octahedra often aggregated together similarly to crystals of NH_4Cl (Marx, *S* 57, 193, Stolba, *D P J*, 364, 371). Crystals of Pb are also obtained by hanging a rod of zinc in a slightly acidulated solution of a Pb salt, or by passing an electric current through

such a solution. Pb vaporises at high temperatures (1400° – 1600°), the vapour is very poisonous. The lustrous surface of Pb quickly tarnishes in ordinary air from formation of a film of oxide (probably Pb_2O). Very finely divided Pb is quickly changed to the suboxide Pb_2O , such finely divided Pb may be obtained by covering a Zn plate with a little PbSO_4 , made into a paste with water, laying another Zn plate on the top, and placing the whole in NaClAq for 9 or 10 days (*v* Bolley, *Ph C* 1850 59). Pb is readily changed to PbO on the surface by melting in air. Pb is not changed in dry air, nor when kept under pure water from which all air has been removed, in contact with water and the atmosphere it becomes gradually covered with a white deposit of 2PbCO_3 , $\text{PbO.H}_2\text{O}$, and small quantities of a soluble Pb salt are also formed (*v* *Reactions*, No 2). Pb is soluble in HNO_3Aq , and is slowly changed to PbCl_2 by the action of HClAq in the air, hot conc. H_2SO_4 forms PbSO_4 , Pb is oxidised by heating with KNO_3 or KHSO_4 . Combines with the halogens, also with S , Se , P , and As , forms alloys with many metals.

The atomic weight of Pb has been determined (1) by finding V D of PbCl_2 (Roscoe, *Pr* 27, 426), and analyses of this compound (Marignac, *Ar Sc* 1, 59, 209, Dumas, *A Ch* [3] 55, 196), (2) by syntheses of $\text{Pb}(\text{NO}_3)_2$ and PbSO_4 (Stas, *Rech* 101, *Chem. Propriétés* 329), (3) by converting Pb , PbO , and PbS into PbSO_4 , and by analysing PbCO_3 (Berzelius, *P* 8, 15, *G A* 37, 259, 265, Turner, *A* 13, 17), (4) by determining SH of Pb (Regnault, *A Ch* [3] 26, 286).

Lead is metallic in its chemical behaviour. The oxide PbO is strongly basic, it is slightly soluble in water, and the solution turns red litmus blue and absorbs CO_2 from the air, it decomposes hot solutions of NH_4 salts with evolution of NH_3 , and it reacts with acids to form salts PbX_2 ($\text{X} = \text{NO}_3$, $\frac{1}{2}\text{SO}_4$, &c.), PbCO_3 is isomorphous with the carbonates of the alkaline earths, on the other hand, PbO dissolves in fairly conc. KOH Aq , probably forming a salt PbO.K_2 , and compounds of PbO with CaO and Ag_2O are known. PbO_2 reacts with strong bases, e.g. KOH , to form unstable salts, e.g. K_2PbO_3 , on the other hand, treatment of Pb_2O_3 with conc. $\text{H}_3\text{PO}_4\text{Aq}$ or $\text{H}_2\text{C}_2\text{O}_4\text{Aq}$ produces solutions which react as if they contained salts of PbO_2 . The greater number of the lead salts correspond with the oxide PbO , $\text{Pb}(\text{CH}_3)_4$, however, is stable as a gas.

Lead is placed in Series II of Group IV, the other odd series members of this group are Si , Ge , and Sn . Pb more nearly resembles Sn than any other member of Group IV, it is more distinctly metallic than the other elements of the group, the other elements of this group form several compounds MX_4 , whereas most of the Pb compounds belong to the form MX_2 . Pb forms many basic and several double salts (*cf* CARBON GROUP OF ELEMENTS, vol 1 p 684, also TIN GROUP OF ELEMENTS, in vol iv). In its physical properties Pb resembles Tl , one series of Tl salts, however, is similar to those of the alkali metals, and the other series resembles those of the earth metals (*v* EARTH METALS OF THE, vol u p 424). The atom of Pb is divalent in the gaseous molecule PbCl_2 , and tetravalent in the gaseous molecule $\text{Pb}(\text{CH}_3)_4$.

Ramsay (*C J* 55, 521) has determined the lowering of vapour pressure of Hg produced by dissolving Pb in Hg, the results obtained make it probable that the molecular weight of lead is the same as the atomic weight, this result assumes the accuracy of Van't Hoff's law, that equal volumes of dilute solutions contain equal numbers of molecules of the dissolved substances, and it also regards the molecular weight of liquid Hg as the same as the atomic weight

Reactions—1 Unchanged in dry air, superficially oxidised (probably to Pb₂O) in moist air, oxidised to PbO by heating in air or oxygen—2 Water quite free from air has no action on Pb at ordinary temperatures (Stalman, *D P J* 180, 366, Bottger, *J* 1866 232, but *v* Muller, *J pr* [2] 36, 317) Water and air together dissolve a little Pb, and at the same time a deposit of 2PbCO₃, Pb(OH)₂ is formed Water charged with CO₂ under a pressure of a few atmospheres dissolves considerable quantities of Pb (perhaps in the form of an acid carbonate) The presence of small quantities of nitrates, especially NH₄NO₃, of soluble alkaline chlorides, and of some other salts, *eg* (NH₄)₂SO₄, increases the solvent action of water on Pb, the amount of Pb in solution is greater after a few days' action than after many days' exposure to the air In these reactions the soluble salt of Pb is probably slowly acted on by the CO₂ of the air with formation and ppn of insoluble 2PbCO₃, Pb(OH)₂ The presence of alkaline carbonates or of a little Ca silicate in water almost wholly stops the solvent action on Pb, probably the insoluble hydrocarbonate is formed as quickly as Pb is dissolved The action of water on lead has been examined by Graham, Miller, a Hofmann, Noad (*C J* 4, 20), Yorke (*P M* [3] 5, 82), Dumas (*C R* 77, 1054), Berthelot (*C R* 77, 1063), Pattison Muir (*C N* 25, 294, 33, 102, 125, 145, 34, 223, 234, 35, 82, 110, *C J* 31, 660), Muller (*J pr* [2] 36, 317), Carnelley a Frew (*S C I* 7, 15, 78), and others Granulated lead slowly decomposes boiling water, evolving H (Stolba, *J pr* 94, 113)—3 Lead is dissolved by several acids, slowly by hot conc HClAq in contact with air, also slowly by conc H₂SO₄ (Calvert a Johnson, *C J* 16, 66), rapidly by HNO₃Aq For account of gaseous products of reaction with HNO₃Aq *v* Ackworth a Armstrong, *C J* 82, 54.

Technical applications of lead—Lead is largely used for vessels and apparatus in which different chemical processes are conducted, *eg* for sulphuric acid chambers, water pipes are usually made of lead Alloys of lead with tin form solder and pewter, an alloy with As is used for making shot, an alloy with Sb is used as type-metal, and emery wheels and grinding tools used by lapidaries are made of an alloy of Pb and Sb, alloys of Pb with Sn and Bi have low melting-points Lead oxide, chromate, acetate, and carbonate are all largely used in manufactures

Detection and Estimation—HClAq ppts white PbCl₂ from solutions of Pb salts, sl sol cold water, fairly sol hot water, nearly insol dilute HClAq Dilute H₂SO₄, or a soluble sulphate, ppts white PbSO₄, almost insol dilute H₂SO₄Aq, quite insol alcohol. K₂CrO₄Aq ppts

yellow PbCrO₄, insol water KIAq ppts yellow PbI₂, sol boiling water, but reppd in yellow crystalline spangles on cooling H₂S ppts brownish black PbS, insol dilute acids, alkalis, or alkaline sulphides, presence of much HCl prevents ppn from rather dilute solutions of salts of lead, in presence of a little HCl pp is sometimes red or yellow red, and consists of xPbS yPbCl₂ (*v* Lead sulphochloride of) The H₂S test is said to detect 1 pt of lead in 100,000 pts of water, the H₂SO₄ test 1 pt in 20,000, and the K₂CrO₄ test 1 pt in 70,000 Pb may be estimated as Pb₂O, by ppn with dilute H₂SO₄Aq adding about 2 vols of alcohol, washing with alcohol, drying and calcining

Supposed allotropic form of lead When lead is deposited on the negative electrode by passing a current through a neutral or acid solution of a lead salt, and the current is continued for some hours, the lead becomes the colour of copper (Wohler, *A Suppl* 2, 135) After washing the red leaflets thus obtained with water and alcohol, they retain their colour on exposure to the air, and are not acted on by dilute HClAq or cold dilute HNO₃Aq, or alkalis, but are readily dissolved by hot HNO₃Aq, heated in H they melt above 200° and become ordinary lead Wöhler regarded the red substance either as an allotypic form of lead, or as a hydride of lead, Stolba (*J pr* 94, 113) suggested that the red colour was due to a film of oxide on the surface of the lead (*v* also Schutzenberger, *C R* 86, 1265)

Lead, alloys of Lead alloys with many metals, the alloys are produced by fusing together the constituent metals, some of them are definite compounds, in these cases either of the constituents is generally soluble in the compound

Alloys with bismuth, and with bismuth and other metals Pb and Bi may be alloyed in all proportions, malleability is diminished by adding more than an equal weight of Bi By fusing together 70 p c Pb, 15 p c Bi, and 15 p c Sb an alloy is obtained which expands on cooling An amalgam of Pb (4 pts) Bi (2 pts), and Hg (1 pt) may be melted by rubbing pieces of it together Alloys of Pb with Bi and Sn are known as fusible alloy, the MP varies from c 90° to c 170°, all these alloys are said to solidify at 98° (For MP and solidification points of these alloys *v* Dallo, *C C* 1865 831, for expansion and SH *v* Spring, *A Ch* [5] 7, 178, also Wiedemann, *W* 3, 237, for thermal conductivity *v* Wiedemann a Fierz, *P* 89, 514, 108, 399, *cf* also vol 1 p 511)

Britannia metal is composed of equal parts of brass, Sn, Sb, and Bi, Queen's metal of 1 pt Pb, 1 pt Bi, 1 pt Sb, and 9 pts Sn Alloys of Pb with Bi, Sn, and Ag are very fusible, melting as low as 45°, the alloy often used for tinning the inside of glass globes, tubes, &c is made by fusing together 1 pt Pb, 1 pt Sn, 2 pts Bi, and adding 10 pts Ag

Alloys with copper *v* vol u p 254

Alloys with mercury Pb easily amalgamates with Hg, by rubbing together Pb filings with Hg or by adding Hg to molten Pb An amalgam containing 33 p c Pb is liquid, an amalgam of equal parts Pb and Hg can be crystallised Contraction occurs during the amalgamation

Alloys with potassium and sodium Prepared either by directly heating the metals together or by fusing PbO with an alkaline flux, *e.g.* with cream of tartar. When distilled with EtI, MeI, &c., compounds of Pb with Et, Me, &c. are obtained.

Alloys with tin These metals may be alloyed in all proportions, the S.G. of the alloy is always a little less than that calculated from the S.G. of the constituents. There are no indications of the formation of definite compounds (*v.* Laurie, *C J* 55, 677). These alloys are very easily ignited and burnt. The two commonest alloys of Pb and Sn are *solder* and *pewter*. *Fine solder* contains 1 pt Pb and 2 pts Sn, *common solder*, equal pts Pb and Sn, *coarse solder*, 2 pts Pb and 1 pt Sn. The M.P. varies from c 340° for 1 pt Pb and 2 pts Sn, to 370° for equ 1 parts Pb and Sn, and 440° for 2 pts Pb and 1 pt Sn, by increasing the relative quantity of Pb the M.P. rises until the alloy of 25 pts Pb to 1 pt Sn melts at c 560°, by increasing the relative quantity of Sn, above 2 pts to 1 of Pb, the M.P. also slightly rises until the alloy of 6 pts Sn to 1 pt Pb melts at c 380°. *Pewter* is an alloy of c 80 pts Pb with 20 pts Sn. According to Rudberg (*P* 18, 240) when Pb and Sn are melted together and allowed to cool, the thermometer always shows a stationary point at 187°, he thinks that a compound PbSn₂ is always formed, that this alloy has a fixed solidification point lower than that of Pb (326°) or Sn (228°), and that the heat produced by the solidification of this alloy acting on the excess of Pb or Sn causes fluctuations in the final solidification point of the whole mixture (*v.* also Pohl, *W A B* 1850 402, Riche, *C R* 55, 143, Wertheim, *P Ergänzzbd*, 2, 75, Matthiessen, *P* 130, 62). It seems that the amount of Pb in *pewter* vessels for domestic use should not exceed 18 p.c., else Pb may be dissolved out by the action of dilute acids, *e.g.* vinegar.

Alloys with tin and copper *Bell metal* is an alloy of 43 p.c. Pb, 80 p.c. Cu, 101 p.c. Sn, and 56 p.c. Zn. A little Pb is sometimes added to Cu and Sn in making *bronze*.

Alloys with palladium A crystalline, greyish white, brittle alloy is obtained by fusing granulated Pb with rather more than its own weight of Pd foil, and removing excess of Pb by the action of acetic acid and CO₂. The alloy has the composition Pd₂Pb, S.G. 11.225 (Bauer, *B* 3, 691, 4, 419).

Alloys with platinum An alloy having the composition PtPb is formed by fusing 3 pts Pb with 1 pt Pt, and exposing the product to the prolonged action of CO₂, O, and acetic acid vapour (Bauer, *B* 3, 691, 4, 449). Steel-grey, crystalline, fairly fusible, S.G. 15.736. Bauer (*l.c.*) also describes an alloy PtPb₂.

Alloys with zinc Pb and Zn alloy in all proportions, all the Zn is removed by heating very strongly. Alloying Pb with Zn increases the hardness of the Pb. Alloys of Pb, Zn, and Sn solidify at 168°, when the ratio is 2Pb Zn 9Sn the alloy solidifies at 168° only, but other alloys show also two higher points.

Lead also alloys with *chromium* and *manganese*.

Lead, antimonate of, *v.* vol 1 p 286

Lead, antimonides of Pb and Sb may be melted together in all proportions, the alloys are crystalline, most of them are formed with expansion. Ordinary *type metal* consists of 83 pts Pb and 17 pts Sb (*v.* Riche, *C R* 55, 143, Matthiessen, *P* 110, 28, Calvert a Johnson, *P M* [4] 18, 354, H v d Planitz, *B* 7, 1664, Rollmann, *P* 84, 277).

Lead, arsenates of, *v.* vol 1 p 308

Lead, arsenides of Pb melted with As takes up c 16 p.c. of the latter, the alloys are brittle and very fusible, some, but not all, of the As is expelled by heating. *Shot metal* is an alloy of Pb with not more than 3 p.c. As.

Lead, arsenites of, *v.* vol 1 p 306

Lead, borates of, *v.* vol 1 p 530

Lead, borofluoride of Pb(BF₄)₂ [= PbF₂, 2BF₃] Prismatic crystals, obtained by dissolving PbO in excess of HBF₄ and evaporating to a syrup. Crystallises with difficulty, partially decomposed by water or alcohol (Berzelius, *P* 2, 113, cf. Borofluorides, vol 1 p 526).

Lead, bromide of PbBr₂. Mol w not certain, but from analogy of PbCl₂ is probably 365.9 (= PbBr). Melts at c 499° (Carnelley, *C J* 33, 278), boils at c 861° (Carnelley a Williams, *C J* 33, 283). S.G. ppd PbBr₂ 6.572 at 19.2° (Clarke's *Table of Specific Gravities* [1888] 32). H.F. [Pb₂Br]₂ = 61,450, [PbBr₂Aq] = 54,410 (*Th*. 3, 337). Obtained by digesting PbO or PbCO₃ with HBrAq, or by ppg the solution of a Pb salt by HBrAq or a soluble bromide. Obtained in well formed, colourless, rhombic needles by slowly evaporating a solution of Pb in alcohol containing Br (Hjortdahl, *Z K* 3, 302). Sol. cold, more sol. hot, water, more sol. acids. From the solution in hot conc. HBrAq (72 p.c. HBr), Ditté says that white lustrous needles, PbBr₂·3H₂O, crystallise (*C R* 92, 718), and that the compound 5PbBr₂·2HBr·10H₂O is obtained by passing HBr into the solution of PbBr₂ in HBrAq. After melting, PbBr₂ solidifies to a yellow, horn like mass. Heated in air, a yellow mass remains containing the *oxybromide* PbOBr₂ (= PbBr₂·PbO), the same oxybromide is obtained by digesting PbBr₂ with Pb(C₂H₃O₂)₂Aq. A compound of PbBr₂ with PbCl₂ has been obtained, also compounds of PbBr₂ with PbI₂, and with Pb(CNS)₂. (*v.* **Lead, bromochloride of**, **Lead, iodobromides of**, and **Lead, bromosulphocyanides of**). André (*A Ch* [6] 3, 104) describes several *double compounds* of PbBr₂ with NH₄Br.

Lead, brom iodides of, *v.* **Lead, iodobromides of**.

Lead, bromochloride of PbBr₂PbCl₂ (= PbBrCl). White needles, observed by Illes in a blast furnace where lead carbonate ores had been reduced (*C N* 43, 216), crystallisable from water, S.G. 5.741.

Lead, bromosulphocyanides of PbBr₂Pb(CNS)₂, brownish crystals, by digesting the constituents together, and then crystallising from boiling water. By acidifying with HBrAq, the salt 8PbBr₂Pb(CNS)₂ is produced (Grisson a Thorp, *Am* 10, 229).

Lead, chlorides of Only one chloride of lead, PbCl₂, has been isolated, but there is strong proof of the existence of the tetrachloride, PbCl₄, in solutions of PbO, in HClAq.

Lead chloride, PbCl₂. (**Lead dichloride**

Plumbous chloride *Horn-lead*) Mol w 277.14 S G 5.805 at 15° (Stolba, *J pr* 97, 503), 5.88 (Brigellmann, *B* 17, 2359) V D 187.1 at c 1080° (mean of 4) (Roscoe, *Pr* 27, 426) Melts at 498°, and boils between 861° and 954° (Carnelley, *C J* 33, 278, and C A Williams, *C J* 35, 564) Crystallises in rhombic system, $a b c = 5943.1 \ 5949$ (Schabus, *W A B* 1850 456) H F [Pb, Cl] = 82,770, [Pb, Cl², Aq] = 75,970 (*Th* 3, 337)

Occurrence—As *cotumite*, found in the crater of Vesuvius after the eruption of 1822, mixed with NaCl and CuSO₄.

Formation—1 By heating Pb in a stream of Cl, combination occurs slowly—2 By dissolving Pb in hot HClAq in contact with air, PbCl₂ separates on cooling—3 By adding solution of a chloride, e.g. NaClAq, to a conc solution of a Pb salt—4 By dissolving PbS in hot HClAq and crystallising

Preparation—PbO or PbCO₃ is boiled with dilute HClAq, and HClAq is added little by little until all is dissolved, the liquid is allowed to cool, and the PbCl₂ which separates is repeatedly crystallised from boiling water

Properties—Large, white, rhombic crystals (*v supra*), which melt at 498° and boil between 861° and 954° (*v supra*) V D at c 1080° agrees with the formula PbCl₂. Sol. sol cold water, S = 95 at 16.5°, solution contains 9414 p.c. PbCl₂ (Bell, *C N* 16, 69), addition of 1 p.c. HCl diminishes S to 347, and with 14 p.c. HCl, S = 09. If more than 14 p.c. HCl is present S increases, S in HClAq S G 1.162 = 2.9 (Bell, *l c*) According to Field (*C J* [2] 11, 575) PbCl₂ is less soluble in NaClAq containing 5 p.c. NaCl than in water (S = 23), but the values of S for water and conc. NaClAq are about the same. The considerably greater solubility of PbCl₂ in conc. than in dilute HClAq is supposed by Ditte to be due to the formation of a compound of PbCl₂ and HCl which is decomposed by water (*C R* 92, 718) Ditte gives the following table—

HCl in 100 pts H ₂ O	PbCl ₂ dissolved in 1000 pts of the liquid in vol 1				
	at 0°	20°	40°	55°	80°
0	8	11.8	17	21	31
5.6	2.8	3	4.6	6.5	12.4
10	1.2	1.4	3.2	5.5	12
18	2.4	4.8	7.2	9.8	19.8
21.9	4.7	6.2	10.4	12.9	23.8
31.5	11.9	14.1	19	24	38
46	29.8	30 (at 17°)	—	—	—

PbCl₂ is readily dissolved by boiling water, but almost all crystallises out on cooling. Insol. alcohol 94 p.c., somewhat sol. in more dilute. Fairly sol. NaC₂H₃O₂Aq, Na₂S₂O₃Aq, and some other salt solutions. After melting and cooling, PbCl₂ appears as a horn-like mass (hence the name *horn-lead* formerly used)

Reactions—1 Heated in the air until white fumes cease to come off, a light yellow oxychloride Pb₂OCl₂ (= PbCl₂ PbO) is formed (*v Lead, oxychlorides of*)—2 Recently pptd PbCl₂ when digested with cold neutral lead acetate produces an oxychloride 2Pb₂OCl₂H₂O (Brandes, *A* 10, 273)—3. The oxychloride

Pb₂OCl₂H₂O is obtained by partially pptg PbCl₂Aq by lime water—4 By adding caustic potash to PbCl₂ suspended in water, the oxychloride PbCl₂ 2PbO is said to be formed (Ditte, *C R* 94, 1180)—5 According to Berzelius, the product of the action of ammonia on PbCl₂ is PbCl₂ 3PbO 4H₂O—6 Heated in carbon monoxide, COCl₂ and Pb are produced—7 Heated, not too strongly, in phosphoretted hydrogen, HCl, P, and Pb are formed—8 Oxidised by alkaline hypochlorites to PbO₂—9 Long continued passage of Cl into PbCl₂ suspended in water most probably produces PbCl₂ PbO₂ being simultaneously pptd (*v Fisher, C J* 35, 284)

Combinations—1 PbCl₂ absorbs ammonia gas, forming 2PbCl₂ 3NH₃, according to H. Rose. All the NH₃ is readily given off on warming—2 Combines with lead monoxide, when the two are heated together, to form oxychlorides xPbCl₂ yPbO (*v Lead, oxychlorides of*)—4 With lead carbonate forms PbCl₂ PbCO₃, obtained by boiling together the two salts, or by the action of CO₂ on PbCl₂ under pressure (Miller, *C J* [2] 8, 37)—5 With lead acetate to form PbCl₂ Pb(C₂H₃O₂)₂, obtained by mixing freshly pptd PbCl₂ with Pb(C₂H₃O₂)₂ and enough glacial acetic acid to form a viscid mass, which soon solidifies, and removing excess of acid by pressure (Carius, *A* 127, 87)—6 With lead phosphate, forms PbCl₂ Pb₃(PO₄)₂ H₂O and PbCl₂ 2Pb₃(PO₄)₂, or according to Gerhardt (*A Ch* [3] 22, 505) PbCl₂ Pb₂H(PO₄)₂. The former is produced by pouring boiling PbCl₂Aq into excess of boiling Na₂HPO₄Aq, the latter by pouring the Na phosphate into the PbCl₂ solution (Heintz, *P* 73, 122) *Pyromorphite* PbCl₂ 3Pb₃(PO₄)₂ occurs native—7 Combines with lead bromide to form PbCl₂ PbBr₂, with lead iodide to form PbCl₂ PbI₂, with lead fluoride to form PbCl₂ PbF₂, and with lead sulphocyanide to form PbCl₂ Pb(CNS)₂ (*v Lead bromochloride, Lead iodobromides, Lead chlorofluoride, and Lead chlorosulphocyanide*)—8 Probably combines with chlorine to form PbCl₄, when Cl is passed into PbCl₂ dissolved in considerable excess of HClAq, or when Cl is passed into PbCl₂Aq mixed with CaCl₂ (*v Lead tetrachloride*)—9 André (*A Ch* [6] 3, 104) describes various double compounds of PbCl₂ with NH₄Cl

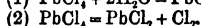
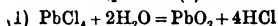
LEAD TETRACHLORIDE (*Plumbic chloride*) No other chloride of lead except PbCl₂ has been isolated, but there is considerable evidence in favour of the existence in HCl solution of tetrachloride PbCl₄.

If a considerable quantity of HClAq is added to PbCl₂, and Cl is passed in, the PbCl₂ dissolves to a reddish brown liquid, on warming Cl is evolved, addition of water causes pptn of PbO₂.

A mixture of PbCl₂Aq with CaCl₂Aq absorbs much Cl and forms a deep yellow liquid, which may be preserved unchanged in a closed vessel, but is decomposed in an open vessel giving PbCl₂ and Cl, a few drops of this liquid added to a large quantity of water gives a ppt of PbCl₂ and PbO₂, alkalis and alkaline carbonates ppt PbO₂, MnCl₂Aq ppts MnO₂ and PbCl₂, Fe, Cu, Zn, gold leaf, and finely divided Pt, are soluble in the liquid with separation of PbCl₂, it rapidly oxidises many organic bodies, PbCl₂ being pptd.

(Sobrero & Salmi, *A Ch* [8] 29, 165, Nicklès, *A Ch* [4] 10, 323)

PbO₂ dissolves in cold fairly conc HClAq. From this yellow solution, alkalis, alkaline carbonates, oxides and carbonates of Ba, Mg, Zn, &c., and alkaline borates and acetates, ppt PbO₂. The solution smells of Cl and is decomposed by heat with evolution of Cl and ppt of PbCl₂. If the HClAq is saturated with PbO₂, the addition of water ppts PbO₂. Fisher (*C J* 35, 282) determined the ratio of Pb pptd from this solution by water as PbO₂ to Cl given off by decomposing the solution with ppt of PbCl₂. The ratio was almost exactly Pb 2Cl, now this is the ratio required on the hypothesis that the decomposition by water proceeds as shown in equation (1), and the decomposition to PbCl₂ and Cl proceeds as shown in equation (2)—



Fisher proved that the whole of the Pb in solution is thrown down by water as PbO₂. A yellow liquid with properties the same as those of the ratio just described is obtained by treating Pb₂O₃ with a considerable quantity of HClAq (Fisher, *l c*)

By the method described above, Sobrero & Salmi (*l c*) obtained a double salt PbCl₂.9NaCl, and Nicklès (*l c*) obtained PbCl₂.16CaCl₂. Evaporation *in vacuo*, over conc KOHAq, of a solution of PbO₂ in HClAq produces crystals of PbCl₂, containing some PbCl₄, according to Bendant & Daguin (*Ann M* [5] 4, 239, *v* also Nikolukine, *J R* 1885 207, abstract in *C J* 50, 123)

Lead, chlorobromide of, *v* Lead, bromo-*hydrides of*

Lead, chlorocarbonate of, *v* Lead chloride, *Combinations No 4*

Lead, chlorocyanide of, *v* Lead cyano-*chloride of*

Lead, chlorofluoride of PbCl₂PbF₂ (=PbFCl). Obtained by ppg boiling PbCl₂Aq by KFAq, or Pb(C₂H₃O₂)₂Aq by a solution of 2 parts NaF with 3 parts NaCl (Berzelius). A white powder, slightly soluble in water without decomposition

Lead, chloronitrides of, *v* Lead, iodo-*chlorides of*

Lead, chlorophosphate of, *v* Lead chloride, *Combinations No 6*

Lead, chlorosulphide of, *v* Lead, sulpho-*chloride of*

Lead, c'lorosulphocyanide of, PbCl₂Pb(CNS)₂, by digesting the constituents together, and crystallising from boiling water, slowly changed to PbO₂H₂ by NH₃Aq (Grisson & Thorp, *Am* 10, 229)

Lead, chromates of, *v* vol II pp 155 and 157

Lead, cyanides of. None has been isolated, but an *oxycyanide*, and also *ferri-* and *ferro-* cyanides, are known, *v* vol II pp 341, 339, and 335

Lead, cyanochloride of PbCl₂.2PbCy₂, by digesting PbCl₂ with KCyAq (Grisson & Thorp, *Am* 10, 229)

Lead, ferricyanide of, *v* vol II p 349

Lead, ferrocyanide of, *v* vol II p 335

Lead, fluochloride of, *v* Lead, chloro-*fluoride of*

Lead, fluoride of PbF₂. Pb is not acted on by HFAq, PbO₂H₂, or PbCO₃, dissolves in HFAq, and on evaporating, and heating to remove HF, PbF₂ remains. PbF₂ is also pptd on adding Pb(C₂H₃O₂)₂Aq to HFAq. A white powder, slightly soluble in water, sol in HNO₃Aq (Berzelius). Gunz (*A Ch* [6] 3, 5) gives H F of PbF₂ as (PbO·H₂·2HF) = 24,300

Lead, hydroxides of, *v* Lead, oxides and *hydrated oxides of*

Lead, iodide of PbI₂. Mol w not certainly known, but from analogy of PbCl₂, probably 459.46 (=PbI₂). S.G. 6.12, 5.6247 molten at 383° (Rodwell, *T* 1882 1144). Melts at 383°, and boils between 861° and 954° (Carnelley, *C J* 33, 277, C a Williams, *C J* 37, 126). For expansion of PbI₂, *v* Rodwell, *Pr* 32, 23. H F [Pb, I] = 39,800 (*Th* 3, 337)

Preparation—Solution of Pb(NO₃)₂ is added to solution of FeI₂, the pp is washed with cold water and dried. KI or CaI₂ may be used instead of FeI₂. If Pb(C₂H₃O₂)₂Aq is added to KIAq oxyiodides are formed when excess of Pb salt is added, and soluble double iodides are formed if excess of KI is present, to prevent both results the solution of Pb(C₂H₃O₂)₂ should be acidified either with HNO₃ or C₂H₃O₂.

Properties and Reactions—Golden yellow crystals, hexagonal, *a c* = 1.3018 (Norden-skjöld). Melts when heated to a brown red liquid, and solidifies to a yellow mass. Heated in the air I is evolved, and oxyiodides (*q v*) are formed. S cold water 03, boiling water 515. Decomposed by Cl. Zn or Fe boiled with PbI₂ under water forms ZnI₂ or FeI₂, and ppts Pb. Na₂S₂O₃ in excess forms Pb sulphite and NaI (Michaelis & Koethe, *B* 6, 999). NH₃Aq added to boiling PbI₂Aq ppts 3PbO.PbI₂.2H₂O. Dry PbI₂ is unchanged by light, when moist and exposed to direct sunshine, in contact with the air, it is slowly changed to PbCO₃ and PbO, with separation of I, this change is hastened by all substances which absorb I (Schmid, *P* 127, 493). PbI₂ is somewhat soluble in conc solution of alkali iodides, but is pptd on dilution.

Combinations—1 Absorbs ammonia, forming a white compound PbI₂.2NH₃.—2 With lead monoxide forms various oxyiodides xPbI₂.yPbO (*v* Lead, oxyiodides of).—3 By dissolving PbI₂ in warm iodohydric acid solution, and allowing to cool, white lustrous needles of PbI₂.2HI = H₂PbI₄ separate. This compound may be called *iodo plumbhydric acid* or *iodo plumbic acid*, it is decomposed by water, gives off HI *in vacuo*, according to Berthelot, the crystals contain 10H₂O (*C R* 91, 1024).—4 PbI₂ combines with potassium iodide to form various double salts, one of which, PbI₂.2KI, seems best regarded as the K salt of H₂PbI₄. Yellow, lustrous leaflets of PbI₂.KI are obtained by dissolving the constituent salts in the ratio PbI₂.KI in boiling water, and allowing to cool, by dissolving this salt in warm KIAq and cooling, the salt PbI₂.4KI is said to be obtained in white needles (Boullay, *A Ch* [2] 34, 366). Remsen (*Am* 11, No 5) asserts that the process described by Boullay does not result in production of PbI₂.4KI, and that the salt formed with most ease when PbI₂ and KI react under different conditions is PbI₂.KI.H₂O (*cf* Ditte, *C R* 92, 1841).—5 By mixing a boiling solution of KI

and NH_4Cl with $\text{Pb}(\text{C}_2\text{H}_3\text{O}_2)_2\text{Aq}$, taking care that no permanent pp was formed, and cooling, Völkel obtained clear yellow lustrous needles of $\text{PbI}_2 \cdot 2\text{NH}_4\text{Cl}$ (P 62, 252), decomposed by water with separation of PbI_2 . By saturating hot NH_4ClAq with PbI_2 , cooling, pouring off from $\text{PbI}_2 \cdot 2\text{PbCl}_2$, which separates, and evaporating, Poggiale (C R 20, 1180) obtained $\text{PbI}_2 \cdot 4\text{NH}_4\text{Cl} \cdot 2\text{H}_2\text{O}$, decomposed by water with separation of PbI_2 . — 6 PbI_2 combines with *silver iodide*. For properties of the PbI_2AgI produced v Rodwell (Pr 32, 540), Bellati a Romanese (Pr 34, 104). — 7 Compounds with *lead bromide* are formed by crystallising PbI_2 from HBrAq (v *Lead, iodobromides of*) — 8 Compounds with *lead chloride*, $\text{PbI}_2 \cdot \text{PbCl}_2 (= \text{PbClI})$ and $\text{PbI}_2 \cdot 2\text{PbCl}_2$, are known (v *Lead, iodochlorides of*) — 9 By boiling PbCO_3 repeatedly with PbI_2Aq , Poggiale (C R 20, 1184) obtained $\text{PbI}_2\text{PbCO}_3$. — 10 $\text{PbI}_2 \cdot \text{Pb}(\text{C}_2\text{H}_3\text{O}_2)_2$ was obtained by Carus (A 127, 87) similarly to the PbCl_2 compound (v *Lead chloride, Combinations No 5*)

Lead, iodobromides of By crystallising PbI_2Aq from HBrAq , Grisson a Thorp obtained the compounds (1) $\text{PbBr} \cdot \text{PbI}_2$, (2) $3\text{PbBr} \cdot \text{PbI}_2$, (3) $6\text{PbBr} \cdot \text{PbI}_2$ (Am 10, 229)

Lead, iodicarbonate of, v *Lead, iodide of, Combinations No 9*

Lead, iodochlorides of The compound $\text{PbClI} (= \text{PbCl}_2 \cdot \text{PbI}_2)$ is obtained, in sulphur yellow prisms, by cooling a solution of PbI_2 in boiling HClAq (Laboure, J Ph [3] 4, 328) According to Engelhardt the compound contains more PbI_2 than required by the above formula (J pr 67, 293) By saturating hot NH_4ClAq with PbI_2 , and allowing to cool, Poggiale obtained yellow needles of $\text{PbI}_2 \cdot 2\text{PbCl}_2$ (C R 20, 1180)

Lead, iodosulphocyanide of, $\text{PbI}_2 \cdot 3\text{Pb}(\text{CNS})_2$, formed with some difficulty by digesting the constituents and crystallising from hot water (Grisson a Thorp, Am 10, 229)

Lead, oxides and hydroxides (or hydrated oxides) of Five oxides of lead have been isolated, viz, Pb_2O , PbO , Pb_2O_3 , Pb_3O_4 , and PbO_2 , one or more oxides intermediate between Pb_2O_3 and PbO_2 perhaps exist Two hydrates of PbO , viz $2\text{PbO} \cdot \text{H}_2\text{O}$ and $3\text{PbO} \cdot \text{H}_2\text{O}$ are known, the hydrates $\text{Pb}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$ and $\text{PbO}_2 \cdot \text{H}_2\text{O}$ have also been isolated The most stable oxide is PbO , Pb_2O is easily oxidised to PbO by heating in air, and Pb_2O_3 , Pb_3O_4 , and PbO_2 are reduced to PbO when strongly heated PbO reacts with acids as a basic oxide, forming salts PbX_2 , $\text{X} = \text{NO}_3$, Cl , $\frac{1}{2}\text{SO}_4$, $\frac{1}{2}\text{PO}_4$, &c., it decomposes NH_3 salts, evolving NH_3 , and combines with CO_2 to form PbCO_3 , PbO also reacts with strongly basic oxides as an acidic oxide forming unstable *plumbites*, e g K_2PbO_2 , PbO reacts as an acidic peroxide, with KOH and NaOH it forms fairly stable *plumbates*, M^+PbO_3 , with most acids it forms salts of PbO , but it appears capable of dissolving without reduction in glacial acetic and phosphoric acids The reactions of Pb_2O_3 and Pb_3O_4 indicate that these oxides are compounds of the form $x\text{PbO} \cdot y\text{PbO}_2$ As none of the oxides of Pb has been gasified the molecular weight of none is known with certainty

LEAD SUBOXIDE, Pb_2O The grey pellicle which forms on the surface of lead exposed to the air consists of Pb_2O , according to Berzelius

Pb_2O is prepared by heating PbC_2O_4 in a retort to near 800° as long as any gas is given off, $2\text{PbC}_2\text{O}_4 = \text{Pb}_2\text{O} + \text{CO} + 3\text{CO}_2$, the residue is allowed to cool in the retort (Dulong, S 17, 229, Pelouze, J pr 25, 486) Pb_2O is a black powder, SG 9 772 (Playfair a Joule, C S Mem 3, 83), heated in air it burns to PbO , heated out of contact with air, PbO and Pb are produced Moist Pb_2O rapidly absorbs O from the air, forming white $\text{PbO} \cdot x\text{H}_2\text{O}$ Dilute acids resolve Pb_2O into PbO , which dissolves, and Pb , saturated $\text{Pb}_2\text{NO}_3\text{Aq}$ produces a similar resolution, but Pb_2O is wholly soluble in dilute $\text{Pb}_2\text{NO}_3\text{Aq}$ This oxide was at one time supposed to be a mixture of PbO and Pb , but this is disproved by the facts that when the oxide is rubbed with Hg either dry or under water, no Pb is removed, and that PbO is not removed from it by boiling with Pb acetate solution

LEAD PROTOXIDE PbO (*Plumbous oxide* *Litharge* *Massicot*) Mol w unknown, as oxide has not been gasified SG 9 277 at 17.8° (Herapath, P M 64, 321), 9 25 (Playfair a Joule, C S Mem 3, 84), 9 17 to 9 88 (Ditte, C R 94, 1310), 8 74 to 9 29 (Geuther, A 219, 60) H F $[\text{PbO}] = 50,300$, heats of neutralisation $[\text{PbO} \cdot 2\text{HClAq}] = 56,830$, $[\text{PbO} \cdot \text{HNO}_3\text{Aq}] = 24,250$ to form basic nitrate (Th 3, 337) Crystallises in rhombic forms $a \cdot b \cdot c = 666 \cdot 1 \cdot 971$ (Nordenskjöld, P 114, 619, v also Mitscherlich, P 49, 403, J pr 19, 451, Grailich, W A B 28, 282, Marx, J pr 3, 217) PbO is dimorphous according to Ditte (C R 94, 1310)

Occurrence — As *lead ochre*, in small quantities in Mexico, Baden, Virginia, and a few other localities

Formation — 1 By heating Pb to low redness in presence of air, if the temperature is kept below the melting point of the PbO , the oxide is obtained as a yellowish powder, known commercially as *massicot*, if the PbO is melted during the preparation, the oxide solidifies to a scaly reddish solid known as *litharge* — 2 By heating Pb with KNO_3 — 3 By strongly heating red lead in air — 4 By strongly heating 'white lead' on an iron plate — 5 By adding boiling Pb acetate solution to boiling KOH Aq , pp is crystalline (Winkelblech, A 21, 21, J pr 10, 227) — 6 By slowly pouring Pb acetate solution into CaOAq at 80° until a crystalline crust appears, then adding a little more, and allowing to cool (Brendecke, R P 55, 118) — 7 By adding PbSO_4 made into a paste with water, to boiling NaOHAq (Wichmann, C C 1860 334) For modifications of these processes v Payen, J pr 13, 485, Behrens, B J 24, 134 8 According to Yorke (P M [3] 5, 82), crystals of PbO are produced by placing Pb in a vessel of water exposed to the air The reaction of steam with Pb at white heat produces PbO

Preparation — 1 Basic Pb nitrate is heated in a Pt crucible until completely decomposed (the neutral salt melts before decomposition is complete) Berzelius recommends the following procedure Equal weights of PbO and Pb_2NO_3 are kept in boiling water until the PbO is wholly transformed into basic nitrate, the basic salt is dissolved in boiling water, the solution is filtered hot and allowed to cool, the basic nitrate which separates is subjected to the action of the boiling solution as long as any solid separates on cooling.

A small quantity of the basic nitrate is made into a paste with water, the inside of a Pt crucible is covered with this paste, and the crucible is dried, the rest of the salt is strongly pressed while moist, then dried, and broken into smaller pieces, which are placed in the crucible, which is then heated until the decomposition is completed. The PbO formed from the pieces of basic nitrate in the middle of the crucible is pure, the outer layers of PbO contain a little Pt—2 PbO is prepared in crystalline form by dissolving PbO, prepared as in 1, in hot KOHAq or NaOHAq, and allowing to cool. The colour and SG of the crystals vary with the conditions. Ditte (C R 94, 1310) obtained a greenish crystalline mass (SG 9 1699) by heating KOHAq c 13 p.c. with PbO \cdot 2H₂O, using 25 p.c. KOHAq he obtained sulphur yellow lustrous crystals (SG 9 2089), brownish needles (SG 9 8835) were obtained by using 30 p.c. KOHAq and allowing to cool, KOHAq 185 p.c., saturated with PbO \cdot 2H₂O when hot, gave deep green, almost black, crystals on cooling (SG 9 5605), saturated KOHAq boiled with PbO \cdot 2H₂O gave deep rose coloured crystals of PbO (SG 9 4223), differing in form from the usual crystals, and becoming yellow when strongly heated (cf. Mitscherlich, J pr 19, 451, Calvert, B J 24, 135, Becquerel, A Ch [2] 51, 105).

Properties—PbO is a yellow to reddish yellow crystalline solid, the colour and SG of the oxide when well crystallised seem to vary with the conditions of formation, *v Preparation*, No 2. Melts between 585° and 630° to a clear dark red liquid. According to Leblanc (B J 26, 193) molten PbO absorbs O, which it gives off on solidifying. PbO is sl sol water, 1 part dissolves in c 7000 water, the solubility is affected by the state of aggregation of the PbO (cf. Bineau, C R 1855 877), if the solution is exposed to the air CO₂ is absorbed, and the PbO as basic carbonate, the PbO is also removed by filtration through paper (Yorke, P M [3] 5, 82). Heated to between 300° and 450° in air, PbO is oxidised to Pb₂O₃, but at higher temperatures Pb₂O₃ is deoxidised to PbO. PbO is soluble in warm conc KOHAq or NaOHAq. Geuther (A 219, 56) says that two varieties of PbO exist, one yellow and the other red, the yellow crystallises in rhombic, the red in hexagonal, forms.

Reactions—1 Heated in air Pb₂O₃ is formed at c 300°–450°, at a somewhat higher temperature PbO is re formed—2 Heated in hydrogen, or carbon monoxide, reduction to Pb occurs at a little above 100°, PbO is also reduced to Pb by heating with C, Na, or KCN—3 Reacts with most acids to form salts PbX₂, X = NO₃, Cl, SO₄, &c., absorbs CO, from the air forming PbCO₃—4 Decomposes alkali salts with separation of the alkali—5 Dissolves in warm fairly conc potash or soda solution, probably with formation of plumbites K₂(Na₂)PbO₂—6 Dissolves in molten potash forming K₂PbO₂, which, on long continued heating in air, is oxidised to K₂PbO₄ (Carnegie, C N 60, 113) (*v PLUMBATES* under LEAD PEROXIDE, p 132)—7 Dissolves also by boiling with milk of lime on evaporating in absence of air a compound separates in needles, probably CaPbO₂—8 Molten PbO dissolves silica, easily forming glass like silicates, these

silicates readily dissolve CaO, Al₂O₃, &c.—9 Heated in chlorine, bromine, or iodine vapour, the compound PbCl₂, PbBr₂, or PbI₂ is formed, heated with Cl and air, Br and air, or I and air, the product is Pb₂Cl₂, Pb₂Br₂, or Pb₂I₂ (Cross a Sigura, C J 33, 405)—10 By boiling excess of PbO with conc calcium chloride solution, filtering, and washing with alcohol the pp that forms on cooling, André (C R 104, 359) obtained 2PbO CaO CaCl₂ 4H₂O, by using strontium chloride he obtained 2PbO SrCl₂ 5H₂O—11 PbO does not react with phosphorus trichloride at 160°, but when heated together over a flame, PbO and PCl₃ react violently forming PbCl₂, Pb(H₂PO₃)₂, and P (Michaels, J pr [2] 4, 449)—12 Dissolves in magnesium chloride solution, on filtering and evaporating, the oxychloride 8PbO PbCl₂ H₂O separates (Voigt, Chem Zeitung, 13, 695).

HYDRATES OF LEAD MONOXIDE—1 3PbO H₂O, obtained by adding c 400 grams KOH to 1000 cc water containing freshly pptd PbO in suspension (Ditte, C R 94, 1310), also by adding basic Pb acetate solution to NH₄Aq at 20°–25°, and digesting the pp under the liquid for some time (Payen, A Ch [4] 8, 302, Behrens, B J 24, 134). This hydrate forms lustrous octahedral crystals, SG 7 592 at 0° (Ditte, C R 94, 1310), loses some H₂O at 130°, and is entirely dehydrated at 145°, a sol caustic alkali solutions—2 2PbO H₂O By pptg Pb(C₂H₃O₂)₂ Aq by KOHAq or NaOHAq, and digesting with excess of pptant, if Pb(NO₃)₂ is used some basic nitrate is always formed (Schaffner, A 51, 175). The hydrates of PbO readily absorb CO₂ from the air, they also combine with NH₃ to form PbO H₂O 2NH₃ and 8PbO 2NH₃ H₂O (Calvert, C R 23, 480).

Plumbites Although PbO is a markedly basic oxide, it nevertheless dissolves in alkali solutions to form unstable salts of the form M₂PbO₂, these salts, plumbites, have not been much investigated, according to Carnegie (C N 60, 113) K plumbite is formed by dissolving PbO in molten KOH, and on long continued heating in air it is oxidised to plumbate K₂PbO₄. The Ag salt is produced by adding NaOHAq to a mixture of a Pb and a Ag salt, Wöhler gives the formula Ag₂O 2PbO Krutwig (B 15, 1264) obtained a yellow salt, Ag₂PbO₂ 2H₂O, by adding AgNO₃Aq to alkaline Pb(NO₃)₂Aq, and washing with hot KOHAq and then with hot water, the salt soon became black in the air.

LEAD SESQUIOXIDE Pb₂O₃ (*Plumbo plumbica oxide*) Mol w unknown, as oxide has not been gashed. A reddish yellow powder resembling PbO, Debray describes Pb₂O₃ as greenish-brown (C R 86, 513). Debray (l.c) prepares this oxide by heating PbO₂ to 350°, or by heating PbO or better PbCO₃ to the same temperature in a stream of air or O, Carnellea Walker (C J 53, 85) say that PbO₂ is changed to Pb₂O₃ at 280°–290°, and that Pb₂O₃ gives off O at c 370°. According to Jacquelin (J pr 53, 153) Pb₂O₃ may be obtained by pouring a solution of red lead, Pb₃O₄, in glacial acetic acid into very dilute NH₄Aq, separating the pp quickly, washing it with hot water containing a very little acetic acid, and drying at 100°. Winkelblech (A 21, 21) obtained this oxide by pouring NaOHAq into Pb(C₂H₃O₂)₂Aq until the pp dissolved, and then

adding cold NaClO_4aq , taking care to avoid excess which would produce PbO_2 . Debray asserts that this method yields only a mixture of PbO and PbO_2 . Pb_2O_3 is not decomposed by heat until c 860° , but a little above this temperature it gives off O, leaving Pb_2O_3 , heated to c 530° , PbO is produced (Carnelley & Walker, *C J* 53, 85). Pb_2O_3 is resolved by acids into PbO_2 and a salt of PbO . Winkelblech's statement that Pb_2O_3 is dissolved unchanged by HClAq , from which solution it may be pptd again by alkalis, is denied by Hausmann (*A* 91, 235). Pb_2O_3 is reduced to PbO by $\text{H}_2\text{C}_2\text{O}_4\text{aq}$ or $\text{H CO}_3\text{HAq}$.

HYDRATE OF LEAD SESQUIOXIDE $\text{Pb}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$
Obtained by adding PbO to conc $\text{K}_2\text{PbO}_4\text{Aq}$, as a red curdy ppt (Seidel, *J pr* [2] 20, 200). The solution of K_2PbO_4 is obtained by dissolving pure PbO_2 in molten KOH with a very little water, and then dissolving the fused mass in a small quantity of water.

RED OXIDE OF LEAD Pb_2O_3 , (*Diphumboplumbic oxide* Red lead *Minium*) Mol w unknown, as oxide has not been gasified SG 9.096 at 15° (Herapath, *P M* 64, 321).

Occurrence—*Minium* is found mixed with other lead ores in Yorkshire, Anglesey, Virginia, the Eifel, &c.

Formation—By heating PbO in air to low redness (not above 450°) for some time, and cooling slowly—2 By adding PbO in KOH Aq to $\text{K}_2\text{PbO}_4\text{Aq}$, washing the ppt with hot water, and heating it—3 Finely divided ppt Pb exposed to air in presence of water and a little NH_4Aq is slowly changed to Pb_2O_3 , mixed with Pb carbonate (Schonbein, *J pr* 74, 323).

Preparation—1 A mixture of 1 pt KClO_3 , 4 pts PbO (prepared by heating PbCO_3), and 8 pts KNO_3 is heated, the mass melts, and PbO_2 is formed and on continued heating is changed to Pb_2O_3 , the fused mass is treated with water and the residue is boiled with dilute KOH Aq , to dissolve unchanged PbO , washed and dried (Levol, *J pr* 22, 38)—2 5 to 6 pts $\text{Pb(NO}_3)_2$ are dissolved in water, and KOH is added until the ppt which forms is dissolved, to this liquid is added 1 pt PbO_2 , and the whole is boiled for some time, unchanged PbO_2 is decomposed by digesting with $\text{H}_2\text{C}_2\text{O}_4\text{Aq}$, and the PbC_2O_4 formed is dissolved in KOH Aq , the residue is washed and dried (Levol, *lc*)—3 Commercial red lead usually contains some PbO , this may be removed by digestion with dilute acetic acid, or better with lead acetate solution Lowe (*D P J* 271, 472) recommends to digest 1 pt commercial red lead with 10–15 pts of a 10 p c solution of $\text{Pb(NO}_3)_2$, for some time at a gentle heat, then to boil, and finally wash well.

Properties—A scarlet crystalline granular powder, when heated it becomes brighter red and then violet, at c 500° – 530° decomposition to PbO and O occurs SG according to different observers varies from 8.94 to 9.1 (*v* Clarke's *Specific Gravity Tables* [2nd ed], p 47). Insol water, reacts with dilute acids to form salts of PbO , and PbO_2 which remains undissolved, with conc H_2SO_4 , or HClAq , gives salts of PbO and O or Cl. Dissolves completely in glacial acetic acid forming a liquid which acts as an energetic oxidiser (*v* *Reactions*, No 6). Pb_2O_3 is readily reduced to PbO , e.g. by SO_4Aq or HNO_3Aq .

Reactions—1 Heated to c 500° – 530° , gives

PbO and O—2 Reduced to PbO by easily oxidised bodies, e.g. SO_4Aq , HNO_3Aq , solution of sugar—3 With stannous chloride gives PbCl_2 , and SnCl_4 —4 Dissolves in considerable quantity of conc cold hydrochloric acid to form a yellow liquid from which alkalis, alkaline carbonates, borates, and acetates, and oxides and carbonates of Ba, Mg, Zn, &c ppt PbO_2 , on heating, the solution decomposes to PbCl_2 and Cl, this solution almost certainly contains PbCl_4 , on warming Cl is evolved and PbCl_2 ppt (Fisher, *C J* 35, 282, *v* *Lead tetrachloride*, p 126). Dilute HClAq forms PbCl_2 , H_2O , and PbO_2 . Warm HClAq in excess evolves Cl and ppts PbCl_2 —5 Dilute sulphuric, nitric, or acetic acid, and other dilute acids ppt PbO_2 and form sulphate, &c of PbO . Hot conc H_2SO_4 forms PbSO_4 , and O—6 Pb_2O_3 dissolves in glacial acetic acid, this solution is strongly oxidising, e.g. it converts SO_4Aq to SO_4Aq , $\text{As}_2\text{O}_3\text{Aq}$ to $\text{As}_2\text{O}_5\text{Aq}$, PbS to PbSO_4 , Pb , Hg , and Cu to oxides, separates I from KI (Schönbein, *J pr* 74, 315), when dropped into dilute NH_4Aq a ppt of Pb_2O_3 is produced (Jacquelain, *J pr* 53, 153, *v* *Lead sesquioxide*, p 129). According to Jacquelain (*loc cit*) a solution of Pb_2O_3 in glacial acetic acid at 40° gives crystals of $\text{PbO}_4(\text{C}_2\text{H}_3\text{O}_4)_2$, $[\text{O}=8, \text{C}=6]$ [old notation], these crystals may be kept under acetic acid but give PbO , and $\text{H}_2\text{C}_2\text{O}_4$, on warming, or with water. Schonbein (*J pr* 74, 315) says that conc acetic acid dissolves about 9 p c of Pb_2O_3 , when shaken for 15 mins with excess of the finely powdered oxide, the solution is easily decomposed by heat or dilution, but is stable at c -18° , by careful addition of H_2SO_4 , PbSO_4 is pptd and an acetate corresponding to Pb_2O_3 remains in solution, this solution deposits PbO_2 , slowly at ordinary temperatures, quickly when heated, addition of KOH Aq does not ppt Pb_2O_3 , but a mixture of PbO and PbO_2 —7 Pb_2O_3 dissolves in conc phosphoric acid, the solution reacts similarly to that of Pb_2O_3 in acetic acid. By using a mixture of phosphoric and sulphuric acids, a solution is obtained containing only the phosphate corresponding to Pb_2O_3 , and free from PbO (Jacquelain, *J pr* 53, 152, *v* *Lead peroxide*, p 131)—8 Conc solutions of arsenic and tartaric acids dissolve Pb_2O_3 , the solutions resemble those in acetic and phosphoric acids, but are more easily decomposed.

Composition and constitution of red lead—When PbO is heated in air, O is slowly and continuously absorbed, when c 176 p c O has been absorbed the product is red, but absorption of O continues, and some specimens of red lead contain 2.67 p c O in excess of that contained in the PbO (Jacquelain, *J pr* 53, 151). An absorption by PbO of 1.79 p c O corresponds with the formation of Pb_2O_3 , the formation of Pb_2O_3 requires 2.39 p c. Older analyses of red lead led to the supposition that it was Pb_2O_3 (Dumas, *B J* 13, 113). Careful syntheses and analyses have, however, shown that a definite red coloured oxide exists having the composition Pb_2O_3 . At the same time the analyses of commercial red leads made by Mulder (*J pr* 50, 438) and Jacquelain (*J pr* 53, 151) show that the percentage of O above that required to form PbO varies from 1.16 to 2.67. Several samples of red lead agreed in composition with the formula Pb_2O_3 .

others approached Pb_2O_3 , (*cf* Löwe, *D P J* 271, 472) It is still doubtful whether a series of oxides exists of the form $x\text{PbO} \cdot y\text{PbO}_2$, x varying from 1 to 3 or 4 and y from 1 to 2 or 3, or whether only two such oxides are definite bodies, viz $\text{PbO} \cdot \text{PbO}_2 (= \text{Pb}_2\text{O}_3)$ and $2\text{PbO} \cdot \text{PbO}_2 (= \text{Pb}_3\text{O}_4)$

Red lead, Pb_3O_4 , may be regarded either as $2\text{PbO} \cdot \text{PbO}_2$ or as $\text{PbO} \cdot \text{Pb}_2\text{O}_3$, the reaction of Pb_3O_4 with acids, the synthesis of Pb_3O_4 by the action of PbO in KOH aq on $\text{K}_2\text{PbO}_2 (= \text{K}_2\text{O} \cdot \text{PbO}_2)$, and the fact that Pb_3O_4 is acidic towards some oxides, point to the constitution $2\text{PbO} \cdot \text{PbO}_2$ rather than $\text{PbO} \cdot \text{Pb}_2\text{O}_3$

LEAD PEROXIDE PbO_2 (*Brown oxide of lead* *Puce oxide of lead* *Lead dioxide*) Mol w unknown, as oxide has not been gasified SG 8 902 at 16.5° (Herapath, *P M* 64, 321), 8 756 to 8 897 (Playfair a Joule, *C S Mem* 3, 84), 9 045 (Wernicke, *P* 141, 109)

Occurrence—As *plattnerite*, at Leadhills, in hexagonal prisms

Formation—1 By boiling a Pb salt with a filtered solution of bleaching powder —2 By digesting red lead with dilute HNO_3 aq, and washing the residue —3 By mixing solutions of Pb acetate (4 pts) with soda crystals ($3\frac{1}{2}$ pts), and passing Cl into the mixture $2\frac{1}{2}$ pts PbO_2 are obtained, PbCl_2 is not formed (Wehler, *J pr* 90, 383, *cf* Geuther, *A* 96, 382)—4 By heating PbO with KClO_3 to less than red heat and washing the mass with water —5 By boiling PbO in H_2O with $\text{K}_2\text{FeC}_7\text{O}_{14}$ aq in presence of KOH ($2\text{K}_2\text{FeC}_7\text{O}_{14} + 2\text{KOH} + \text{PbO} = 2\text{K}_2\text{FeC}_7\text{O}_{14} + \text{PbO}_2 + \text{H}_2\text{O}$, Overbeck, *Ar Ph* [2] 85, 5)—6 By fusing PbO with KOH for some time (Becquerel, *A Ch* 51, 504) Carnegie (*C N* 60, 113) obtained $\text{PbO}_2 \cdot \text{H}_2\text{O}$ by long continued fusion of PbO in KOH solution in water, and careful neutralisation by dilute H_2SO_4 aq (*v Hydrates of lead peroxide, infra*) —7 By oxidising PbO by an alkaline solution of KMnO_4 (Reynoso, *C R* 32, 646)—8 By electrolysis strongly alkaline solutions of Pb salts, the hydrate $\text{PbO} \cdot \text{H}_2\text{O}$ ($q v$) is deposited on the positive electrode (Becquerel, *A Ch* [3] 8, 405) By slowly electrolysis a solution of 1 pt $\text{Pb}(\text{NO}_3)_2$ in 8 pts water, Wernicke obtained PbO_2 (*P* 141, 109)—9 PbO_2 is obtained, mixed with PbO , by shaking basic Pb acetate with H_2O_2 aq (Schonbein, *J pr* 75, 85)

Preparation—1 Very finely powdered Pb acetate is boiled with a filtered solution of bleaching powder, the liquid being poured off, and fresh solution added from time to time until no Pb is found in solution, the residue is washed with very dilute HNO_3 aq, and then with hot water, and dried at a low temperature (Böttger, *J pr* 73, 493)—2 Fehrmann (*B* 15, 1882) recommends to add a filtered solution of bleaching powder to conc PbCl_2 aq at 50° to 60° , until a few drops of the supernatant liquid cease to show a brown colour when filtered, and to wash the pp out of contact with air

Properties—A dark puce brown powder, black according to Fehrmann (*B* 15, 1882) The native compound forms metal like, lustrous, iron-black, hexagonal crystals, SG 9 392 to 9 448 (Breithaupt, *J pr* 10, 508) When heated, PbO_2 gives Pb_2O_3 and O , and at a higher temperature it is resolved into PbO and O Readily

parts with O to oxidisable bodies, *eg* rubbed with P or S vivid combustion occurs, sugar and gallic acid &c are ignited by trituration with PbO_2 . Acids generally react with PbO_2 to form salts of PbO , cold conc HCl aq seems to produce PbCl_2 , and solutions of PbO in glacial acetic and phosphoric acids probably contain salts corresponding with PbO_2 (*v Reactions*, No 9) PbO_2 reacts with strongly basic oxides as an acidic oxide to form plumbates M_2PbO_4 (*v infra*, *Plumbates*) On account of its readiness to part with O , PbO_2 is largely used for forming a surface on which to strike 'safety' matches

Reactions—1 Heated, easily goes to PbO , and O , and at a higher temperature to PbO and O —2 Readily gives up $\frac{1}{2}$ of its O to easily oxidised bodies, either at the ordinary temperature or by slightly heating, *eg* when PbO_2 is rubbed with an equal weight of $\text{H}_2\text{C}_2\text{O}_4$, H_2O , CO_2 , and PbCO_3 are formed, PbO_2 rubbed with $\frac{1}{2}$ pt sugar or $\frac{1}{2}$ pt tartaric acid oxidises these rapidly, causing ignition, P is oxidised and ignited when rubbed with PbO_2 (*v Böttger*, *A* 34, 94)—3 Sulphur dioxide is rapidly absorbed by PbO_2 with rise of temperature and production of SO_2 —4 Potassium iodide is decomposed with liberation of I —5 PbO_2 reacts with a dilute solution of potassium ferrocyanide to form K_2FeCy_6 provided the KOH produced in the reaction is neutralised by CO_2 , with conc potassium ferrocyanide solution in presence of KOH , K_2FeCy_6 is formed —6 By digesting PbO_2 with ammoniac solution, PbO , H_2O , and NH_4NO_3 are produced —7 Not acted on by cold nitric or sulphuric acid, with hot conc H_2SO_4 gives PbSO_4 and O —8 Cold conc hydrochloric acid in excess dissolves PbO_2 , producing a yellow liquid which evolves Cl on heating and gives a pp of PbCl_2 , alkalis, alkaline carbonates, and alkaline earth oxides and carbonates ppt PbO from this solution, the solution almost certainly contains PbCl_2 (*v Lead tetrachloride*, p 126) Hot HCl aq evolves Cl and forms PbCl_2 —9 A basic acetate derived from PbO , is said by Jacquelin (*J pr* 53, 152) to be obtained by dissolving red lead in glacial acetic acid at 40° and cooling, J gives the formula $\text{PbO} \cdot (\text{C}_2\text{H}_3\text{O}_2)_2$, [$\text{O} = 8$, $\text{C} = 6$] [old notation] to the crystals which form These crystals are unchanged in acetic acid, on drying between paper, or on adding water, they are resolved into PbO and acetic acid Similar salts derived from PbO , are obtained by dissolving red lead in very conc phosphoric and arsenic acids, but the salts are very unstable (*J l c. cf Red oxide of lead, Reactions*, No 6, p 130)—10 Heated with phosphorus trichloride, PbCl_2 , POCl_3 , and $\text{Pb}(\text{HPO}_3)_2$ are formed (Michaelis, *J pr* [2] 4, 449)—11 Heated in chlorine, bromine, and iodine, the compounds $\text{Pb}_2\text{Cl}_2\text{O}$, $\text{Pb}_2\text{Br}_2\text{O}$, and $\text{Pb}_2\text{I}_2\text{O}$, respectively, are obtained, heated in chlorine and air, bromine and air, and iodine and air, the products are $\text{Pb}_2\text{Cl}_2\text{O}$, $\text{Pb}_2\text{Br}_2\text{O}$, and $\text{Pb}_2\text{I}_2\text{O}$, respectively (Crosa a. Sigura, *C J* 33, 405)—12 PbO_2 reacts with alkalis to form compounds in which PbO forms the negative radicle, *eg* the salt K_2PbO_4 may be obtained from a solution in conc hot KOH aq (*v infra*, *Plumbates*)

HYDRATES OF LEAD PEROXIDE (1) $\text{PbO}_2 \cdot \text{H}_2\text{O}$ This hydrate is formed as a brown pp, by dis

solving PbO in molten KOH, heating in air for a considerable time, dissolving the K_2PbO_2 produced in cold water, and *exactly neutralising* by dilute H_2SO_4 (Carnegie, *C N* 60, 113). It is also probably obtained, at the positive electrode, by the electrolysis of strongly alkaline solutions of Pb salts (Becquerel, *A Ch* [3] 8, 405). By electrolysis of an alkaline solution of Pb Na tartrate, Wernicke (*P* 141, 109) obtained a lustrous blue-black film, which had the composition $PbO_2 \cdot H_2O$, SG 6 267. (2) According to Carnelley & Walker (*C J* 53, 85), ppd PbO, after drying in the air for 10 days has very nearly the composition $3PbO_2 \cdot H_2O$, at 230° this hydrate is changed to PbO_2 .

Plumbites.— PbO_2 reacts with acids as a basic peroxide forming salts corresponding with PbO , in some cases unstable salts derived from PbO_2 appear to be produced (*vs supra*, *Lead peroxide*, *Reactions* 8 and 9). PbO_2 also reacts with alkalis as an acidic oxide to form salts M'_2PbO_3 , known as *plumbites*. Fremy (*C R* 15, 1109) obtained colourless crystals of $K_2PbO_3 \cdot 3H_2O$ by dissolving PbO_2 in boiling very conc KOH, adding a little water, and placing over H_2SO_4 . Seidel (*J pr* [2] 20, 200) obtained the salt by adding PbO_2 to molten KOH containing a little water, dissolving the fused mass in water, and evaporating over H_2SO_4 *in vacuo*. The crystals form quadratic octahedra, $a:c = 1:1.2216$, they are decomposed by water with separation of PbO_2 . A corresponding Na salt is known. Solutions of these salts are said to yield pps of plumbites when added to solutions of various metallic salts. K_2PbO_3 is also produced by dissolving PbO in molten KOH and heating in air (Becquerel, *A Ch* 51, 504, Carnegie, *C N* 60, 113). Crum obtained a plumbite of Ca by digesting $Pb(NO_3)_2$ with CaO and bleaching powder (*A* 55, 213). For PLUMBATES *vs* KASSNER (*Ar Ph* 232, 375 [1894]).

Lead, oxybromides of Several oxybromides of Pb are known. The compound $PbO \cdot PbBr_2$ is formed by heating $PbBr_2$ in air so long as white fumes are evolved, also by digesting $PbBr_2$ with solution of Pb acetate. By heating PbO in Br vapour, Cross & Sigura obtained $Pb_2Br_3O_2$, and by heating the same oxide in a mixture of Br and air they obtained $Pb_2Br_3O_2$ (*C J* 33, 405), from PbO_2 they obtained $Pb_2Br_3O_2$ by heating in Br.

Lead, oxychlorides of $PbCl_2$ and PbO very readily combine when heated together to form oxychlorides from which HNO_3 dissolves out PbO .

$PbCl_2 \cdot PbO$ occurs native as *matlockite*, SG 7 21 (Greg, *P M* [4] 2, 120, Rammelsberg, *P* 75, 141), formed by heating $PbCl_2$ in air until fumes are no longer evolved, also by digesting freshly ppd $PbCl_2$ with cold Pb acetate solution (the oxychloride thus formed is $(PbCl_2 \cdot PbO)_2 \cdot H_2O$ according to Brandes, *A* 10, 273), also (with H_2O) by partial ppn of $PbCl_2$ by CaO (thus prepared it is used as a pigment (*Pattinson's white lead*)).

$PbCl_2 \cdot 2PbO$ occurs native as *mendipite*, in yellow white trimetric prisms, SG 7 to 7.1, formed, according to Ditté (*C R* 94, 1180), by gradually adding KOH to $PbCl_2$ suspended in water, or by adding KCl to $PbO \cdot xH_2O$.

$PbCl_2 \cdot 3PbO$. the hydrate with $4H_2O$ is

formed by adding NH_4Aq to $PbCl_2$ (Berzelius); also by mixing $NaCl$ with PbO , making into a paste with water, and allowing to stand, the product of this reaction, when washed and heated, gives a yellow powder, known as *Turner's yellow*, which is probably $PbCl_2 \cdot 3PbO$. The hydrate $PbCl_2 \cdot 3PbO \cdot H_2O$ was obtained by Voigt (*Chem Zeitung*, 13, 6⁶⁵) by dissolving PbO in $MgCl_2$ Aq, filtering, and evaporating.

$PbCl_2 \cdot 7PbO$ a lustrous, golden-yellow, crystalline mass, obtained by melting together 1 pt pure NH_4Cl and 10 pts pure PbO , pouring off from a little Pb (formed by the reducing action of NH_3 set free by the PbO acting on NH_4Cl), and allowing to cool, known as *Cassel yellow*.

By heating PbO in a mixture of Cl and air, Cross & Sigura obtained Pb_2Cl_3O (*C J* 33, 405), by heating PbO_2 in Cl the product was Pb_2Cl_3O , and by heating PbO in a mixture of Cl and air the compound Pb_2Cl_3O was obtained.

According to André (*A Ch* [6] 3, 104) $PbCl_2 \cdot PbO$, $PbCl_2 \cdot 2PbO$, and $PbCl_2 \cdot 3PbO$ are obtained by ppg $PbCl_2$ by KOH Aq.

Lead, oxycyanide of, *vs vol* 11 p 341

Lead, oxyiodides of Several oxyiodides of lead are known. $PbI_2 \cdot PbO \cdot H_2O$ is formed by adding $KIAq$ to excess of Pb acetate solution and digesting the pp in the liquid, it is a yellowish-white powder, insol water (Brandes, *A* 10, 269, cf Ditté, *C R* 92, 145). Other oxyiodides, described by Kuhn (*Ph C* 1847 593) and Denot (*J Ph* 20, 1) are $PbI_2 \cdot 2PbO \cdot H_2O$, $PbI_2 \cdot 3PbO \cdot 2H_2O$, and $PbI_2 \cdot 5PbO$. Cross & Sigura (*C J* 33, 405) describe the oxyiodides $Pb_{11}I_{10}O_{10}$, $Pb_2I_3O_3$, $Pb_3I_4O_4$, and $Pb_4I_5O_5$, obtained by heating PbO_2 and PbO , in I and air, and in I only.

Lead, phosphide of No phosphide of Pb has been isolated with certainty. Pelletier (*A Ch* [2] 13, 114) said that Pb takes up c 15 p c P when P is thrown on to melted Pb, or when Pb filings are strongly heated with glacial phosphoric acid, or $PbCl_2$ with P, but the experiments of Cloud (in Percy's laboratory) make it doubtful whether Pb and P combine. According to H Rose (*P* 24, 326) a brown pp, consisting of a phosphide of Pb, is obtained by passing PH_3 into Pb acetate solution.

Lead, salts of Compounds formed by replacing the H of acids by Pb. Pb forms one series of well marked salts, PbX_n , where $X = Cl, NO_3, \frac{1}{2}SO_4, \frac{1}{2}CO_3, \frac{1}{2}PO_4$, &c. $PbCl_2$ very probably exists in a solution of PbO_2 , or Pb_2O_3 , in cold conc $HClAq$, and $PbO(C_2H_3O_2)_2$ is said to have been isolated, these two salts belong to the series PbX_n , corresponding with the oxide PbO . Very many basic salts of Pb are known, and a considerable number of double salts. The formulæ of the Pb salts are determined from analyses, comparison with the salts and compounds of Sn, Ge, and Si, and from the vapour densities of $PbCl_2$ and $PbMe$. The following are the chief salts of lead (v CARBONATES, NITRATES, SULPHATES, &c.). *Antimonate*, *arsenate* and *ite*, *borate*, *bromate*, *carbonates*, *chlorate* and *ite*, *chromates*, *dithionate*, *hypophosphite*, *iodate*, *molybdate*, *nitrates* and *ites*, *pentathionate*, *perchlorate*, *periodates*, *phosphates* and *ites*, *sulphates* and *ites*, *selenate* and *ite*, *silicates*, *thiosulphate*, *trithionate*, *tungstates*, *urates*, *vanadates*.

Lead, selenide of, PbSe Occurs native as *clausthalite* in the Hartz, &c, the Pb being sometimes partly replaced by Co, Cu, Hg, Ag, &c, S G 7 to 88 (H Rose, P 2, 416, 3, 281, Stromeyer, P 2, 403, Kersten, P 46, 265) PbSe is produced by heating the constituents in the ratio Pb Se, it forms a porous, grey, soft mass (Berzelius), melted under borax small regular crystals are obtained, S G 8154 (Little, A 112, 212) Heated in air Se is vaporised, then a little PbSe, and a residue of basic selenite of Pb is left, cold HNO₃ dissolves Pb, leaving Se

Lead, selenocyanide of, Pb(SeCy)₂, v vol u p 348

Lead, silicofluoride of, Pb(BF₄)₂ Long prismatic crystals, by dissolving PbO in HBF₄ and evaporating to a syrup, partially decomposed by water or alcohol (Berzelius, P 2, 113)

Lead, sulphides of Besides PbS, which is a well marked compound, two subsulphides, Pb₂S and Pb₃S, probably exist, there are also indications of the formation of a persulphide

LEAD SULPHIDE (Lead monosulphide Plumbous sulphide) PbS

Occurrence—Native as *galena*, crystallised in monometric octahedra, and cubic and rhombic dodecahedra, S G 725 to 77 (v Neumann, P 23, 1)

Preparation—1 By mixing S with molten Pb—2 By heating PbO with excess of S—3 By the action of H₂S or an alkaline sulphide on Pb salts—4 By passing CS over Pb heated a little above redness, crystals of PbS are formed (Pb thiocarbonate seems to be also produced) (Gautier a Hallopeau, C R 104, 111)—5 Well formed crystals of PbS are obtained by passing H₂S into a solution of c 3 grams Pb(NO₃)₂ in 250 c c 10 p c HNO₃ at c 15°, if only c 1 p c HNO₃ is present the pp is amorphous (Muck, Z [2] 4, 211)—6 Crystalline PbS is produced by the action of PbO on thiourea (Reynolds, C J 45, 162) Addition of H₂S to a Pb salt in presence of HClAq sometimes produces a red pp of 3PbS PbCl₂, on warming and passing in more H₂S, PbS is formed—7 Winssinger (Bl [2] 49, 452) obtained an aqueous solution of colloidal PbS by ppg by H₂S a very dilute solution of a Pb salt, and dialysing, the solution was reddish brown

Properties—Lead grey crystalline solid, as prepared by fusion of Pb and S, S G 7505 (Karsten, S 65, 394) A brown black powder, as prepared by ppg by H₂S S G 6924 at 4° (Playfair a Joule, C J 1, 137), 677 (Schneider, J pr [2] 2, 91) Melts at full red heat out of contact with air, and is said to sublime unchanged Insol in dilute acids, caustic alkalis, and alkaline sulphide solutions

Reactions—1 Heated in air evolves SO₂, and forms Pb and P¹ SO₄, and some PbO (Descotils, A Ch [2] 55, 441)—2 Heated with *lead monoxide* forms Pb and SO₂, Pb and SO₂ are also produced by heating with *cad sulphate*, if little PbS is heated with much PbSO₄, the chief products are PbO and SO₂ (v LEAD, Formation, Nos 1 and 2, p 122)—3 Melted with *scrap iron* FeS and Pb are formed—4 Heated in *steam* gives H₂S, and at first PbO and afterwards Pb—5 Heated with *alkaline carbonates* half the Pb of

the PbS is separated—6 Boiled with *dilute nitric acid* Pb(NO₃)₂, S, and NO are produced, *fuming nitric acid* forms PbSO₄, and S, and sometimes Pb(NO₃)₂—7 *Conc hydrochloric acid* forms PbCl₂, and H₂S—8 *Aqua regia* forms PbCl₂, and PbSO₄—9 Slowly reacts with *chlorine* to form PbCl₂, and S₂Cl₂—10 By long *fusion with potash and nitre* Pb₂O₃ is produced (Carnegie, C N 60, 113)—11 Heated with lead, subsulphides (q v) Pb₂S and Pb₃S are formed

Combinations—With *lead chloride* to form the sulphochloride (q v) 3PbS PbCl₂

LEAD SUBSULPHIDES Pb₂S and Pb₃S Said to be formed by heating together PbS and Pb in the proper proportions (v Bredberg, P 17, 274) Pb₂S is also formed, according to Berthier (A Ch 3] 22, 240), by heating PbSO₄ in a crucible lined with charcoal

LEAD PERSULPHIDE Addition of K penta-sulphide to solution of a Pb salt gives a blood-red pp which is quickly changed, even in the liquid in which it is produced, into PbS and S Berzelius gives the formula PbS₅ to the persulphide

Lead, sulphochloride of 3PbS PbCl₂ Obtained as a red pp by passing a little H₂S into a dilute Pb solution acidified by HCl, Renisch (J pr 13, 130) obtained the compound by passing H₂S into a solution of 1 part Pb acetate in 200 parts water to which 20 parts HClAq, S G 1168, were added Also produced as a yellowish red solid by digesting freshly ppg PbS with PbCl₂ Aq Decomposed by H₂S with formation of PbS Boiling water partly dissolves out PbCl₂

Lead, sulphocyanide of Pb(SCy)₂, v vol u p 350 Combines with PbBr₂, and PbCl₂, v. *Lead bromosulphocyanide*, and *Lead chlorosulphocyanide*, pp 125, 127

Lead, thiocarbonate of PbCS Formed by adding solution of an alkali thiocarbonate to a Pb salt (cf THIOCARBONATES, vol 1 p 703)

M M P M

LEAD, WHITE White lead is a hydrocarbonate of Pb varying in composition, v vol u p 639

LEAD ORGANIC DERIVATIVES

Lead trimethyl salts Chloride PbMe₃Cl Formed by the action of boiling aqueous HCl on PbMe, (Cahours A Ch [3] 62, 257, A 122, 48) Long needles resembling PbCl₂ May be sublimed M sol boiling water and alcohol Converted into PbCl₂ by long boiling with HClAq

Bromide PbMe₃Br More soluble than the chloride

Iodide PbMe₃I Obtained by adding iodine to PbMe, till the odour becomes permanent White mass, crystallising from water in long colourless needles which may be sublimed Sl sol water, v sol alcohol On distillation with solid KOH it yields PbMe₃OH, a strong basic oil smelling like oil of mustard solidifying in prisms

Lead tetra-methyl, PbMe₄, (110°) (B) V D 952 (calc 925) S G ° 2034 From MeI and an alloy of lead (5 pts) and sodium (1 pt), or, better, from PbCl₂ and ZnMe₂ (Cahours) Colourless liquid, having a camphor like odour, insol water, sol alcohol and ether (Cahours) According to Butlerow it smells like strawberries or mould It does not unite directly with chlorine or iodine, but is decomposed thereby, e.g.

$\text{PbMe}_2 + \text{I}_2 = \text{PbMe}_2\text{I} + \text{MeI}$ Acids convert it into OH , and a salt of PbMe_2 .

Di-plumbic hexa ethide, Pb_2Et_6 . *Lead triethyl SG 2 1471* Formed by adding sodium (1 pt) to molten lead (3 pts) and mixing the cooled and powdered alloy with EtI in flasks provided with inverted condensers. When the first violent action has ceased the product is freed from EtI by distilling at 100° , and the residue extracted with ether (Klippel, *J pr* 81, 287, cf Löwig, *J pr* 60, 304, *A* 88, 318).

Properties — Yellowish mobile oil, insol water, \sqrt sol alcohol and ether. Decomposed on distillation. By exposure to light, or by prolonged boiling with water, it decomposes with separation of metallic lead. When exposed to the air in ethereal solution it absorbs O and CO_2 with formation of $(\text{PbEt}_2)_2\text{O}$ and $(\text{PbEt}_2)_2\text{CO}_2$. It combines with iodine forming PbEt_2I .

Lead triethyl salts *Chloride* PbEt_2Cl Obtained by adding BaCl_2 to a solution of $(\text{PbEt}_2)_2\text{SO}_4$ in alcohol, or by treating $(\text{PbEt}_2)_2\text{O}$, $(\text{PbEt}_2)_2\text{CO}_2$, or PbEt_2 with HCl . Beautiful long needles, giving off a pungent odour when heated, presently decomposing with separation of PbCl_2 and metallic lead — $\text{PbEt}_2\text{HgCl}_2$, white nacreous scales (from hot alcohol) — $(\text{PbEt}_2\text{Cl})_2\text{PtCl}_4$, copper-red crystals, sl sol water, m sol alcohol and ether.

Bromide PbEt_2Br Long needles (from ether).

Iodide PbEt_2I From Pb_2Et_6 and iodine, or from $(\text{PbEt}_2)_2\text{SO}_4$ and KI . Sol ether and very unstable, the ethereal solution quickly depositing PbI_2 .

Cyanide PbEt_2Cy From PbEt_2Cl and KCy in alcoholic solution at 100° . Prisms (from ether).

Hydroxide $\text{PbEt}_2(\text{OH})$ From the chloride by treatment with moist Ag_2O or by distillation with KOH . Needles, sl sol water, \sqrt sol alcohol and ether. Has strong alkaline reaction and caustic taste. Saponifies fats, ppts ferric, cupric, and argentic oxides from their salts. Ppts alumina and zinc oxide, the pp dissolving in excess of the precipitant. Expels NH_3 from its salts. Absorbs atmospheric CO_2 . Fumes with HCl . When heated it gives off white vapours which powerfully excite sneezing.

Carbonate $(\text{PbEt}_2)_2\text{CO}_3$ Small hard crystals, nearly insol water, sl sol alcohol and ether. Has a burning taste.

Nitrate PbEt_2NO_3 , unctuous crystalline mass, decomposed by heat with slight detonation. \sqrt sol alcohol and ether — $(\text{PbEt}_2)_2\text{SO}_4$, from the oxide and H_2SO_4 . White crystalline pp, nearly insol water, alcohol, and ether, \sqrt sol alcohol mixed with H_2SO_4 or HCl , from which solution it separates in octahedra.

Phosphate $(\text{PbEt}_2)_2\text{PO}_4$, stellate groups of crystals, \sqrt sol water, alcohol, and ether.

Sulphocyanide PbEt_2SCy crystals, sol water, alcohol, and ether.

The formate $\text{PbEt}_2\text{CHO}_2$, acetate $\text{PbEt}_2\text{C}_2\text{H}_3\text{O}_2$, butyrate $\text{PbEt}_2\text{C}_4\text{H}_7\text{O}_2$, oxalate $(\text{PbEt}_2)_2\text{C}_2\text{O}_4$, aq, tartrate $\text{PbEt}_2\text{C}_4\text{H}_4\text{O}_6$ (dried at 100°), and benzoate are crystalline.

Lead tetraethide, PbEt_4 . Mol w 323 (σ 200°), (152° at 190 mm) (B) SG 1.62.

Preparation — PbCl_2 is added to ZnEt_2 , as long as reaction occurs. The PbEt_4 is distilled

with steam $2\text{ZnEt}_2 + 2\text{PbCl}_2 = 2\text{ZnCl}_2 + 2\text{PbEt}_4$, and then $2\text{PbEt}_4 = \text{Pb} + \text{PbEt}_2$ (Buckton, *P M* [4] 18, 212, 17, 282, *A* 109, 218, 112, 220, Frankland & Lawrance, *C J* 35, 244).

Properties — Oil, not decomposed by water or by gaseous NH_3 , CO_2 , CO , C_2N_2 , NO , O , or H_2S . Partially decomposes at 200° .

Reactions — 1 Gaseous SO_2 reacts forming diethyl sulphone and lead ethyl sulphinate $\text{PbEt}_2 + 3\text{SO}_2 = \text{Et}_2\text{SO}_2 + (\text{EtSO}_2)_2\text{Pb}$ — 2 Iodine forms PbEt_2I and EtI — 3 Conc HCl forms PbEt_2Cl and ethane.

Di-plumbic hexa-iso-amylide, $\text{Pb}_2(\text{C}_5\text{H}_{11})_6$. Prepared like the corresponding Pb_2Et_6 , using isoamyl iodide (Klippel). Oil, with irritating vapour. Detonates with HNO_3 .

Lead tri-isoamyl salts *Chloride* $\text{Pb}(\text{C}_5\text{H}_{11})_3\text{Cl}$ from the oxide and HCl . White needles.

Iodide $\text{Pb}(\text{C}_5\text{H}_{11})_3\text{I}$ from $\text{Pb}_2(\text{C}_5\text{H}_{11})_6$ in ether by adding iodine [100°]. Needles, insol water — $\text{Pb}(\text{C}_5\text{H}_{11})_3\text{HgI}$, golden laminae, insol water, sl sol alcohol and ether — *Hydroxide* $\text{Pb}(\text{C}_5\text{H}_{11})_3\text{OH}$, viscid mass, insol water, sol alcohol and ether. Feebly alkaline, ppts ferric, but not cupric and silver salts.

Lead tetra-phenyl $\text{Pb}(\text{C}_6\text{H}_5)_4$, [225°] SG 22 1.530 Prepared by boiling 500 g of an 8 p.c alloy of sodium and lead with 500 g of bromo benzene and 20 c.c of acetic ether for sixty hours (Polis, *B* 20, 716, 3331). Small needles or dimetric prisms, $a:c=1.381$. Decomposes at 270° . \sqrt sol sl alcohol, ether, and HOAc , m sol benzene, CHCl_3 and CS_2 . Heated with HClAq at 230° it yields PbCl_2 , benzene, and chloro benzene.

Lead diphenyl salts *Chloride* PbPh_2Cl_2 Formed by passing chlorine into a solution of lead tetra phenyl in CS_2 , or by ppg $\text{PbPh}_2(\text{NO}_3)_2$ with KCl . White powder, insol alcohol and ether, \sqrt sl sol benzene.

Bromide PbPh_2Br_2 From lead di phenyl nitrate and KBr . White powder, which decomposes before fusion.

Iodide $\text{Pb}(\text{C}_6\text{H}_5)_2\text{I}_2$ [c 103°]. Formed by adding iodine to a solution of lead tetraphenyl in chloroform (Polis). Golden plates, sol chloroform, benzene, and alcohol.

Nitrate $\text{Pb}(\text{C}_6\text{H}_5)_2(\text{NO}_3)_2$ 2aq. Obtained by gradually adding lead tetra phenyl to boiling HNO_3 (SG 1.4). Small lustrous plates (from water). \sqrt sol hot water and alcohol. Detonates when heated. Boiling its aqueous solution produces a pp of the basic salt $\text{Pb}(\text{C}_6\text{H}_5)_2(\text{NO}_3)(\text{OH})$ as a white powder which detonates when heated.

Oxide $\text{Pb}(\text{C}_6\text{H}_5)_2\text{O}$ Obtained by boiling the nitrate with aqueous NaOH (Polis, *B* 20, 3332). White powder, insol alcohol, ether, and benzene.

Oxy cyanide $\text{Pb}(\text{C}_6\text{H}_5)_2\text{Cy}(\text{OH})$ From the nitrate in aqueous solution and KCy . White powder, insol water, alcohol, and ether.

Sulphocyanide $\text{Pb}(\text{C}_6\text{H}_5)_2(\text{SCy})_2$ Formed by adding ammonium sulphocyanide to an aqueous solution of the acetate. White powder, insol water and ordinary solvents.

Phosphate $(\text{PbPh}_2)_2(\text{PO}_4)_3$ Formed by adding sodium phosphate to an aqueous solution of the nitrate. White pp, insol. usual solvents.

Oxy-carbonate $(\text{PbPh}_2\text{OH})_2\text{CO}_3$. Ppd. as

a white insoluble powder by adding Na_2CO_3 to solutions of salts of lead diphenyl

Chromate $\text{Pb}(\text{C}_6\text{H}_5)_2\text{CrO}_4$ From the nitrate and $\text{K}_2\text{Cr}_2\text{O}_7$. Yellow pp, insol water
Sulphide $\text{Pb}(\text{C}_6\text{H}_5)_2\text{S}$ From the acetate and H_2S . Yellow prisms, sl sol alcohol, v sol benzene. Decomposes between 80° and 90°

Formate $\text{Pb}(\text{C}_6\text{H}_5)_2(\text{OHO})_2$, aq. From lead tetra phenyl and concentrated formic acid. Needles. Melts above 200° , with decomposition

Acetate $\text{Pb}(\text{C}_6\text{H}_5)_2(\text{C}_2\text{H}_3\text{O}_2)_2$, 2aq. From lead tetra phenyl and glacial HOAc (Polis, B 20, 3333). Prisms, v sol. water containing acetic acid. Melts at 193° when anhydrous

Lead tetra-*p*-tolyl $\text{Pb}(\text{C}_6\text{H}_4)_4$ [240?] S G 22 1433. Prepared by heating a mixture of 8 p.c. sodium lead alloy (300 g), *p* bromo toluene (120 g), toluene (40 g), and acetic ether (4 c.c.) for thirty hours in an oil bath. Repeated crystallisation of the product from benzene and alcohol yields lead tetra tolyl and di-*p* tolyl (A. Polis, B 21, 3424). Small needles. Decomposes at 254° . More sol. benzene, CS_2 and CHCl_3 than lead tetra phenyl. When heated with HCl at 200° it gives PbCl_2 and chloro toluene. Cold conc. HNO_3 causes decomposition with blackening, but on adding lead tetra tolyl to boiling nitric acid (S G 14) the products are lead di tolyl nitrate $\text{Pb}(\text{C}_6\text{H}_4)_2(\text{NO}_3)_2$ and nitro toluene

Lead di-*p*-tolyl salts

Chloride $\text{Pb}(\text{C}_6\text{H}_4)_2\text{Cl}_2$. Formed by passing Cl_2 into a solution of lead tetra tolyl in CS_2 . White powder, insol alcohol and ether, sl sol CHCl_3 , benzene, and CS_2 . Decomposed when heated. Converted into the corresponding nitrate by AgNO_3 in alcoholic solution

Bromide $\text{Pb}(\text{C}_6\text{H}_4)_2\text{Br}_2$. From $\text{Pb}(\text{C}_6\text{H}_4)_4$ in CS_2 and bromine. Resembles the chloride

Iodide $\text{Pb}(\text{C}_6\text{H}_4)_2\text{I}_2$. From $\text{Pb}(\text{C}_6\text{H}_4)_4$ and iodine, both dissolved in CS_2 . Yellow powder, more sol. chloroform and CS_2 than the chloride or bromide

Nitrate $\text{Pb}(\text{C}_6\text{H}_4)_2(\text{NO}_3)_2$, 2aq. Prepared as above. White powder composed of very slender needles, sol. water, containing HNO_3 , sl sol alcohol. Detonates when heated

Oxy nitrate $\text{Pb}(\text{C}_6\text{H}_4)_2(\text{OH})(\text{NO}_3)$. Formed by heating the nitrate with water, or by adding ammonia to a solution of the nitrate in dilute HNO_3 . White amorphous powder

Chromate $\text{Pb}(\text{C}_6\text{H}_4)_2\text{CrO}_4$. Insol yellow powder, ppd, by adding $\text{K}_2\text{Cr}_2\text{O}_7$ to a solution of the acetate

Sulphide $\text{Pb}(\text{C}_6\text{H}_4)_2\text{S}$. Formed by passing H_2S into a solution of the acetate. Transparent yellow plates (from alcohol benzene), sl sol alcohol and ether, v sol benzene, CS_2 , and CHCl_3 . Begins to turn brown at 90° and melts at 98° . It is decomposed by light

Formate $\text{Pb}(\text{C}_6\text{H}_4)_2(\text{CHO}_2)_2$. Formed from $\text{Pb}(\text{C}_6\text{H}_4)_4$ and formic acid. Beautiful white needles, which at 233° blacken and decompose

Acetate $\text{Pb}(\text{C}_6\text{H}_4)_2(\text{C}_2\text{H}_3\text{O}_2)_2$, 2aq. [183 5?] Formed by adding $\text{Pb}(\text{C}_6\text{H}_4)_4$ to boiling HOAc . Small white needles, more sol. dilute HOAc than the corresponding phenyl compound

LECANORIC ACID $\text{C}_{10}\text{H}_{16}\text{O}_7$. (a)-*Orsellic acid* *Diorsellic acid*. [168° cor] S 04 at

100° (Schunck) S (80 p.c. alcohol) 7° at 15° (Schunck), S, (ether) 4 at 20° (Hesse), 125 at 15° (Schunck). Discovered by Schunck in 1842 and extracted from several lichens of the genera *Lecanora*, *Varicolaria*, and *Roccella* (Schunck, A 41, 157, 54, 261, 61, 72, Roehleder & Heldt, A 48, 1, Stenhouse, A 68, 61, 70, 218, Strecker, A 68, 108, Laurent & Gerhardt, A Ch [3] 24, 310, Robiquet, A Ch 42, 236, Hesse, A 139, 22). Obtained by exhausting the lichens with ether, dissolving the greenish white crystalline residue left on evaporation in milk of lime, ppg the filtrate with H_2SO_4 , washing the pp with water, and recrystallising it from hot alcohol (Hesse). Stellate needles (containing aq). Almost insol cold water. Decomposed by heat with evolution of CO_2 . A solution of barium lecanorate is not decomposed by CO_2 . FeCl_3 colours its alcoholic solution dark purple red. An ammoniacal solution of lecanoric acid forms white pps with lead subacetate and with AgNO_3 , but the silver salt is quickly reduced. An alcoholic solution gives no pp with alcoholic solutions of lead acetate, HgCl_2 , or AgNO_3 , but with cupric acetate it gives an apple green pp. When a solution of lecanoric acid is boiled it gives orsellic acid $\text{C}_{10}\text{H}_8\text{O}_4 + \text{H}_2\text{O} = 2\text{C}_5\text{H}_4\text{O}_2$. A solution of barium lecanorate yields, on boiling, first barium orsellate, and finally orcin $\text{C}_8\text{H}_6\text{O}_4$ and BaCO_3 . An alcoholic solution yields, on boiling, orsellic ether. An ammoniacal solution acquires, on exposure to air, a splendid purple colour, through formation of orcein. Bleaching powder gives a red tint, quickly changing to brown

Salt — BaA'' small stellate needles

Di-bromo-lecanoric acid $\text{C}_{10}\text{H}_{12}\text{Br}_2\text{O}_7$ [179°] From lecanoric acid and Br in ether (Hesse). White crystals (from alcohol), insol water. FeCl_3 colours its alcoholic solution violet. Gives off CO_2 on fusion

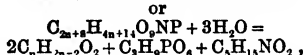
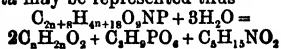
Tetra-bromo lecanoric acid $\text{C}_{10}\text{H}_8\text{Br}_4\text{O}_7$ [157°] From lecanoric acid and excess of Br in ether. Pale yellow prisms (from alcohol), v sol alcohol, ether, and alkalis. FeCl_3 colours its alcoholic solution purple. Boiling baryta-water decomposes it

LECITHIN $\text{C}_{44}\text{H}_{88}\text{O}_6\text{PN}$ (Von Lippmann, B 20, 3201), $\text{C}_{44}\text{H}_{88}\text{O}_6\text{PNaq}$ (Diaconoff) *Protagon*. Occurs in the eggs and milt or soft roe of the carp and herring, the yolk of poultry eggs, the brain of man, sheep, and domestic fowl, bile of pigs, men, and oxen, retina of oxen, blood, yeast, milk, butter, radicles of sprouting lupin seeds, maize, peas, wheat, beet root, &c (Gobley, J Ph [3] 9, 1, 83, 161, 11, 409, 12, 5, 17, 401, 18, 107, 19, 406, 21, 241, 30, 241, 33, 161; Strecker, A 123, 356, Z [2] 4, 437, Kodweiss, A 59, 261, Diaconoff, Z [2] 4, 154, Liebreich, A 134, 29, Cahn, H 5, 215, Hoppe Seyler, H 2, 427, 3, 378, J 1866, 698, 744, Hermann, Z 1866, 250, Schmidt Mulheim, J Th 1883, 166, Schulze a Barbieri, J pr [2] 27, 358, H 13, 365, Heckel a Schlagdenhauffen, C R 103, 388, Copeman a Winston, J Physiol 10, 213).

Preparation — 1 Yolk of egg is extracted with alcohol ether, and an alcoholic solution of platinum chloride is added. A compound of leorhin and platinum chloride is ppd, and this is agitated with Ag_2O , the filtrate freed from silver by H_2S and evaporated (Strecker) — 2 Brain is

exhausted with ether, and the residue extracted with absolute alcohol at 40°. The alcoholic extract is corded at 0° and filtered, and the pp of lecithin and cerebrin collected, washed with alcohol, and treated with ether. The ethereal solution of lecithin is evaporated (Diaconoff) — 3 Beet root is extracted with ether alcohol, the ether evaporated and the residue dissolved in alcohol and ppd by platinum chloride. The pp is decomposed by H_2S (Lippmann).

Properties — Hygroscopic wax-like substance, which swells up in water and is v sol alcohol, ether, chloroform, and oils. Boiling with baryta water produces glycoero-phosphoric acid, neurine, and a fatty acid (stearic, oleic, or palmitic). According to Diaconoff lecithin from brain may be separated by dissolving in alcohol and cooling to -10° , into a stearo lecithin which separates and an oleo lecithin which remains in solution. According to Strecker the decomposition by baryta may be represented thus —



thus stearo lecithin would be $C_{11}H_{20}O_8NP$, while oleo lecithin would be $C_{11}H_{20}O_8NP$ and might be written

$(C_{15}H_{33}O_2)_2C_2H_5OPO(OH)OC_2H_4N(CH_3)_2OH$. The particular lecithin examined by Strecker appeared, however, to be $C_{15}H_{33}O_8NP$ or to contain one oleyl and one palmityl radicle in place of the two oleyls in the above formula. That the neurine is not present as a base appears from the observation that lecithin is hardly attacked by dilute (10 p c) sulphuric acid (Gilson, *H* 12, 585). The neurine salt of distearyl glycoero phosphoric acid appears therefore not to be a true lecithin (v GLYCERIN). According to Lippmann the base obtained by boiling beet root lecithin with baryta is sometimes neurine and sometimes betaine. The presence of lecithin enables a solution of glucose to absorb oxygen from the air in the same way as an alkaline solution of that sugar (Nencki & Sieber, *J pr* [2] 26, 17).

Salts — The hydrochloride is a waxy mass — $(C_{15}H_{33}O_8PN)_2PtCl_6$ yellow flocculent pp v sol ether, chloroform, and benzene, but ppd from these solutions by alcohol (Strecker) — $CdCl_2$ gives a pp in a solution of lecithin in ether-alcohol.

LEDITANNIC ACID $C_{15}H_{26}O_8$. Occurs in the leaves of the marsh wild rosemary (*Ledum palustre*) (Willkg, *A* 74, 363, Rochleder & Schwartz, *A* 74, 366, *Z* 1866, 382, Thal, *J* 1883, 1401). Prepared by ppg the alcoholic decoction of the leaves with water, and the filtrate with lead acetate. The pp is dissolved in dilute HOAc, filtered, heated to boiling, and ppd with lead subacetate. The pp is suspended in water and decomposed by H_2S , and the filtrate evaporated in a current of CO_2 . Reddish powder, v sol water and alcohol. $FeCl_3$ colours its aqueous solution dark green. Lead acetate and $SnCl_4$ give yellow pps. Boiling dilute H_2SO_4 forms ledixanthin but no sugar.

Ledixanthin $C_8H_{10}O_2$ is a reddish-yellow powder, v. sol alkalis. Its alcoholic solution

gives a reddish-brown pp with lead acetate. On dry distillation it yields pyrocatechin.

LEDUM OIL. An oil obtained by steam distillation from the leaves of *ledum palustre*. According to Willkg (*Sitz W* 9, 302) it consists of a terpene and of an oxygenated oil $C_{15}H_{26}O_2$. Grassmann (*Rep Pharm* 38, 53) obtained a volatile oil (1 pt) and ledum camphor (2 pts). Fröhde (*J pr* 82, 181) found the oil to be acid and to contain a terpene (160°) and an oxygenated oil (241°). According to Trapp (*Russ Zeit Pharm* 1874, 289) the oil when recently prepared is yellowish, viscid, lighter than water, and possesses a pungent odour, when exposed to air the greater part solidifies to a crystalline mass soluble in alcohol. The solution of this solidified oil (ledum camphor) in H_2SO_4 is turned violet by HNO_3 . Ivanoff (*Russ Zeit Pharm* 1876, 577) found ledum oil to consist of a white crystalline mass saturated with oil. By repeated crystallisation from alcohol white prisms of $C_{15}H_{26}O_2$ were obtained. These were insol water, v sol alcohol, ether, chloroform, and benzene, melted at 101° , boiled at 174° , and were dextrorotatory in alcoholic solution. According to Hjelt & Collan (*B* 15, 2500) ledum camphor melts at 101° , and has a vapour density 12.33 corresponding with the formula $C_{15}H_{26}O_2$ (calc 13.02), and sublimes in long white needles. Rizza (*J R* 19, 319) gives the following properties for ledum camphor, $C_{15}H_{26}O_2$ [101°], (292° 1 V), VD 8.10, and states that Ac_2O at 150° converts it into a sesquiterpene $C_{15}H_{24}$ (264°), SG \pm 935.

LEGUMIN v PROTEINS

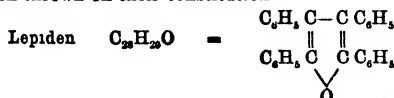
LEKENE C_2H_4 [79°] SG = 939. Forms the chief constituent of ozokerit or mineral wax, from which it is isolated by distilling the wax *in vacuo*, dissolving the distillate in benzene, and precipitating with absolute alcohol (Beilstein & Wiegand, *B* 16, 1547). White glistening crystals S (96 p c alcohol at 16°) = 0.105, S (benzene at 15°) = 8, sparingly sol in most other cold solvents, e sol hot benzene and chloroform, insol acetone. It decomposes on distillation under ordinary pressure, but can be distilled *in vacuo*. It is very stable towards oxidising agents.

LEMON OIL. Obtained by pressure from the peel of the fruit of *citrus medica*. Contains a terpene $C_{10}H_{16}$ (166°) (Blanchet & Sell, *A* 6, 281, Soubeiran & Capitaine, *A* 34, 317) or (175°) (Regnault, *J* 1863, 70, *A* 52, 171). SG 85 at 15° – 22° . The terpene is dextrorotatory, $[\alpha]_D = 109^\circ$ (Kanonnikoff, *Bn* 3, 299). According to Wallach (*A* 227, 290) oil of lemon contains pinene and hesperidene (cf Deville, *A* 71, 348, Oppenheim, *B* 5, 628). In examining the essential oil obtained by steam distillation from lemon peel, Tilden (*Ph* [3] 9, 654) found the SG to be 85.2 at 20° , and the rotation + 59° in a column 100 mm long. In this oil he found terbenzene $C_{10}H_{16}$, citrene $C_{15}H_{24}$ (constituting 70 p c of the whole), cymene (6 p c), a dextrorotatory body $C_{15}H_{26}O$ resembling terpineol (over 200°), a compound ether $C_{15}H_{24}O$ decomposed by heat into HOAc and $C_{10}H_{16}$, and $C_{15}H_{24}O$ (116°) Tilden & Dick, *C J* 57, 32). The citrene (176°) forms a terpene hydrate when treated with HNO_3 . Conc H_2SO_4 converts it into an inactive hydrocarbon (176°). Bouchardat & Lafont (*C R* 101, 383) find in essence of lemon a terpene $C_{10}H_{16}$ (178°) with a rotatory power + 105° which yields

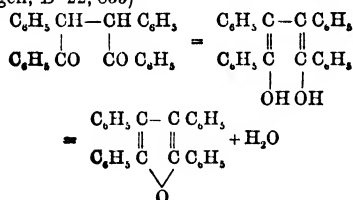
a solid inactive hydrochloride Flavitzky (*Bl* [2] 85, 171) found in a commercial sample of lemon oil ($[\alpha]_D = +60^\circ$) a levorotatory terpene (165°), S G 22 857, $[\alpha]_D = -29.8^\circ$ which formed a levorotatory hydrochloride $[\alpha]_D = -25.9^\circ$, but this was possibly an adulterant. The oil from the peel of *citrus bigamalia* contains a terpene (178°) S G 22 852 which is strongly dextro rotatory, and forms a crystalline hydrochloride $C_{10}H_{16} \cdot 2HCl$ (De Luca, *J* 1857, 481)

LEPARGYLIC ACID v AZELAIC ACID

LEPIDEN AND ITS DERIVATIVES Most of these complex compounds were first described by Zinin¹. It is only recently that light has been thrown on their constitution



(*Tetraphenylfurfuran*) (Zinin, *Z* 1867, 313, Japp a Klingemann, *B* 21, 2934 n, cf Dorn, *A* 153, 358) Formed, along with benzil and oily matters, when benzoil is heated with 1½ times its weight of conc hydrochloric acid (saturated at 8°) for 7-8 hours at 130° . The product is washed with ether, which removes the oil and benzil, after which it is recrystallised from boiling glacial acetic acid (Zinin). Also formed when bidesyl (hydro oxylepiden), or isobidesyl, is heated with conc hydrochloric acid for 2-3 hours at 130° - 140° (Maganini a Angeh, *B* 22, 855)



—Laminæ [175°] Volatilises at 220° . Insol water, sol 170 pts of boiling alcohol, 52 pts of ether at 17° , 28 pts of boiling glacial acetic acid, 8 pts of cold benzene. Boiling alcoholic potash is without action on it. Nitric acid or chromic acid oxidises it to 'acicular oxylepiden' $C_{28}H_{40}O_2$. Chlorine has the same action, but bromine converts it into dibromlepiden. With phosphorus pentachloride it yields dichlorlepiden

Chlorlepiden $C_{28}H_{38}ClO$ By treating an alcoholic solution of dichlorlepiden with sodium amalgam for 24 hours (Dorn, *A* 153, 355) — Large needles [143° - 146°], insol ether, sl sol alcohol, v sol benzene

Dichlorlepiden $C_{28}H_{36}Cl_2O$ Cannot be obtained by passing chlorine into a boiling solution of lepiden, as under these circumstances 'acicular oxylepiden' is formed (Zinin, *J R* 7, 893) By heating together equal weights of lepiden and phosphorus pentachloride (Zinin, *J R* 5, 22) — Needles [169°], sol 20 pts boiling

glacial acetic acid, 66 pts boiling alcohol (95° p c), v sol ether

A dichlorlepiden described by Dorn (*A* 153, 355), and stated by him to melt at 156° , is probably identical with the foregoing

Isodichlorlepiden $C_{28}H_{36}Cl_2O$ Formed, to gether with hydrodichloroxylepiden, by boiling an acetic acid solution of 'acicular dichloroxylepiden' [202°] (Zinin, *J R* 7, 331), or of 'sparingly soluble dichloroxylepiden', with zinc dust (Zinin, *J R* 7, 194) — Needles [166°], sol 174 pts boiling alcohol (95° p c), 12.5 pts boiling glacial acetic acid, sol ether. Oxidation converts it back into 'acicular dichloroxylepiden'

Pentachlorlepiden $C_{28}H_{34}Cl_5O$ By heating oxylepiden with excess of a mixture of phosphorus pentachloride and oxychloride for half an hour at 200° (Dorn, *A* 153, 355) — Indistinct crystals [186°], v sl sol alcohol, ether, and glacial acetic acid, v sol benzene

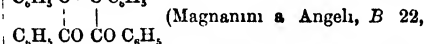
Hexachlorlepiden $C_{28}H_{32}Cl_6O$ By heating dichloroxylepiden with the foregoing mixture for 1 hour at 200° (Dorn) — Yellow amorphous substance [80° - 90°], v sol alcohol, ether, and benzene

Octochlorlepiden $C_{28}H_{30}Cl_8O$ By heating dichloroxylepiden with the foregoing mixture for 7 hours at 210° (Dorn) — Orange yellow amorphous substance [97°], v sol alcohol, ether, and benzene

Dibromlepiden $C_{28}H_{38}Br_2O$ Prepared by heating a solution of lepiden in acetic acid with bromine (Zinin, *Z* 1867, 315) Formed, along with hydrodibromoxylepiden, when 'acicular dibromoxylepiden' is treated with zinc and acetic acid (Zinin, *J R* 7, 330) — Laminæ (from acetic acid) [190°] (Zinin), [185°] (Berlin), sol 480 pts alcohol (94° p c), 44 pts boiling, and 66 pts cold, acetic acid, 50 pts ether

Oxylepiden $C_{28}H_{40}O_2$ Zinin has prepared three isomeric oxylepidens, which he distinguishes as 'acicular oxylepiden', 'tabular oxylepiden', and 'octahedral oxylepiden'

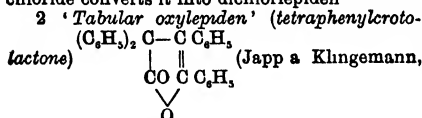
1 'Acicular oxylepiden' (dibenzoylstilbene)



$C_6H_5-CO-CO-C_6H_5$, 854, cf Japp a Klingemann, *C S Pr* 1889, 136) By oxidising lepiden with nitric acid (Zinin, *Z* 1867, 314) By treating thionessal $C_{28}H_{40}S$ (Berlin, *A* 153, 131) or toallyl sulphide $C_{28}H_{40}S_2$ (Dorn, *A* 153, 352) with hydrochloric acid and potassium chlorate By digesting benzoil with dilute sulphuric acid (Limpnich a Schwanert, *B* 4, 337) In order to prepare it, 1 pt of lepiden is suspended in 10 pts of boiling glacial acetic acid, and a mixture of 1 pt of acetic acid with 3 pts of nitric acid (S G 15) is added (Zinin, *lc*) — Yellow needles [220°], insol water, almost insol ether, sol 200 pts of boiling alcohol (94° p c) and 22 pts boiling glacial acetic acid, v sol benzene When heated to 340° it is converted into 'tabular oxylepiden' along with a little 'octahedral oxylepiden' Chromic anhydride in acetic acid solution oxidises it to dioxylepiden $C_{28}H_{40}O_3$. Zinc and acetic acid reduce it to lepiden Heating with hydriodic acid has the same effect (Dorn), also distillation with zinc dust (Berlin) In the formation of lepiden by reduction, the oxylepiden is probably first converted into hydro oxylepiden, which then parts with water, yielding lepiden

¹ Zinin's work appeared for the most part in the *Journal of the Russian Chemical Society*. The author of the present article is indebted, for his account of this portion of the subject, to Beilstein's *Handbuch der organischen Chemie*.

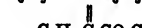
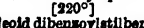
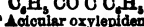
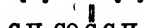
(*v supra*). Sodium amalgam, acting on the acetic acid solution, reduces it to hydroxy-lepiden $C_{28}H_{42}O_2$ (Zinnin). When heated with benzoin it is reduced to lepiden, whilst the benzoin yields benzil (Lamprecht a Schwanert, *B 4*, 388). Alcoholic ammonia at 200° converts it into a mixture of two imides of the formula $C_{28}H_{41}NO$ —one derived from 'acicular,' the other from 'tabular,' oxylepiden—whilst with methylamine it yields a methylimide $C_{28}H_{41}NO$ [161°] (Klingemann a Laycock, *private communication*). Heating with phosphorus pentachloride converts it into dichlorlepiden.



C S Pr 1889, 187) By heating 'acicular oxylepiden' to 340° and recrystallising the product, first from ether and afterwards from alcohol Tabular crystals are deposited first, then microscopic octahedra of the third modification (Zinnin, *J R* 5, 16). Tabular crystals [136°], insol water, sol 14.5 pts boiling alcohol (95 p c), and in its own weight of boiling glacial acetic acid. Dissolves readily in hot alcoholic potash solution (distinction from 'octahedral oxylepiden'), forming a salt of oxylepidenic acid $C_{28}H_{42}O_2$ (*infra*). Zinc and acetic acid are without action on it, but sodium and boiling amyl alcohol reduce it to tetraphenylbutyrolactone $C_{28}H_{42}O_2$. Alcoholic ammonia at 200° converts it into an imide $C_{28}H_{41}NO$, with alcoholic methylamine at 150° it yields the methylamide of oxylepidenic acid $C_{28}H_{41}O_2(NHCH_3)$. When heated with phosphorus pentachloride at 200° it forms chloroxy-lepiden $C_{28}H_{41}ClO_2$.

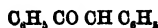
3 'Octahedral oxylepiden' Only 2 p c of this compound is formed when 'acicular oxylepiden' is heated to 340°. Most readily obtained by boiling 'acicular oxylepiden' with an alcoholic solution of caustic potash or caustic soda. Four parts of 'acicular oxylepiden' are boiled for 12–15 hours with 60 parts of alcohol (95 p c) and 3 parts of caustic soda, employing a reflux condenser. The product is washed successively with alcohol, water, and ether, and is finally recrystallised from boiling acetic acid (Zinnin, *J R* 7, 186, *J* 1875, 409). Yellowish, microscopic octahedra [232°]. Sol 76 parts boiling glacial acetic acid, almost insol alcohol. When heated to incipient boiling it is totally converted into 'tabular oxylepiden'. Boiling the acetic acid solution with zinc reduces it to hydroxy-lepiden, part of which loses water, yielding lepiden. A solution of chromic anhydride in acetic acid oxidises it to isodichlorlepiden $C_{28}H_{42}O_2$. Alcoholic potash is without action on it.

As regards the constitution of 'octahedral oxylepiden,' it is probable that this compound is a stereometric isomeride of 'acicular oxylepiden,' the latter representing the *maleoid*, the former the *fumaroid* form of dibenzoyl stilbene.



(Maleoid dibenzoylstilbene) (Fumaroid dibenzoylstilbene)

(Japp a Klingemann, *C S Pr* 1889, 188) Both yield on reduction the same hydro oxylepiden (bidesyl)



which may be converted by dehydration into lepiden.

The three oxylepidens yield by destructive distillation the same so called *isolepiden* $C_{28}H_{42}O$ (*infra*).

Chloroxy-lepiden $C_{28}H_{41}ClO_2$. By heating 'tabular oxylepiden' with a mixture of phosphorus pentachloride (1 part) and phosphorus trichloride ($\frac{1}{2}$ part) at 180°–200° (Zinnin, *J R* 5, 21—Crystals [185°], sol 22.8 parts boiling glacial acetic acid.

Dichloroxy-lepiden $C_{28}H_{41}Cl_2O_2$. 'Acicular dichloroxy-lepiden' is obtained by heating lepiden with four times its weight of phosphorus pentachloride and treating the product with water (Zinnin, *J R* 5, 23). By the oxidation of dichlorlepiden [169°] (Zinnin, *J R* 7, 332, *J* 1876, 426)—Needles [202°], sol 146 parts cold, 13.7 parts boiling acetic acid, 90 parts boiling ether. When boiled with zinc and acetic acid it yields a mixture of hydrodichloroxy-lepiden and isodichlorlepiden [166°].

When the foregoing 'acicular dichloroxy-lepiden' is heated to near its boiling point it is converted into a mixture of two new isomeric dichloroxy-lepidens, which may be separated by the difference in their solubility in ether. 'Readily soluble dichloroxy-lepiden,' which forms the chief product, is very soluble in alcohol, ether, and acetic acid, and is deposited from these solutions as a soft resin. Alcoholic caustic potash converts it into a salt of dichloroxy-lepidenic acid $C_{28}H_{42}Cl_2O_2$. The other isomeride, 'sparingly soluble dichloroxy-lepiden,' is best prepared by boiling 4 parts of 'acicular dichloroxy-lepiden' with three parts of caustic soda and 40 parts of alcohol for from 20 to 24 hours. It forms a granular powder consisting of microscopic prisms [230°], almost insol alcohol and ether, sol 36 parts boiling glacial acetic acid. Boiling with acetic acid and zinc converts it into isodichlorlepiden [166°] (Zinnin, *J R* 7, 191).

These three compounds, as their mode of formation and reactions denote, are to be regarded as dichlor substitution products of 'acicular,' 'tabular,' and 'octahedral' oxylepiden respectively.

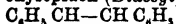
A fourth *dichloroxy-lepiden* has been obtained by Dorn (*A* 153, 353) by treating *dichlorthionessal* $C_{28}H_{41}Cl_2S$ with hydrochloric acid and potassium chlorate—Small needles [178°], readily sol alcohol, benzene, and glacial acetic acid. Not attacked by alcoholic potash at 150°. Zinc and acetic acid convert it into a dichlorlepiden [156°] (*v supra*), whilst with hydriodic acid at 100° it yields dichlorlepiden, lepiden, and oxylepiden (?).

Dibromoxy-lepiden $C_{28}H_{41}Br_2O_2$. Three dibromoxy-lepidens, corresponding with the first three of the foregoing dichloroxy-lepidens, have also been described by Zinnin (*J R* 7, 329, *J* 1876, 425). In their modes of preparation, properties, and reactions, they resemble the chlorine compounds. 'Acicular dibromoxy-lepiden' is obtained either by brominating 'acicular oxylepi-

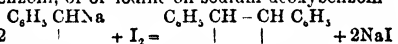
den' or by oxidising dibromolepiden in acetic acid solution with nitric acid. Needles [222°], sol 40 parts boiling acetic acid. Heated above its melting point it yields two isomeric dibromoxyepidenes—'readily soluble dibromoxyepiden', which is resinous, v sol alcohol, ether, and acetic acid, and is converted by alcoholic caustic potash into a salt of dibromoxyepidenic acid $C_{22}H_{22}Br_2O_2$, and 'sparingly soluble dibromoxyepiden'. This latter is more easily obtained by boiling the acicular modification with a quantity of alcoholic potash insufficient to dissolve it. Crystallises from alcohol in lemon yellow rhombic tables [239°], almost insol ether, sol 66 parts boiling acetic acid and 1,000 parts boiling alcohol (95 p c). Heated above its melting point it is converted into 'readily soluble dibromoxyepiden' (Zinnin, *J R* 7, 329, *J* 1876, 425).

Both 'acicular' and 'sparingly soluble' dibromoxyepiden, when reduced with zinc and acetic acid, yield the same hydrodibromoxyepiden (dibromidesyl), part of which undergoes dehydration, forming dibromlepiden [190°].

Hydroxyepiden (*Bidesyl*)

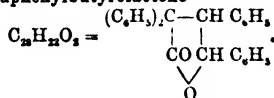


$C_{22}H_{22}O_2 =$ $C_6H_5 \cdot CO \quad CO \cdot C_6H_5$ (Magnanini & Angeli, *B* 22, 853) Formed, along with lepiden, by the action of zinc and acetic acid on 'octahedral oxyepiden', or of sodium amalgam on 'acicular oxyepiden'. The product of the reaction is ppt with water and treated with ether, which dissolves only the lepiden (Zinnin, *J R* 7, 188, *J* 1875, 409). By the action of desyl bromide, $C_6H_5 \cdot CHBr \cdot CO \cdot C_6H_5$, on sodium deoxybenzoin, or of iodine on sodium deoxybenzoin



$C_6H_5 \cdot CO \quad CO \cdot C_6H_5$ (Knovenagel, *B* 21, 1358) —Long flat needles (from acetic acid), [251°] (Zinnin), [254°–255°] (Knovenagel, also Magnanini & Angeli). Almost insol cold alcohol and ether, sol 112 pts boiling acetic acid, sol hot benzene. Heated with hydrochloric acid at 130°–140° it parts with water, yielding lepiden (Magnanini & Angeli, *B* 22, 853).

Tetraphenylbutyrolactone



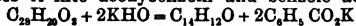
By treating a solution of 'tabular oxyepiden' (tetraphenylbutyrolactone) in boiling amyl alcohol with sodium (Klingemann & Laycock, *private communication*) —Microscopic, white needles [221°], sol boiling, sl sol cold, alcohol.

Hydrodichloroxyepiden $C_{22}H_{22}Cl_2O_2$. Formed, along with dichlorlepiden, when 'sparingly soluble dichloroxyepiden' is boiled for a long time with zinc and 'acetic acid, or, more readily, by boiling an alcoholic solution of 'acicular dichloroxyepiden' with sodium amalgam, keeping the liquid acid with acetic acid (Zinnin, *J R* 7, 195, *J* 1875, 413). —Flat needles (from acetic acid) [261°], insol alcohol and ether, sol 205 pts boiling acetic acid.

Hydrodibromoxyepiden $C_{22}H_{22}Br_2O_2$. By the action of zinc and acetic acid on 'acicular' or 'sparingly soluble' dibromoxyepiden (Zinnin,

J R 7, 330, *J* 1876, 425) —Slender needles, v sl sol alcohol and ether, sol 172 pts boiling acetic acid.

Dioxyepiden $C_{22}H_{22}O_2$. By warming 25 pts of 'acicular oxyepiden', suspended in 20 pts of acetic acid, with a solution of 12–15 pts of chromic anhydride dissolved in 150 pts of acetic acid (Zinnin, *Z* 1871, 483) —Rhombic tables [157°], sol 24 pts boiling alcohol (95°). Reducing agents are without action on it. By further action of chromic anhydride it yields benzil and benzoic acid. Alcoholic potash hydrolyses it into deoxybenzoin and benzoic acid.

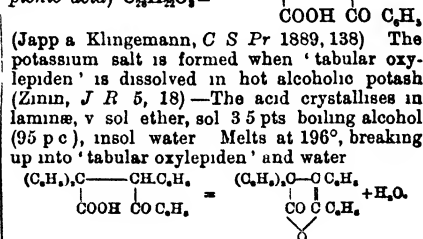


Isodioxyepiden $C_{22}H_{22}O_2$. By boiling 'octahedral oxyepiden' with a solution of chromic anhydride in acetic acid (Zinnin, *J R* 7, 190, *J* 1875, 410) —Laminæ [164°], sol 10 pts boiling alcohol, less sol ether, sol 4 pts boiling acetic acid. Boiling alcoholic potash is without action on it (distinction from foregoing isomeride).

Oxyepidenimide $C_{22}H_{21}NO$. By heating 'acicular oxyepiden' with alcoholic ammonia for 5 hours at 200°, it yields a mixture of oxyepidenimide and the isomeric tetraphenylpyrrolone. The oxyepidenimide is separated by means of its greater solubility in alcohol (Klingemann & Laycock, *private communication*) —Yellow prisms (from benzene) [180°–182°]. Heated to 310° it changes into tetraphenylpyrrolone (*v infra*).

Oxyepiden methylimide $C_{22}H_{21}O(NCH_3)$. By heating 'acicular oxyepiden' with an alcoholic solution of methylamine for 2 hours at 200° (K. & L.) —Small pale yellow plates [161°] from alcohol, well shaped crystals from carbon bisulphide. V sol benzene, readily sol carbon bisulphide, moderately sol boiling alcohol.

Oxyepidenic acid (*Benzoyl triphenyl pro*)
 $(C_6H_5)_3C - CH \cdot C_6H_5$
ponic acid) $C_{22}H_{22}O_2 =$

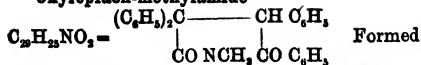


The so called *isoxylepidenic acid* described by Zinnin (*J* 1877, 397) is identical with the foregoing (Japp & Klingemann, *C S Pr* 1889, 139).

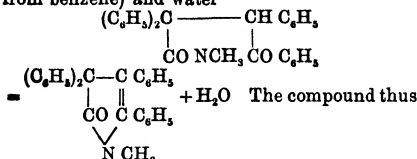
Tetraphenylpyrrolone
 $(C_6H_5)_3C - C \cdot C_6H_5$
 $C_{22}H_{21}NO =$ $\begin{array}{c} || \\ CO \quad C_6H_5 \\ | \\ NH \end{array}$ Formed, along

with the isomeric oxyepidenimide, when 'acicular oxyepiden' is heated with alcoholic ammonia at 200°. By heating oxyepidenimide to 310° as sole product when 'tabular oxyepiden' is heated with alcoholic ammonia at 200° (Klingemann & Laycock) —Small pale yellow plates [208°], sparingly sol alcohol.

Oxylepiden-methylamide



when 'tabular oxylepiden' is heated with an alcoholic solution of methylamine at 150° $\text{C}_{22}\text{H}_{22}\text{O}_2 + \text{NH}_4\text{CH}_3 = \text{C}_{22}\text{H}_{22}\text{NO}_2$ (Klingemann a Laycock) — Yellowish laminae [287°] from acetic acid, short needles from alcohol. Sol boiling alcohol, more sol boiling acetic acid. Decomposed by distillation *in vacuo* into methyl-tetraphenylpyrrolone (prismatic crystals [158°] from benzene) and water

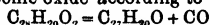


formed appears to be isomeric, not identical, with oxylepiden-methylamide (K a L)

Dichloroxyepidenic acid $\text{C}_{22}\text{H}_{20}\text{Cl}_2\text{O}_2$. By dissolving 'readily soluble dichloroxyepiden' in boiling alcoholic potash (Zinn, *J R* 7, 191, *J* 1875, 411) — Rhombic laminae [182°], from acetic acid. Sol 16 pts boiling acetic acid. Heated to 200° it parts with 1 mol water, and is reconverted into 'readily soluble dichloroxyepiden'.

Dibromoxyepidenic acid $\text{C}_{22}\text{H}_{20}\text{Br}_2\text{O}_2$. By dissolving 'readily soluble dibromoxyepiden' in boiling alcoholic potash (Zinn, *J R* 7, 330, *J* 1876, 425) — Six sided laminae from acetic acid.

So called isolepiden and its derivatives — Under the name of *isolepiden*, Zinn (*J R* 5, 20) has described a compound which he obtained by the destructive distillation of oxylepiden, and which he regarded as isomeric with lepiden. Japp a Klingemann have, however, shown (*C S Pr* 1889, 139) that this compound has the formula $\text{C}_{27}\text{H}_{20}\text{O}$, instead of $\text{C}_{22}\text{H}_{20}\text{O}$, and that it is formed from oxylepiden with elimination of carbonic oxide according to the equation



The name 'isolepiden' is consequently inappropriate, but will be retained here, as the constitution of the compound is unknown, and a systematic name cannot be given. The formulae of the various hydro 'isolepidens' and oxy 'isolepidens' described by Zinn must also be written with 27 instead of with 22 atoms of carbon, although these compounds have not yet been re-investigated.

Isolepiden $\text{C}_{27}\text{H}_{20}\text{O}$ (Japp a Klingemann, *lc*) By the destructive distillation of oxylepiden (Zinn, *J R* 5, 20, cf preceding paragraph) The three oxylepidens all yield this compound on distillation, but in reality it is only 'tabular oxylepiden' which yields it, as the other two oxylepidens are previously transformed into 'tabular oxylepiden' at the temperature of the reaction. The distillate is washed with ether and then recrystallised, first from alcoholic potash, afterwards from alcohol. The ether extracts a little 'tabular oxylepiden,' which is carried over undecomposed (Japp a Klingemann, *C S Pr* 1889, 139) — Isolepiden forms yellow tabular crystals [150°]. Decomposes on redistillation. Sol 18 pts boiling alcohol (95 p c), and in 2 pts boiling acetic acid (Zinn, *J* 1877,

394). Not attacked by alcoholic potash. It is reduced by zinc dust and acetic acid to dihydro isolepiden $\text{C}_{27}\text{H}_{22}\text{O}$, and by sodium amalgam to tetrahydro isolepiden $\text{C}_{27}\text{H}_{24}\text{O}$. Oxidising agents convert it into oxy isolepiden, and ultimately into a mixture of benzophenone and benzil.

Dihydro-isolepiden $\text{C}_{27}\text{H}_{22}\text{O}$. By reducing a solution of isolepiden in from 5 to 6 pts of acetic acid with zinc dust. The product is poured into water, and the pp is washed with ether and recrystallised from alcohol (Zinn, *J* 1877, 394) — Small rectangular prisms [182°]. Sol 12.5 pts boiling alcohol (95 p c), v sl sol ether.

Tetrahydro isolepiden $\text{C}_{27}\text{H}_{24}\text{O}$. By the action of sodium amalgam on a boiling alcoholic solution of isolepiden (Zinn). Separates from ether as a soft resinous mass which soon becomes crystalline [182°]. Readily sol. alcohol and acetic acid, less sol ether. Chromic anhydride oxidises it in acetic acid solution, even in the cold, to dihydro isolepiden.

Oxyisolepiden $\text{C}_{27}\text{H}_{20}\text{O}_2$. Three pts of isolepiden are dissolved in 40 pts of acetic acid, and oxidised with a solution of 3 pts of chromic anhydride in 30 pts of acetic acid (Zinn, *J* 1877, 395) — Short slender needles [161°]. Sol 40 pts boiling, and 600 pts cold alcohol, and in 4 pts boiling acetic acid. Boiling alcoholic potash is without action on it. Zinc and acetic acid reduce it to dihydro isolepiden. Excess of chromic acid oxidises it, in acetic acid solution, to benzophenone as chief product, together with benzil and benzoic acid.

Oxy isolepiden, like oxylepiden, may be converted into isomeric compounds. Thus, when it is boiled with a quantity of alcohol or alcoholic potash insufficient to dissolve it, it is transformed into *cuneiform oxy isolepiden* [162°]. When this is heated above its melting point, or when ordinary oxy isolepiden is distilled, *tabular oxy-isolepiden*, a third isomeride, is formed. Rhombic tables [152.5°] from acetic acid. Sol 13.5 pts boiling acetic acid, 80 pts boiling alcohol. F R J

LEPIDINE v (Py 1) MFTHYL QUINOLINE

LEUCANILINE is described under TRIAMIDO DI PHENYL TOLYL METHANE.

Para-leucaniline is described as TRIAMIDO-TRI PHENYL METHANE.

LEUCAURIN v TRI OXY TRI PHENYL MFTHANIC

LEUCATROPIC ACID $\text{C}_{17}\text{H}_{12}\text{O}_3$ [74°]. Occurs in belladonna (Kunz, *Ar Ph* [3] 23, 722). Minute satiny needles. Insol cold, m sol boiling water, and alcohol.

LEUCINES v PROTEIDS

LEUCIC ACID Described as OXY HEXOIC ACID.

LEUCINDIGO v INDIGO

LEUCINE $\text{C}_6\text{H}_7\text{NO}_2$ i e

$\text{CH}_3\text{CH}_2\text{CH}_2\text{CH}(\text{NH}_2)\text{CO}_2\text{H}$ a *Amido n-hexoic acid*. Mol w 131 [170°] (Schwanert) S G 12 1 293 (Engel a Vilmain, *Bl* [2] 24, 279) S 22 at 18° (Schulze, *H* 9, 254), 37 in the cold (Zollkofer). S (alcohol of S G 828) 152 in the cold (Mulder). S (96 p c alcohol) 096 in the cold (K). S (98 p c hot alcohol) 125 (K). $[\alpha]_D = +14.1^\circ$ in a 15 p c HCl solution, $+5.6^\circ$ in a 25 p c solution in NH_4Aq (Reese, *A* 242, 11). Discovered by Proust (*A Ch* [3] 10, 40) as a product of the putrefaction of cheese, and called 'caseous oxide'. Braconnot

(*A Ch* [2] 18, 119) found it among the products of the action of H_2SO_4 on animal substances Mulder (*J pr* 16, 290) showed the two substances thus obtained were identical

Occurrences—In old cheese (Proust), in fresh calf's liver (Liebig, *Chem Briefe*, ed 3, 453), in diseased but not in healthy human liver (Frerichs & Stadel, *J* 1854, 675, 1856, 702, 1858, 550), in the tissue of the lungs (Cloetta, *A* 92, 289), in the thymoid and thyroid glands, and in the pancreas (Gorup-Besanez, *A* 98, 7, Radziejewsky, *Z* 1866, 416), in the brain of oxen (W Muller, *A* 103, 131), in the pancreas of oxen (Scherer, *J* 1859, 610), in the liver and urine in cases of typhus, smallpox, leucæmia, and poisoning by phosphorus (Salkowsky, *J Th* 1880, 457, Valentiner, *J* 1854, 675, Sotnitschewsky, *H* 3, 391), in the stomachs and intestines of the pupæ of butterflies (Schwarzenbach, *J* 1857, 538), in *Agaricus muscarius* (Ludwig, *J* 1862, 516), in the juice of vetches germinated in the dark (Gorup Besanez, *B* 7, 146, cf Cossa, *G* 5, 314), in *chenopodium album* in young pumpkin plants (Schulze a Barbieri, *B* 11, 1233), and in beet root molasses (Lippmann, *B* 17, 2837)

Formation—1 In the putrefaction of proteids and gelatin (Bopp, *A* 69, 20) and by boiling these substances with dilute H_2SO_4 , or by fusing them with potash (Hinterberger, *Sitz W* 9, 450, *A* 71, 75, Zollikofer, *A* 82, 174, Gossmann, *A* 91, 130, Leyer a Koller, *A* 83, 332, Schlossberger, *Z* 1860, 424, Erlenmeyer a Schoffer, *Z* 1859, 315, Hochstetter, *J pr* 29, 36, Ritt hausen a Kreuzler, *J pr* [2] 3, 307)—2 In the pancreatic digestion of gelatin (Nencki, *B* 7, 1593, Jeanneret, *J pr* [2] 15, 353)—3 By acting on a bromo hexoic acid with ammonia (Hufner, *J pr* [2] 1, 6, *Z* [2] 4, 391, 616)

Preparation—Horn shavings (2 lbs) are boiled with H_2SO_4 (5 lbs) and water (13 lbs) for 24 hours with inverted condenser The product is mixed with lime, filtered, and evaporated to a smaller bulk (12 lbs) Oxalic acid is then added to acid reaction, and the liquid filtered and evaporated till a crystalline film forms on the surface Leucine mixed with tyrosine is deposited in groups of yellowish laminae On recrystallisation from water tyrosine is deposited first, and the mother liquor is then decoloured by animal charcoal and evaporated The leucine is recrystallised from alcohol (Schwanert, *A* 102, 221, cf Hinterberger, *A* 71, 72 Waage, *A* 118, 295) Leucine may be detected and isolated by means of its sparingly soluble copper salt ($C_6H_{11}NO_2$)₂Cu, although the precipitation of this salt is hindered by free acids and by some organic bodies (Hoffmeister, *Sitz W* 75, 469)

Properties—Soft nacreous scales (from alcohol) resembling cholesterol It may be sublimed (Mulder) Decomposed on fusion, giving amylinine, CO_2 , and NH_3 Lævotatory (Lewkowitch, *B* 17, 1439, cf Mauthner, *H* 7, 223) Sol water and alcohol, insol ether Its solubility in water is increased by the presence of acetic acid or an alkaline acetate When heated with baryta water at 160° it becomes inactive. The inactive leucine is changed to an active variety, lævotatory when dissolved in aqueous HCl, by the action of *Penicillium glaucum* (Schulze a Bosshard, *B* 18, 388).

Reactions—1 An alkaline solution exposed to the action of ozone yields CO_2 , butyric acid, and NH_3 (Gorup-Besanez, *A* 125, 210)—2 Chlorine passed into water in which leucine is suspended forms CO_2 and valerionitrile, as well as chloro valerionitrile (Schwanert) Chlorine passed into an alkaline solution of leucine forms oxy-hexoic acid—3 Nitrous acid converts it into a oxy hexoic (leucic) acid—4 Distillation with MnO_2 and dilute H_2SO_4 yields CO_2 and valerionitrile—5 Distillation with water and PbO_2 yields butyric aldehyde and NH_3 (Liebig, *A* 70, 313)—6 Aqueous $KMnO_4$ yields NH_3 , oxalic acid, and valeric acid (Neubauer, *A* 106, 59)—7 Potash fusion gives NH_3 , hydrogen, and potassium valerate (Liebig, *A* 57, 127)—8 When heated with fuming HNO_3 at 140° it gives hexoic acid and ammonia (Hufner)—9 With KOH (2 mols) and MeI (3 mols) it forms potassium di methyl amido-hexoate methyl iodide $C_6H_{11}(NMe_2)CO_2K$, which, when heated with moist Ag_2O , yields methylamine, a salt $C_6H_{11}O_2K$, and potassium leucinate $C_6H_{11}O_2K$ (Körner a Menozzi, *G* 13, 353)—10 Leucine gives off more nitrogen when its solution is decomposed by $NaBrO$ in presence of NH_3 , than when the NH_3 is absent (E Schulze, *J pr* [2] 31, 236)

Salts— $HA \cdot HCl$ crystals, v sol water (Laurent a Gerhardt, *A Ch* [3] 24, 321, *A* 68, 365)— $(HA)_2 \cdot HCl$ laminae (Schwanert)— $(HA)_2 \cdot H_2PtCl_6$ yellow crystalline pp— $HA \cdot HNO_2$ colourless needles, v e sol water— CuA_2 pale blue scales S 033 in the cold, 07 at 100° (Hoffmeister)— $HgA_2 \cdot aq$ laminae Mercurio nitrate gives a white flocculent pp in a solution of leucine (R Hoffmann, *A* 87, 183)— $PbA_2 \cdot aq$ ppd by adding NH_3 aq to an aqueous solution of leucine and lead acetate (Strecker, *A* 72, 89)

Benzoyl derivative $C_6H_5 \cdot BzNO_2$ i.e. $C_6H_5(NHBz)CO \cdot H$ From leucine and $BzCl$ at 100° (Destrem, *Bl* [2] 30, 481) Granules, sol alcohol and ether In the preparation of leucine anhydride there is also formed the anhydride ($C_6H_5(NHBz)CO$)₂O, which is an amorphous body [85°] insol water and ether, v e sol alcohol Decomposed by boiling water into benzoic acid and the anhydride of leucine

Phthaloxyl derivative $C_6H_5(CO \cdot H)NH \cdot CO \cdot C_6H_5$ [132°] From leucine, alcoholic KOH , and phthalyl chloride (Reese, *B* 21, 277)— K_2A'' concentric groups of small slender needles

Anhydride (C_6H_5NO)₂O When the product of the action of $BzCl$ on leucine at 100° is treated with alcohol leucine anhydride remains undissolved, while its di benzoyl derivative (v supra) passes into solution (Destrem, *C R* 86, 484) Leucine anhydride is a white amorphous body, insol alcohol, but becoming gelatinous when boiled therewith It is not easily converted into leucine by boiling water

LEUCINDIN SULPHONIC ACID v INDIN

LEUCINIMIDE $C_6H_{11}NO$ Obtained in small quantity, together with leucine and tyrosine, by boiling proteids with dilute H_2SO_4 (Limpricht a Hesse, *A* 116, 201, Erlenmeyer, *A* 119, 17, Thudichum, *C J* 23, 409) Formed also when proteids are treated with bromine water (Hlasiwetz a Hebermann, *A* 159, 328), and, together with benzoyl-leucine, by heating leucine with $HOBr$ at 200° (Destrem, *Bl* [2] 30, 481). Mi-

amorphous, sl sol water— $\text{Ba}_2(\text{C}_2\text{H}_3\text{O}_2)_2$, flocculent pp— $\text{Pb}_2(\text{C}_2\text{H}_3\text{O}_2)_2$, pp— $\text{Ag}_2\text{C}_2\text{H}_3\text{O}_2$ (W)

Penta oxim $\text{C}_5(\text{NOH})_5$, Yellow crystalline solid Formed by the action of hydroxylamine upon croconic or leuconic acid. Decomposes suddenly at 172° — $\text{C}_5\text{H}_4\text{N}_5\text{O}_5\text{K}_2$, very explosive The penta oxim can be reduced by

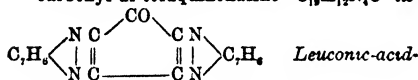
SnCl_2 to $\text{CH}(\text{NH}_2)$ $\begin{matrix} \text{C}(\text{NH}_2) & \text{C}(\text{NH}_2) \\ | & | \\ \text{C}(\text{NH}_2) & \text{C}(\text{NH}_2) \end{matrix}$ which forms crystalline salts

Tetra acetyl derivative of the penta-oxim $\text{C}_5(\text{NOH})(\text{NOAc})_4$, sq (from benzene) From the penta oxim and Ac_2O at 45° (Nietzki a Rosemann, *B* 22, 916) Plates (from Ac_2O) or needles containing chloroform (from chloroform) Sl sol hot benzene Decomposed at 100°

Tetra-oxim $\text{C}_4\text{H}_4\text{N}_4\text{O}_4$, s

$\text{CO} \begin{matrix} \text{C}(\text{NOH}) & \text{C}(\text{NOH}) \\ | & | \\ \text{C}(\text{NOH}) & \text{C}(\text{NOH}) \end{matrix}$ When potassium croconate (30 g) is gradually added to a cooled mixture of HNO_3 (45 g of S G 1.39) and water (40 g), and the product is diluted with water (500 cc) and heated with hydroxylamine hydrochloride (180 g) for some hours at 45° and finally at 100° there is formed a pp of mixed oxims This is dissolved in aqueous Na_2CO_3 and CO is passed in, whereupon the penta oxim is ppd while the tetra oxim can be ppd from the filtrate by HCl Yellow pp Explodes at 160° Its sodium derivative is v sol water, but is ppd on adding alcohol or NaCl , apparently in the form of $\text{Na}_2\text{C}_4\text{H}_4\text{N}_4\text{O}_4$

Carbonyl-di-toluquinoxaline $\text{C}_{18}\text{H}_{12}\text{N}_4\text{O}$ s



di tolylene-o diamide [above 300°] Formed by adding a salt of tolylene-o diamine to a cold aqueous solution of leuconic acid Golden yellow needles V sol warm chloroform, sl sol hot alcohol and acetic acid, insol water Weak base

Phenyl hydrazide $(\text{C}_6\text{H}_5\text{N})_2\text{C}_2\text{C}_6\text{H}_5\text{HPh}$ Red needles (from acetic acid), v sol chloroform, sl sol alcohol Weak base The hydrochloride forms a dark-green pp (Nietzki a Ben kiser, *B* 19, 776)

LEUCOTIN v COTO BARR

LEUCOTURIC ACID $\text{C}_8\text{H}_8\text{N}_2\text{O}_6$ Oxalantin

Formed when a solution of alloxanic acid is rapidly boiled down to a syrup, cold water is then added which leaves the leucoturic acid undissolved (Schlieper, *A* 56, 1) Formed also by reducing parabanic acid with zinc and HClAq in the cold (Limpricht, *A* 111, 134) White crystalline powder, insol cold, m sol hot, water Decomposes alkaline carbonates on boiling Decomposed by heating with KOH , giving off NH_3 , and forming oxaluric acid Its ammoniacal solution is ppd by AgNO_3 , and on boiling reduction takes place Boiling conc HNO_3 does not attack it

LEVONIC ACID $\text{C}_8\text{H}_8\text{O}_8$ 3aq An acid said by Wiederhold (*C* 1884, 971) to be obtained by boiling levulose with baryta water Yellowish brown powder, v sol alcohol and water Decomposed by heat.

LICARENE $\text{C}_{10}\text{H}_{16}$ (168° – 172°) S G 1.835 An inactive terpene obtained by the action of ZnCl_2 or P_2O_5 on the essential oil of *Lucaria Kanak* (Morin, *A Ch* [5] 25, 427) Conc HClAq forms $\text{C}_{10}\text{H}_{16}\cdot 2\text{HCl}$, a colourless liquid, S G 1.1069, inactive to light, and decomposed on distillation into hcareene and hydrochloric acid The essential oil $\text{C}_{10}\text{H}_{16}\text{O}$ might be looked upon as hcareene hydrate and exhibits the following properties—(198°) at 755 mm S G 1.568, $[\alpha]_D = -19$ at 15° It is sol alcohol, ether, and glycerin

LICHENINE $\text{C}_8\text{H}_{10}\text{O}$, Occurs in Iceland moss

Preparation—*Cetraria islandica* or other similar lichen is heated for several hours with a 2 p c solution of K_2CO_3 The aqueous solution is ppd by alcohol (Honig a Schubert, *M* 8, 460, cf Knop a Schnedermann, *A* 55, 164, Maschke, *J pr* 61, 7, Davidson, *N Ed P J* 28, 260, Errera, *Bn* 1, 882, Bauer, *J pr* [2] 34, 49, Klason, *B* 19, 2541)

Properties—Gelatinous pp, v sl sol cold water Boiling water forms an opalescent solution, reppd on cooling or on adding alcohol Gives no blue colour with I and H_2O_2 Dissolves in HCl , from which solution it is ppd in snow-white flocks by alcohol Weak hot acids convert it after some time into a dextro rotatory sugar $[\alpha]_D = +55^\circ$ Glucose is also formed Two carbohydrates, resembling soluble starch in their properties, are present in the aqueous solution

LICHENO-STEARIC ACID $\text{C}_{18}\text{H}_{34}\text{O}_2$ [c 120°] An acid occurring in Iceland moss (Schnedermann a Knop, *A* 55, 149), and probably also in the fly agaric or toadstool (Bolley, *A* 86, 50) Obtained by boiling Iceland moss for 15 minutes with dilute alcohol and some K CO_3 , filtering, adding HClAq and 4 volumes of water The pp is boiled with alcohol of 45 p c, whence a mixture of lichenostearic and cetraric acids separates on cooling The lichenostearic acid is extracted from the mass by petroleum and recrystallised from alcohol Mass of radiating needles which soon change to delicate pearly laminae Has a rancid taste Insol water, v sol alcohol, ether, and oils Is not attacked by AcCl On oxidation it yields CO_2 and decolic acid (Hilger a Buchner, *B* 23, 461) The ammonium salt forms a jelly containing extremely slender needles— BaA' , greivish white pp which cakes together in boiling water— PbA' , flocculent pp fusing under water— AgA'

LIEBERMANN'S REACTION A blue or green colour obtained by warming phenol with H_2SO_4 containing nitrous acid It may be used as a test for phenol or for nitrous acid Various derivatives of phenol may be used instead of phenol, while nitroso and oximido compounds usually react like nitrous acid

LIGHT For an account of the applications of optical methods of inquiry to chemical problems, v PHYSICAL METHODS, section OPTICAL

LIGNIFICATION v LIGNONE

LIGNO-CERIC ACID $\text{C}_{18}\text{H}_{16}\text{O}$ or $\text{C}_{22}\text{H}_{18}$, CO H [81°] Occurs in the paraffin obtained from beech wood tar (Hell, *B* 13, 1709) Occurs also in the product of the saponification of earth-nut oil, (Kreiling, *B* 21, 880) Colourless felted needles or plates (from alcohol)

Salts— A'Na white powder— AK white

powder, sol alcohol—A'Ag white, pp [c 155°]
—A'Cu green powder, sol ho. benzene—
A'Pb white powder, [117°] v sol hot benzene

Methyl ether A'Me [58°] White glistening plates Sol CS₂, CHCl₃, C₆H₆, ether, and ligron, sl sol alcohol Distils undecomposed at a high temperature

Ethyl ether A'Et [55°], (310° at 20 mm, 860° at 760° mm) Glistening plates

Chloride C₂H₅COCl [48°–50°] Plates
Sol ether

LIGNONE (*Lignum, Lignose, Lignification*)
Lignification, or the process of wood formation, is one of the principal of the modifications of the cell-wall, by which it and the tissues containing it are differentiated for fulfilling their several functions. The history of a lignified cell, or rather of the substance of the cell wall, is usually stated to consist of (1) the elaboration of the primary cell wall from materials in protoplasm, and formed at its limiting film as an envelope of pure cellulose, (2) the induration of this cell-wall, *ie* lignification, by the infiltration of substances, which when deposited are known as 'lignin,' or more vaguely as 'incrustative' substance. Lignification is recognised by the morphological changes with which it is associated, as well as by the very characteristic reactions of the product (Goodale, *Phys Botany*, 1885). These products, although presenting a wide range of differences, corresponding with variations in structural and other characteristics of the tissues which they compose, fall naturally, and as regards their chief constituent, into a homogeneous chemical group, designated by the term lignocellulose, of which the typical features are represented in the substance of the jute fibre (vol 1 p 719).

It is the purpose of this article to give a brief account of more recent researches into the constitution of this typical lignocellulose, and to show the bearings of the results arrived at upon the chemistry of the woods, the most numerous and important section of the group. The advantages of the jute fibre over the latter as a subject of study are that it is a simple tissue, whereas the woods are complex aggregates, that as an isolated fibred it is much more easy of penetration by reagents, and that as a product of only a few months' growth it has not undergone such secondary changes in composition as necessarily take place in the substance of a perennial stem or true wood. Such characteristics mark out the jute fibre substance as a natural basis for the general solution of the chemical problem of lignification.

Evidence has already been given (vol 1 *loc cit*) for regarding this product as chemically homogeneous, which view is further developed in the investigations referred to (*C J* 1889 199). In all reactions in which hydroxyl groups only are brought into play—hydrolysis and etherification—it behaves as an integral compound (lignocellulose), of which the formula C₁₂H₁₀O₅ is an approximate empirical expression. The limit of 'nitration' is the tetranitrate, indicating a number of OH groups in the original less by two than in the molecule of cellulose, similarly represented, *ie* as C₁₂H₂₀O₁₀. The product is of a bright gold colour, with a silky lustre. In further contradistinction to

cellulose, the OH groups of the lignocellulose react with acetic anhydride at its boiling point. Its reaction with chlorine has been studied quantitatively, 1 gm of the purified fibre substance takes up 54–55 c c Cl (calc at 0° and 760 mm), or approximately 16 p c by weight the quantity of Cl as HCl formed being equal to that entering the molecule. The chlorinated product *n*(C₁₂H₁₀Cl₂O₅),¹ a simple substitution-derivative of the non-cellulose constituent (which we may regard as *n*(C₁₀H₈Cl₂O₄)), contains mairgallol in combination with a body not yet completely studied, but yielding furfural on hydrolysis. In addition to the molecular groupings thus indicated, the presence of methyl groups is proved by the formation of methyl chloride on heating this compound, and of acetic acid on destructive distillation of the fibre substance, as also by dissolving it in sulphuric acid, diluting and distilling. These results afford a general view of the constitution of the 'lignin' substance, as it has been termed hitherto. We may regard it as containing closed C₆ chains, further characterised by the presence of ketone or quinone oxygen, and united to the furfural yielding body, which is probably related to 'wood gum' ('Holzgummi'). Tollens has, in fact, isolated xylose from the jute fibre (*B* 22, 1046), though in very small quantity, and we would note here the increase in the proportion of the parent substance, wood-gum, with the more advanced lignification of perennial stems, as additional evidence for this view. It is difficult to localise the methyl groups, but they would appear to be independent of the above, and may be regarded as existing in an acetic residue in combination. For a substance of marked ketonic and aldehydic characteristics, the terms *lignum* and *lignose* are obviously unsuited, and should give place, according to present views, to *lignone*, by which, therefore, we shall designate the non cellulose component of the lignocelluloses generally.

The lignone of plant cells generally in the earlier stage of growth, *ie* lignification, appears to be not merely similar to but identical with that of jute. Proof of this has been afforded by the particular investigation of such widely differing structures as the stony concretions of pears—Erdmann's glycydrupose—the fibrovascular bundles of *Musa Paradisiaca* (Monocot), and the bast fibres of the *Sida* species (*C J* 1882, 108, 1883, 19, 1889, 212).

But very few of the woods have been particularly investigated in regard to the constitution of the wood substance, and for the most part only in regard to the products of hydrolysis (v vol 1 p 719). Thus Erdmann concluded from his study of coniferous wood that its chief constituent—'glycolignose'—is a chemical individual C₃₀H₄₄O₂₁, a species of glucoside resolved by treatment with boiling acids into 'lignose' C₁₈H₂₆O₁₁ and a fermentable sugar (glucose). The presence of closed C₆ chains in the 'lignose' molecule was evidenced by its yielding protocatechuic acid on fusion with alkaline hy-

¹ A more direct conversion of the lignone into definitely aromatic products is that which takes place spontaneously when masses of jute are exposed to moisture and heat. From specimens of fibre 'rotted' under such conditions Cross and Bevan obtained an astringent substance C₁₀H₆O₄, which yielded phloroglucinol and protocatechuic acid on fusion with potash (*C J* 1882, 93).

drates It was also concluded that the wood of the widely different poplar species was similar if not identical in composition (*A Suppl* 5, 228)

The later researches of F Bente (*D P J* 217, 235), although modifying these views, chiefly in the variation in the results of hydrolysis, nevertheless in the main confirm them. The general conformity of the woods to the types above described as representative is shown

(1) In the close similarity of their characteristic reactions. Of these we may mention (a) the colouration produced with solutions of the aromatic amines (golden yellow), and of solutions of the phenols in hydrochloric acid, most characteristic of which is the reaction with phloroglucin (crimson), (b) the powerfully reducing action of the wood substance upon the oxides of copper, silver, gold, and mercury—showing the presence of aldehydic groups, (c) the reactions with the halogens yielding, in the case of chlorine, substitution products of definite quinone chloride characteristics, attended by complete resolution into lignone (chloride) and cellulose with bromine and iodine less definite compounds, but constant under constant conditions, (d) with nitric and sulphuric acids yellow-coloured explosive nitrates, (e) with acetic anhydride at its boiling point and with benzoyl chloride in presence of alkalis, the corresponding ethereal derivatives, (f) with solutions of the caustic alkalis at 160°–190°, with bisulphites (of the alkaline earths) at 150°–170° and with sulphurous acid (7 p.c. solution) at 90°–105°, attended by complete resolution into lignone (soluble derivatives) and cellulose (insoluble)

(2) In their empirical composition, which shows a remarkable uniformity throughout the group. This is illustrated in the appended table of results of analyses and determinations of calorific equivalents (Gottlieb, *J pr* 2] 28, 385) —

Wood	Ash	Nitrogen	Carbon	Hydrogen	Calorific equivalents per 1 gram
Oak	0.37	—	50.16	6.03	4620
Ash	0.57	—	49.18	6.17	4711
Hornbeam	0.50	—	48.99	6.20	4728
Beech	0.57	0.09	49.06	6.11	4777
Birch	0.29	0.10	48.88	6.06	4771
Fir	0.28	0.05	50.36	5.92	5035
Pine	0.37	0.04	50.31	6.20	5085

An investigation by N Schuppe of the chemical composition of a number of woody tissues (*Pharm J* [3] 14, 52) led to the following conclusions (a) that the woods are uniform in their characteristics, being composed of cellulose and 'lignin' in somewhat variable proportions, (b) the cellulose when isolated (Schultze's process) having the composition $C_6H_{10}O_5$, and (c) 'lignin,' being represented by the empirical formula $C_{11}H_{18}O_6$, which is closely similar to that obtained for the 'lignone' of jute. Further, G W Hawes has examined the woods of typical acrogens, e.g. lycopodium, equisetum, and aspidium, and finds that they do not differ essentially in composition from exogenous woods (*Am S* [3] 7, 585)

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(3) In the products of their destructive distillation, which, when carried out under uniform conditions, gives results which are very similar for the several woods. A very elaborate series of such distillations was carried out by M Senff (*B* 18, 60), the results being given in percentages of the wood under (1) total distillate, (2) tar, (3) crude vinegar, (4) anhydrous acid (acetic), (5) charcoal, (6) gases. The following numbers represent the limits of the percentages observed: (1) 40–50, (2) 3–8, (4) 2–6, (5) 20–30, (6) 17–35. The woods of all the more important species, e.g. Quercus, Populus, Betula, Fagus, and Pinus, were included in the research, and shown by the results to resemble one another very closely in composition. The products may be more particularly classified as regards the light which they throw upon the molecular grouping of the parent substance into (a) members of the fatty series alcohols, ketones, aldehydes, and acids generally of low molecular weights, (b) furfural and its homologues, (c) a group of aromatic bodies, constituting 'creosote,' which may be described as a mixture of guaiacol and creosol, containing besides, methyl creosol and the dimethyl ethers of pyrogallol and its homologues in varying proportions (Schorlemmer, 'History of Creosote,' *S C I* 4, 152). These divisions, it will be noted, correspond with the ascertained grouping of the complicated lignone molecule (*supra*), of which they afford additional confirmation. It is obvious that the process of destructive distillation must be attended with complications arising from secondary reactions, but the temperatures in the case of wood are sufficiently low to reduce these to a minimum, thus the yields of furfural are at a maximum when the temperature does not exceed 200° (Heil, *B* 10, 936), and the acetic acid is increased considerably beyond the percentages given by Senff (*supra*), viz. from 2–6 to 5–10, by raising the temperature very gradually through 150°–300° (W Rudnew, *D P J* 264, 88 & 128), no doubt at the expense of the methylation of the aromatic derivatives. This subject, however, except in regard to the outlines which we have sketched, belongs rather to the general theory of destructive distillation.

The fundamental tissue of the woods we regard, therefore, in all cases as a lignocellulose, of which the lignone portion, while possessing the typical characteristics common to the group, is no doubt variously differentiated with the specialised characteristics of families, and indeed species and individuals.

In addition to the fundamental tissue the woods contain other constituents, which from their nature are seen to be more or less adventitious. They are, in fact, generally removable by the action of solvents (in the case of resins, gum resins, balsams, &c.), or by simple hydrolysis. In this group we may mention such definitely aromatic derivatives as coniferin and vanillin (M Singer, *M* 3, 395), the tinctorial constituents of the dye woods, also the very characteristic and important constituent of most exogenous woods known as 'wood-gum' (Holzgummi), first described by Thomsen (*J pr* [2] 19, 146), who found quantities varying from 8 p.c. to 20 p.c. of the woods examined. This body yields on hydrolysis the C₆ sugar, xylose (Tollens, *Unters a d Agr Chem Lab Göttingen An-*

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malen, 254, 325), which by further resolution yields furfural. Whether the wood gum is a product of resolution of the lignone molecule has not yet been disclosed, but it would appear to be probable. A similar remark applies to the aromatic derivatives above mentioned.

In conclusion we must briefly notice a recent contribution to the subject of the 'constitution of lignin' (Lange, *Ztschr Physiol Chem* 14, 217). This consists in a study of the soluble products of resolution of certain woods—beech, oak, and fir—by heating with strong solutions of the alkaline hydrates at 185°. In addition to formic and acetic acids, traces of higher fatty acids, oxalic acid and small quantities of pyrocatechol and protocatechuic acids, there were obtained certain amorphous bodies of brown colour, which the author terms lignic acids. These were resolved by treatment with alcohol into (a) soluble, (b) insoluble in this menstruum. The empirical composition of these derivatives is subjoined

	(a)	(b)
Lignic acids	Beech C 61.3 H 5.4 Oak C 60.9 H 5.4 Fir C 61.5 H 5.0	C 59.0 H 5.4 — C 60.4 H 5.0

The yield of these bodies is from 12 to 15 p.c. of the weight of the wood. The insoluble residues from the original alkaline digestion are described as 'celluloses,' but the description is doubtful. These results have an empirical value, but throw little light on the constitution of the wood substance. They afford additional evidence, however, of the general similarity of woods of various origin.

Digested with alkalis at higher temperatures (200°–250°) than those employed in the researches detailed above, the lignocelluloses are resolved for the most part into oxalic and carbonic acids. With potassium hydrate—which appears to give the maximum yield—the following proportions of oxalic acid have been obtained, the yield being calculated on the dry woods: pine, 94.7 p.c.; poplar, 93.2 p.c.; oak, 83.4 p.c. (W Thorne, *D P J* 210, 24). It is evident that the oxalic acid is derived from both lignone and cellulose, which are therefore probably similarly constituted as regards the arrangement of the C atoms.

The action of the alkalis, however, at the point at which they resolve the lignocellulose is too severe to afford any trustworthy evidence, from the study of the products, as to the constitution of the original substance. The problem can only be solved by first studying those reactions which yield definite substitution or other real derivatives; these are chiefly chlorination, conversion into nitrates, acetates, and benzoates, and the reaction with bisulphites (solutions) at high temperatures.

In this article we do not attempt a special description of the woods or their constituents, our endeavour is rather to generalise what is known concerning the wood substance proper, that which resists mechanical solvents altogether, and hydrolytic agents up to a certain degree of intensity. C F C

LIGRON The mixture of homologues of methane obtained by collecting the portion of petroleum that boils below 100°.

LIGULIN A crimson colouring matter in ripe privet berries (Nicklés, *J Ph* [3] 35, 328),

sol water and alcohol, insol ether. Does not contain nitrogen. Coloured green by alkalis.

LIGUSTRIN A yellow hygroscopic bitter mass extracted from leaves of the privet (*Ligustrum vulgare*), insol ether and alcohol, sol water and dilute alcohol. Conc. H_2SO_4 gives an indigo blue solution (Polex, *Ar Ph* [2] 17, 75).

LIGUSTRON [c 100°] (260°–280°). Occurs in privet bark (Reinsch, *J* 1847, 787). Needles, v sol water, alcohol, and ether. Tastes bitter. Reduces ammoniacal AgNO_3 .

LIME Oxide of calcium, CaO , v vol 1 p 368.

LIME, CHLORIDE OF A name sometimes given to *bleaching powder*, v BLEACHING POWDER, vol 11 p 17.

LIME LEAF OIL The fragrant oil obtained by distilling the leaves of *Citrus Limetta* with steam contains a citrene (c 176°), 1 active to light, and with refractive index for red rays 1.4611 at 30°, terpinol, methyl ennyl ketone, and a colophene (F Watts, *C J* 49, 316).

LIMES, OTTO OF Obtained by rasping and squeezing from the unripe peel of the fruit of *Citrus limetta*. Contains a terpene (176°) and a soft resin, not volatile at 250°. On standing the resin deposits $\text{C}_{10}\text{H}_{16}\text{O}$, [102°] (Wright a Piesse, *C J* 32, 548).

LIMETIC ACID $\text{C}_8\text{H}_8\text{O}_4$. Obtained by the action of H_2SO_4 and $\text{K}_2\text{Cr}_2\text{O}_7$ on the oil of lime (from *Citrus Limetta*) and on oil of rosemary (Vohl, *N Ber Arch* 74, 16). White crystalline body, may be sublimed. Has no taste or smell. Sol water, v sol alcohol—Ag. A powder, sl sol water, blackening on exposure to light.

LIMETTIN $\text{C}_8\text{H}_8\text{O}_4$, [122°]. A neutral body occurring in oil of limes (Tilden, *C J Proc* 6, 30). Tufts of pale yellow needles (from alcohol). Not acted upon by AcCl or by phenylhydrazine. Bromine forms colourless scales of $\text{C}_{10}\text{H}_{11}\text{Br}_2\text{O}_6$. Boiling conc. NaOH forms NaOAc and crystalline $\text{C}_8\text{H}_{11}(\text{OH})\text{O}$.

LIMONENE A terpene occurring in oil of lemon and in many other essential oils, v TERPENE.

LIMONIN $\text{C}_{15}\text{H}_{24}\text{O}$, [275°] (Paterno a Ogialoro, *G* 9, 64), [245°] (Hoffmann *Ar Ph* [3] 14, 839). A bitter substance contained in the pips of oranges and lemons (Bernays, *Buchner's Rep* [3] 21, 306, *A* 40, 317, Schmidt, *A* 51, 338). The pips are exhausted with boiling water, alcohol is added, and the mixture is boiled with inverted condenser. After filtering, and distilling off the alcohol, limonin is left, together with a fatty substance which may be removed by CS_2 . 1500 g of pips yield 80 g of limonin (P a G). Laminæ, sl sol water, ether, and NH_4Aq , v sol alcohol, HOAc , KOH aq, and baryta water. The barium salt is not decomposed by CO . Conc. H_2SO_4 forms a red liquid, from which the limonin is ppd by water.

LINALOES OIL The essence of linaloes, obtained from the wood of *Lycari kanali*, the white cedar of Cayenne, is a slightly coloured liquid with an odour like rose and lemon. After distillation over CaCl_2 it consists of $\text{C}_{10}\text{H}_{16}\text{O}$ (198°) SG 1.868 [α_D^{20}] = -19 at 15°. It is sol alcohol, ether, and glycerin. When distilled with ZnCl_2 it yields a terpene, v LIGARENE. By prolonged treatment with saturated HCl aq and

exposure to light it yields $C_{18}H_{35}BrO_2$, an optically inactive liquid with a camphor like odour, and SG 1.069 When distilled with lime it yields leucarene $C_{18}H_{36}$ (ca 170°), SG 1.835 (Morin, *C R* 92, 998, 94, 733)

LININ $C 62.9 p c H 47 p c$ AcrySTALLISABLE substance obtained by digesting *Linum catharticum* with milk of lime, filtering, ppg with HCl, and extracting with ether. (Pagenstecher, *A* 40, 322, C Schröder, *N Rep Pharm* 10, 11) Silky crystals, v sl sol water, v e sol alcohol and ether, m sol chloroform and HOAc The alcoholic solution is intensely bitter

LINOLEIC ACID $C_{18}H_{32}O_2$ *Linolic acid* SG 1.492 Occurs as glyceryl ether in linseed oil, hemp oil, poppy oil, olive oil, nut oil, cotton seed oil, earth nut oil, almond oil, oil of sesame, palm oil, cacao butter, and probably in most fixed vegetable oils (Pelouze & Boudet, *A Ch* [2] 59, 43, Laurent, *A Ch* [2] 65, 150, 298, Liebig, *A* 33, 113, Sacc, *A* 51, 214, Schuler, *A* 101, 252, Oudemans, *J* 1858, 304, Hazura & Grussner, *M* 9, 944, 10, 242, Benedikt & Hazura, *M* 10, 353) It does not occur in animal oils, so that when the acids obtained by saponifying animal oils are oxidised by $KMnO_4$, no sativic acid will be formed, but di oxy stearic acid will be among the products, this being derived from oleic acid (B & H)

Preparation—Crude linseed oil or hemp oil is evaporated with aqueous NaOH, the sodium soap is decomposed by H_2SO_4 , and the crude acid dissolved in alcohol, neutralised with NH_3 , and ppd by $BaCl_2$. The barium salt is dissolved in ether, HCl is added, the ether decanted from ppd $BaCl_2$, evaporated, and the liberated acid dried *in vacuo* over H_2SO_4 (Schuler, Bauer & Hazura, *M* 7, 216) The acid so obtained is a mixture of oleic, linoleic, linolenic, and iso linolenic acid (Hazura) By treatment with bromine at 0° and reduction of the product $C_{18}H_{32}Br_2O_2$ with zinc and HCl pure linoleic acid is got

Properties—Faintly-yellow lumpid oil Does not solidify at -18° Insol water, v sol ether, m sol alcohol When distilled under 90 mm pressure one third passes over at 290° , and the distillate contains sebacio acid $C_{16}H_{30}O_2$ [130°] and an oily mixture (Hazura & Grussner, *M* 9, 206, cf Norton & Richardson, *B* 20, 2735)

Reactions—1 Potash fusion gives myristic, acetic, and formic acids, with traces of azelaic acid—2 $KMnO_4$, hydrogen peroxide, and MnO_2 with H_2SO_4 oxidise it to azelaic acid—3 $KMnO_4$ in presence of KOHAq converts it into sativic acid, some azelaic acid also being formed But alkaline $KMnO_4$ oxidises crude linoleic acid from linseed oil (100 g) into sativic acid $C_{18}H_{32}(OH)_2O_2$ [173°] (6.5 g), linusic acid $C_{18}H_{30}(OH)_2O_2$ [204°] (4.5 g), isolinusic acid [175°] (15.8 g), and di oxy stearic acid $C_{18}H_{34}(OH)_2O_2$ [137°] (1.2 g) (Hazura, *M* 9, 180 cf Dieff & Reformatsky, *B* 20, 1211) It appears therefore that crude linoleic acid consists of linoleic acid $C_{18}H_{32}O_2$ (which gives sativic acid on oxidation), linolenic acid $C_{18}H_{30}O_2$ (which gives linusic acid), isolinolenic acid (which gives isolinusic), and oleic acid (which gives di oxy stearic acid) The formation of sativic acid in this manner may be made use of as a test for linoleic acid—4 Bromine at ordinary temperatures forms with crude linoleic

acid solid $C_{18}H_{30}Br_2O_2$ [177°] Bromine at 0° forms a tetrabromide $C_{18}H_{26}Br_4O_2$ [115°] (Hazura, *M* 8, 147, Hazura & Friedrich, *M* 8, 155, 265) Of these two compounds the former is formed from linolenic, the latter from linoleic acid Pure linoleic acid gives only $C_{18}H_{32}Br_2O_2$ [115°]—5 Fuming HIAq and amorphous phosphorus converts it into stearic acid (Peters, *M* 7, 552)

Salts—The salts are not crystalline With the exception of the salts of the alkalis they are insol water They are sol ether With the exception of the Pb, Mn, Na, and NH_4 salts they are insol alcohol— BaA' (Peters)

Linolenic acid $C_{18}H_{30}O_2$ Obtained by treating the compound $C_{18}H_{32}Br_2O_2$ [177°] (*v supra*) dissolved in alcohol with zinc and HCl (Hazura, *M* 8, 267) Yields on oxidation by alkaline $KMnO_4$ no solid acid except linusic (hexa oxy stearic) acid [201°] Bromine forms only $C_{18}H_{30}Br_2O_2$ [177°] Linolenic acid 'dries' rapidly when exposed to air, through oxidation Its salts behave in like manner The more glyceryl linolenate there is in an oil the more rapidly does it dry Glyceryl linolenate and isolinolenate also possess drying properties, but glyceryl oleate does not The product produced by exposing the acids to air is an anhydride, insol ether, but furnishing soluble acids when heated with alkalis

Isolinolenic acid $C_{18}H_{30}O_2$ An acid assumed to exist in crude linoleic acid on account of the formation of isolinusic acid on its oxidation When crude linoleic acid is oxidised by $KMnO_4$ in alkaline solution, and the product ppd by H_2SO_4 , there is obtained a mixture of fatty acids whence cold ether extracts di oxy stearic acid, sativic acid remaining undissolved The filtrate from the ppd acids is neutralised with NaOH, evaporated, again ppd with H_2SO_4 , the pp extracted with ether, and the residue crystallised from alcohol and then from water, whereby it may be separated into linusic and isolinusic acids

LINSEED OIL The oil expressed from the seeds of flax (*Linum usitatissimum*) Like other drying oils when exposed to the air it dries up to a transparent resinous mass Linseed oil is composed of the glyceryl ethers of oleic, linoleic, linolenic, and isolinolenic acids **LINOLEIC ACID** Linseed oil dissolves some oxide of lead when heated therewith, being decolourised and rendered more easily drying (boiled oil)

LINUSIC ACID *v* HEXA OXY STEARIC ACID

Isolinusic acid *v* HEXA OXY STEARIC ACID

LIQUIDAMBAR A balsam obtained from a large tree, *Liquidambar styraciflua*, growing in Florida and Mexico It resembles balsam of Peru, containing cinnamyl cinnamate, styrene, and cinnamic acid (Harrison, *Ar Ph* [3] 6, 541, Maisch, *Ar. Ph* [3] 6, 545)

LIQUIDS, diffusion, dispersion, osmose, refraction, transpiration, of, *v* PHYSICAL METHODS

LITHIUM Li At w 7.01 Mol w probably 7.01 (*v p* 149) [180°] (Bunsen, *J* 8, 324) SG 578 to 589 (Bunsen, *l c*) SH 27° to $100^\circ = 9408$ (Regnault, *A Ch* [3] 68, 11) EC at 20° (Hg at $0^\circ = 1$) 10.69 (Mathiessen *P M* [4] 12, 199, 13, 81) Characteristic lines in emission spectrum are L_{α} in the red 6705.2, and a

weaker line in the orange $\delta 102$, and a weak line in the blue 4602.7 (Thalen, 1868, v also L de Boisbaudran, *Spectres Lumineux*, p 55, Schön, W 10, 143, Liveing & Dewar, T 1883 187) S V S 119

Occurrence—Salts of Li are very widely distributed, but occur only in small quantities. Several phosphates of Fe and Al contain Li phosphate, especially *traphylina*, which contains c 7 p c Li_2O , many silicates of Al and Fe contain small quantities of Li silicate, especially *lepidolite*, which sometimes contains from 3 to 5 p c Li_2O . Tourmalines, borates, &c, often contain traces of lithia. Very many mineral springs contain lithia, according to Kirchhoff a Bunsen lithia is present in almost all mineral waters (P 113, 357). Truchot found lithia in the soil of L'imague in the Auvergne, nearly all the plants growing on this soil take up lithia (C R 78, 1022). Dieulaufait has found traces of Li in sea water from all parts of the globe, in the water of marshes, in mineral springs, in primary rocks, in gypsum of different formations (A Ch [5] 17, 377). Lithia is found in many plants, and in all sorts of tobacco, but not in raw sugar, cocoa, coffee, or tea (Focke, *Der Naturforscher*, 1872 307, Grandeau, A Ch [3] 67, 216). Lithia has also been found in the milk and blood of cows (Bunsen & Kirchhoff), in different parts of the human organism (Bence Jones, P M [4] 29, 394), and in normal urine (Schiaparelli & Peroni, G 10, 390).

While investigating various silicates in 1817, Arfvedson (S 22, 93, 34, 214) found a new alkali base with a molecular weight smaller than that of soda or potash. Berzelius gave the name *lithia* to the new base (*lithios*) supposing that, unlike soda and potash, it was to be found only in minerals. The new alkali was decomposed by electrolysis in 1818 by Davy, also in 1820 by Brandes (S 8, 120), but the metal was first prepared approximately pure and in considerable quantity by Bunsen and Matthiessen in 1855 by electrolyzing fused LiCl (A 94, 107).

Preparation—The metal is obtained by electrolyzing molten LiCl . Bunsen & Matthiessen (A 94, 107) passed the current from 4-6 Bunsen cells through LiCl kept molten in a thick walled porcelain crucible, using a cylindrical rod of rector graphite as positive electrode, and an iron wire the thickness of an ordinary knitting needle as negative electrode. The metal separated on the iron wire in small pellets, which were quickly removed by an iron spoon and placed under petroleum. Some of the metal was always oxidised, occasionally with ignition. To obviate this, Hiller (*Neues Handwörterbuch der Chemie*, 8, 534) passed the negative electrode (iron wire) through the stem of a tobacco pipe, which he connected with an apparatus supplying pure dry H_2 , he allowed H_2 to pass through the pipe until air was completely expelled, then plunged the bowl open end downwards, with the end of the wire inside, into the molten LiCl , stopped the H_2 , and sent the current through the LiCl , when sufficient Li had collected inside the bowl, the pipe was broken and the metal collected under petroleum. To prevent any action between the Li and the silica in the pipe, the inside of the bowl is covered with a thin layer of graphite, this is done by mixing powdered graphite with

dilute LiCl aq so as to form a thick paste, spreading this inside the bowl, and drying first in air and then at a moderate red heat. It is advantageous to mix the LiCl before fusion with some NH_4Cl .

There are many methods for preparing LiCl from Li containing minerals, the methods vary according to the composition of the mineral dealt with, all seek to prepare a solution containing only the alkalis, from which Li may be separated by taking advantage of the comparatively small solubility in water of Li_2CO_3 . LiCl is obtained by dissolving Li_2CO_3 in HCl aq, evaporating, and drying the crystals which separate. *Lepidolite* is the usual starting point, different specimens contain from less than 1 to c 5 p c Li_2O . The mineral is very finely powdered and triturated with water, the finest powder is dried and heated to redness with twice its weight of lime, the cold mass, in which the SiO_2 is combined with lime, is treated with HCl aq, CaO is added from the solution by H_2SO_4 aq and evaporation, the filtrate is evaporated to dryness, and the solid is heated until H_2SO_4 is all removed, the residue is dissolved in water, the solution is digested with CaCO_3 to remove Al_2O_3 , and Ca is removed by pptn with $(\text{NH}_4)_2\text{C}_2\text{O}_4$, the filtrate is evaporated to dryness, and the residue is strongly heated, the Li_2SO_4 thus obtained is dissolved in water, and the liquid is added by Ba acetate, after filtration the Li acetate is strongly heated and so transformed into Li_2CO_3 (Arfvedson, S 22, 93, 34, 214).

The method recommended by Hauer (J pr 68, 310) consists in strongly heating for 2 hours a mixture of equal parts of very finely powdered *lepidolite* and gypsum, lixiviating the mass with water, filtering, evaporating until CaSO_4 and K_2SO_4 crystallise out, adding to the mother-liquor a mixture of NH_4Aq , NH_4HSAq , and $(\text{NH}_4)_2\text{C}_2\text{O}_4$ aq, whereby all bases are pptd except the alkalis, after filtering, Li_2CO_3 is pptd from the warm solution by $(\text{NH}_4)_2\text{CO}_3$.

The process adopted in Schering's manufactory at Berlin is described by Filsinger (D P J 219, 183, 222, 321, 385). Finely ground and sifted *lepidolite* is mixed with conc H_2SO_4 in a warm brick trough to the consistence of a thin paste, which is heated with slight stirring till it forms into lumps, the lumps are calcined in a reverberatory furnace, and, while warm, are lixiviated with water, the liquid is mixed with enough K_2SO_4 to convert all Al_2O_3 into alum, which separates on boiling, the residual Al_2O_3 being removed by milk of lime, the salts in the filtrate are converted into chlorides by pptn with BaCl_2 aq, and the liquid is evaporated to dryness, digestion with absolute alcohol dissolves the chlorides of Li and Ca , after distilling off alcohol, Ca is pptd by $(\text{NH}_4)_2\text{C}_2\text{O}_4$ aq, the liquid is filtered, and a little NH_4HSAq is added to ppt any Fe, &c, still present, the filtrate is boiled to remove NH_4HS and evaporated to dryness in a silver dish, pure LiCl is thus obtained. The LiCl may be converted into Li_2CO_3 by dissolving in water, adding NH_4Aq and $(\text{NH}_4)_2\text{CO}_3$, and washing the ppt with alcohol of 60 p c.

Schrötter's method is said to be one of the best (J pr 93, 275). *Lepidolite* is melted, at full red heat, with frequent stirring, the molten mass is ladled out by an iron spoon into water,

when cold the solid is powdered and triturated with water, HClAq S G 12 is added little by little to the pasty mass, care must be taken that sufficient water is present to prevent the whole mass from solidifying, after standing for 24 hours, with frequent stirring, the semi liquid substance is heated nearly to boiling, and a little more HClAq S G 12 is added, the total quantity of HCl used should be c 2 parts to 1 part lepidolite, after a few hours most of the SiO_2 has separated, a little of the filtered liquid should be so acid that no permanent pp is formed on addition of a few drops of $\text{Na}_2\text{CO}_3\text{Aq}$, a little HNO_3Aq is now added to completely oxidise FeCl_2 to FeCl_3 , the liquid is filtered from ppd SiO_2 (which separates as a powder), and Fe_2O_3 , Al_2O_3 , CaO , MgO , &c, are ppd by careful addition of $\text{Na}_2\text{CO}_3\text{Aq}$ to the boiling liquid. The alkaline filtrate is nearly free from all salts except chlorides of the alkalis, it is evaporated until the small quantities of MgCO_3 , MnCO_3 , &c, still present separate out, and Li_2CO_3 is ppd from the filtrate by addition of Na_2CO_3 and evaporation.

Commercial Li_2CO_3 generally contains small quantities of salts of Mg, Ca, K, Na, &c, it may be purified by one of the foregoing methods. For the other methods of preparing Li_2CO_3 from lepidolite, &c v Hugo Muller, *J pr* 58, 148, Fuchs, *J pr* 5, 319, Troost, *A Ch* [3] 51, 103, Mallet, *A* 101, 389, Lunglmayr, *D P J* 171, 293, Allen, *J pr* 87, 480, Reichardt, *D P J* 172, 447, L Smith, *A* 159, 82, Stolba, *D P J* 198, 225, L de Boisbaudran, *Bl* [2] 17, 551. A detailed criticism of various methods will be found in *D P J* 219, 183, 222, 271 385.

Properties — A silver white metal, very soft, but harder than K or Na, when freshly cut, the surface appears yellowish, when melted and at once pressed between glass plates it forms a silver like mirror. Li makes a grey streak on paper. It may be drawn into wire, but shows very little tenacity. Li is the lightest known solid, S G c 59, it swims on rock oil. Melts at 186° . Is not acted on by dry O at its M P, heated in air to c 200° it burns with a very brilliant white flame. May be vapourised in H at full red heat. Li decomposes cold water without itself melting, it combines rapidly with Cl, Br, I, S, O, it burns when heated in dry CO_2 . Li reacts with most acids to form salts, conc HNO_3Aq oxidises it with great rapidity, the metal usually melts, and is sometimes ignited. Li dissolves in liquid NH_3 , and on evaporation of the NH_3 it is left unchanged (Seeley, *C N* 23, 169).

The atomic weight of Li has been determined (1) By converting LiCl into AgCl (Arfvedson, *S* 22, 43, Mallet, *Am S* [2] 22, 349, Troost, *A Ch* [3] 51, 108, Stas, *Nouv R* 268), (2) by determining O in Li_2O (Berzelius, *P* 17, 379), (3) by ppg Li_2SO_4 by BaCl_2 (Berzelius, *P* 17, 379, Heumann, *P* 15, 482, Hagen, *P* 48, 361, Diehl, *A* 121, 97), (4) by determining CO_2 in Li_2CO_3 (Hermann, *P* 15, 480, Troost, *A Ch* [3] 51 108, Diehl, *A* 121, 93), (5) by converting Li_2CO_3 into Li_2SO_4 (Troost, *A Ch* [3] 51, 108), (6) by converting LiCl into LiNO_3 (Stas, *Nouv R* 274), (7) by determining SH of Li (Regnault, *A Ch* [3] 68, 11).

Ramsay (*O J* 55, 521) has endeavoured to

determine the mol w of Li by measuring the lowering of vapour pressure of Hg produced by dissolving Li in Hg the results make it probable that the mol w of Li is the same as the at w. This result is based on the assumption that Van't Hoff's law holds good, viz, that equal volumes of dilute solutions contain equal numbers of molecules of the dissolved substance, it also presupposes that the molecular weight of liquid Hg is the same as the atomic weight.

As the V D of no Li compound has yet been determined, the valency of the atom Li in gaseous molecules is not certainly known, but from the close analogy between Li and the other alkali metals there can be little doubt that the atom of Li is monovalent in gaseous molecules.

Li is a strongly positive metal, it belongs to the group of alkali metals, none of which shows any tendency to enter into the negative radicle of salts. Li shows closer resemblances to the alkaline earths than are exhibited by any other metal of the alkalis, LiOH , Li_2CO_3 , and Li_3PO_4 are much less soluble in water than the corresponding compounds of Na, K, Rb, and Cs, Li does not form an alum. The position and analogies of Li are discussed in the article ALKALIS, METALS OF THE, vol 1 p 114.

Reactions and Combinations — 1 Li decomposes cold water rapidly without itself melting. Thomsen (*Th* 3, 227) gives the thermal data $[\text{Li}, \text{H O Aq}] = 48,970$ (to form $\text{LiOH Aq} + \text{H}$) — 2 Very rapidly oxidised by conc nitric acid — 3 Slowly acted on by conc sulphuric acid, rapidly dissolved by dilute $\text{H}_2\text{SO}_4\text{Aq}$, also by dilute hydrochloric acid (Matthiessen, *A* 94, 10) — 4 Burns when heated in dry carbon dioxide or sulphuretted hydrogen — 5 At temperatures lower than its melting point Li acts on silica, alkaline silicates, iron, gold, silver, and platinum — 6 Combines readily with sulphur, phosphorus, and the halogens.

Detection — Li compounds give a red colour to a non luminous flame. Examination by the spectroscope will detect 000009 mgrms Li present as LiCl (Bunsen). To detect Li in silicates, the powdered mineral is treated with HFAq, the liquid is poured off, and the residue is evaporated with addition of a little H_2SO_4 , the residue is extracted with absolute alcohol, and the liquid is evaporated to dryness, the residue is again treated with absolute alcohol, and this solution is again evaporated, and the residue ($\frac{1}{10}$ mgm is sufficient) is examined in the spectroscope, if the mineral is non-siliceous, the treatment with HFAq may be omitted. Li is estimated in the form of Li_2SO_4 .

Lithium, antimonate of LiSbO_3 . By adding LiCl to KSbO_3Aq (cf ANTIMONATES, vol 1 p 285).

Lithium, borate of $\text{Li}_2\text{B}_2\text{O}_5$, and hydrates with $5\text{H}_2\text{O}$, $6\text{H}_2\text{O}$, and $10\text{H}_2\text{O}$, by adding Li_2CO_3 to boric acid solution (Arfvedson, *A Ch* [2] 10, 82, Finsinger, *Ar Ph* [3] 8, 198, cf BORATES, vol 1 p 529).

Lithium, borofluoride of By double decomposition between $\text{Ba(BF}_4)_2\text{Aq}$ and $\text{Li}_2\text{SO}_4\text{Aq}$ and evaporation at 40° , large deliquescent prisms are obtained, al sol water, these crystals are probably LiBF_4 , but they have not been accurately examined (Berzelius).

Lithium, bromide of LiBr HF [Li, Br, Aq] = 91,810 (*Th* 3, 227), [Li, Br, Aq] = 11,850 (Bodisko, *J R* 1889 [1] 7) SG 3 102 at 17° (Clarke, *Am S* [3] 13, 293) A white crystalline, very deliquescent, mass Obtained by dissolving Li_2CO_3 in HBrAq and evaporating (Troost, *A Ch* [3] 51, 108), or by ppg excess of CaBr_2Aq by K_2CO_3 , after 24 hours adding enough Li_2CO_3 to ppt all the Ca , filtering, and evaporating (Klein, *A* 128, 239) S 143 at 0° , 196 at 34° , 222 at 59° , 244 at 82° , 270 at 103° (Kremers, *P* 103, 65)

Lithium, chloride of LiCl SG 2 074 at 39° (Schroder, *P* 106, 226), 1998 at 0° , 1 515 at M P (Quincke, *A* 138, 141), SG fused 1 575 (Wermicke, *P* 138, 141) HF [Li, Cl] = 93,810, [Li, Cl, Aq] = 102,250 (*Th* 3, 227) SH 282 (Regnault)

Preparation—1 By dissolving Li_2CO_3 in HClAq and evaporating—2 By decomposing $\text{Li}_2\text{SO}_4\text{Aq}$ by BaCl_2Aq , filtering from BaSO_4 , and evaporating

Properties—Crystallises from aqueous solution in regular octahedra, very deliquescent, more so than CaCl_2 , tastes like NaCl , melts at dark red heat to a clear liquid, which gives off some Cl and becomes alkaline when heated for a long time in the air, the same change occurs to a slight extent when LiClAq is evaporated E sol alcohol, also in a mixture of ether and alcohol in which KCl and NaCl are nearly insoluble Volatilised at white heat S 63 7 at 0° , 80 7 at 20° , 104 2 at 65° , 115 at 80° , 129 at 96° , 139 at 140° , 145 at 160° Gerlach (*Fr* 8, 279) gives the following —

SG LiClAq	Pc LiCl	SG LiClAq	Pc LiCl
1 008	1	1 148	25
1 030	5	1 182	30
1 058	10	1 219	35
1 086	15	1 256	40
1 117	20		

B P of saturated LiClAq = 171° (Kremers, *P* 103, 65)

Reactions—1 Heated in air for some time is partially decomposed with evolution of Cl , residue is alkaline (Schulze, *J pr* [2] 21, 407) Evaporation of LiClAq is accompanied by slight decomposition—2 Completely decomposed by heating in steam, with evolution of HCl , decomposition is rapid in presence of silica, but is prevented by admixture of NH_4Cl (Kunheim, *J* 1861 149)

Combinations—1 With water to form two hydrates (1) $\text{LiCl} \cdot 2\text{H}_2\text{O}$, obtained as quadratic crystals by evaporating LiClAq under 10° Dried between paper, the crystals become opaque and powdery, when warmed, melts in water of crystallisation, then solidifies, and then the dry LiCl melts again at red heat, (2) by evaporating a solution of LiCl in aqueous alcohol, Rammelsberg obtained the monohydrate $\text{LiCl} \cdot \text{H}_2\text{O}$ (*P* 66, 79)—2 With alcohol to form $\text{LiCl} \cdot 2\text{C}_2\text{H}_5\text{O}$, and with methyl alcohol to form $2\text{LiCl} \cdot 3\text{CH}_3\text{O}$, obtained by evaporating solution of LiCl in the respective alcohols (Simon, *J pr* [2] 20, 871)—3 With platinic chloride, to form $\text{Li}_2\text{PtCl}_6 \cdot 6\text{H}_2\text{O}$, orange-red salt, sol in water, alcohol, and ether-alcohol

Lithium, chromate and dichromate of, v. vol u pp 155, 157

Lithium, fluoride of LiF Small crystalline tablets, by dissolving excess of Li_2CO_3 in HFAq , filtering, and evaporating Slightly sol in water, melts at red heat (Berzelius, *A* 1, 17) SG 2 295 at 21.5° (Clarke, *Am S* [3] 13, 292) By solution of LiF in HFAq and evaporation, crystals of LiF HF are obtained, when heated LiF and HF are formed Fluckiger (*A* 87, 261) describes the double salt $2\text{LiF} \cdot \text{SbF}_5$, LiF combines with SiF_4 to form the *silico-fluoride* Li_2SiF_6 (*v* LITHIUM, SILICOFLUORIDE *ov*, p 151)

Lithium, haloid compounds of Lithium combines directly with the halogens, the haloid compounds, LiX , are generally prepared by dissolving Li_2CO_3 in the respective acids and evaporating As the V D of none of the compounds has been determined, their molecular weights are not known with certainty, but from the close similarities between compounds of Li , K , Na , and Cs there can be little doubt that the formula LiX ($\text{X} = \text{F}, \text{Cl}, \text{Br}, \text{I}$) expresses the composition of the molecules of the haloid compounds of Li

Lithium, hydrosulphide of LiSH Obtained by reducing Li_2SO_3 by C , and passing H_2S into a solution of the product Only known in solution (Berzelius, *P* 6, 439) Thomsen gives HF [$\text{Li}, \text{S}, \text{H}, \text{Aq}$] = 66, 120 (*Th* 3, 227)

Lithium, hydroxide of LiOH Obtained by boiling Li_2CO_3 with CaOAq in a silver dish (Pt cannot be used as it is acted on by LiOH), filtering, evaporating, and heating to 100° , better by ppg $\text{Li}_2\text{SO}_4\text{Aq}$ by an equivalent quantity of BaOAq , filtering, evaporating, and heating to 100° Also obtained by dissolving Li_2O (*q v*) in H_2O and evaporating Thomsen (*Th* 3, 227) gives HF [$\text{Li}, \text{O}, \text{H}, \text{Aq}$] = 117, 440, and heats of neutralisation [$2\text{LiOHAq}, \text{H}_2\text{SO}_4\text{Aq}$] = 31,290, [$2\text{LiOHAq}, \text{H}^+\text{Cl}^+\text{Aq}$] = 27,700 Beke toff (*Bl* 41, 312) gives [$\text{Li}^+\text{O}, \text{Aq}$] = 13,000 (to form LiOHAq)

A white crystalline mass which melts when heated, without decomposition, sol in water, but less so than KOH or NaOH , insol in ether alcohol Gmelin obtained small crystals of LiOH by evaporating a solution *in vacuo*, according to Muretow (*B* 5, 331) the crystals are a *hydrate of lithium hydroxide*, $\text{LiOH} \cdot \text{H}_2\text{O}$ (*cf* Dittmar, *S C I* 7, 730)

Lithium, iodide of LiI Obtained by saturating HIAq , containing a little H_3PO_4 , with Li_2CO_3 , warming the ppt Li_3PO_4 with BaI_2 and a trace of H_2SO_4 , filtering, adding enough Li_2CO_3 to decompose excess of BaI_2 present, filtering again, evaporating, crystallising, and drying by pressure between filter paper (Liebig, *A* 121, 222) Liebig (*lc*) also recommends to neutralise half of an HI solution containing a little H_3PO_4 by BaO or CaO , to add the other half of the acid, neutralise by Li_2CO_3 , filter from Ba or Ca phosphate and evaporate The crystals are generally yellowish from a little separated I , this is removed by quickly pressing between filter paper LiI forms small, colourless, deliquescent crystals SG 3 485 at 23° (Clarke, *Am S* [3] 13, 293) HF [$\text{Li}, \text{I}, \text{Aq}$] = 76, 100 (*Th* 3, 227) S 151 at 0° , 164 at 19° , 179 at 40° , 200 at 59° , 263 at 75° , 435 at 80° , 476 at 99° , 588 at 120° (Kremers, *P* 103, 65) By evaporating a solution of Li_2CO_3 in HIAq over H_2SO_4 , Ram

melberg obtained the hydrate $\text{Li}_2\text{S}\cdot\text{H}_2\text{O}$ (P 66, 79)

Lithium, oxide of Li_2O Obtained by burning small quantities of Li, in a small iron vessel, in dry O at 200° , cooling in O, and heating in the air to decompose Li peroxide (Troost, *A Ch* [3] 51, 103) Also prepared by heating Li_2CO_3 with C in a Pt crucible, and by heating LiNO_3 to redness in a Ag dish, best mixed with Cu turnings (H Muller, *J pr* 58, 148)

A white crystalline solid, S G 2 102 at 15° (Brauner & Watts, *P M* [5] 11, 60) Not decomposed by heating with C or Fe Does not act on Pt at high temperatures, corrosion of the Pt vessel in the preparation of Li_2O indicates the presence of H_2O or Cs O Reacts with Cl, S, and P Heated in O, Li_2O is superficially changed to peroxide Thomsen gives $[\text{Li}_2\text{O}, \text{Aq}] = 166.5^\circ$ (Th 3, 227), and Beketoff (*Bl* 41, 312) gives $[\text{Li}_2\text{O}, \text{Aq}] = 13,000$, hence $[\text{Li}_2\text{O}] = 153,520$

Lithium peroxide is said to be formed by heating Li_2O , or Li_2CO_3 , for some time in air or O, but to be decomposed at a little above the temperature of formation

Lithium, phosphide of According to Troost (*A Ch* [3] 51, 103) Li and P combine, when heated together, to form a brown substance which is decomposed by water with evolution of inflammable P hydride

Lithium, silicofluoride of $\text{Li}_2\text{SiF}_6\cdot 2\text{H}_2\text{O}$ Transparent monoclinic crystals, obtained by adding H_2SiF_6 aq to Li acetate or carbonate evaporating, treating the residue with water, filtering, and crystallising (Stolba, *J pr* 91, 456) S 526 at ordinary temperature, sol alcohol, insol ether S G 233 Dehydrated at 100° , melts at a higher temperature with evolution of SiF₄

Lithium, salts of Compounds produced by replacing H of acids by Li The Li salts belong to one series Li_xX , where $\text{X} = 2\text{Cl}, 2\text{NO}_3, \text{SO}_4, \text{CO}_3, \frac{2}{3}\text{PO}_4$, &c, they are generally obtained by dissolving Li_2CO_3 in the different acids, some are prepared by double decomposition from LiCl or Li_2SO_4 Most of the salts of Li are sol in water, but Li_3PO_4 and Li_2CO_3 are considerably less sol than the corresponding salts of the other alkali metals, LiOH is also less soluble than the other alkalis (cf ALKALIS, METALS OF THE, vol 1 p 114) Few, if any, basic salts of Li are known The chief Li salts are the following (v CARBONATES, NITRATES, SULPHATES, &c) *Antimonate, arsenate, borate, bromate, carbonate, chlorate, chromates, ditimonate, hypochlorite, hypophosphite, iodate, nitrate and nitrite, perchlorate, periodate, phosphates, selenate and selenites, silicates, sulphates and nitrite, tellurate and nitrite, thio-arsenate*

Lithium, sulphide of Li_2S Li and S combine when heated together, the solution in water is yellow from presence of polysulphides Li_2S is obtained by reducing Li_2SO_4 by an equivalent quantity of C at full red heat, excess of C makes the product pyrophoric (Berzelius, P 6, 439) Naudin & Mantholon (*C R* 83, 53) say that Li_2S may be prepared by the long-continued passage of H_2S through Li_2CO_3 suspended in water Li_2S is easily sol water and alcohol $[\text{Li}_2\text{S}, \text{S}, \text{Aq}] = 115,260$ (Th 3, 227)

Lithium polysulphides are said to be obtained by melting LiOH with S (Vauquelin,

A Ch 7, 284), they closely resemble the polysulphides of the other alkali metals

Lithium, sulphate of, v LITHIUM, HYDROSULPHIDE OF, p 150 M M P M

LITHO-BILIC ACID $\text{C}_{20}\text{H}_{30}\text{O}_8$ [199?] Occurs, together with lithofelic acid, in Oriental bezoar, and is prepared by decomposing the barium salt with hydrochloric acid and recrystallisation from alcohol Long pale-yellow needles, insol water, v sol alcohol, m sol ether Its alcoholic solution is dextrorotatory It resembles lithofelic and the bilary acids in its behaviour with Pettenkofer's reaction and its distillation products (Roster, *G* 9, 462, Grattarola, *J* 1880, 831) — BaA'_6 aq Ppd by adding BaCl_2 to a warm aqueous solution of crude sodium lithofellate, usually a yellowish semi-transparent resin, was obtained on one occasion in minute monoclinic crystals

LITHOFELIC ACID $\text{C}_{20}\text{H}_{30}\text{O}_8$ [205?] S (alcohol) 3.4 at 20° , 15 at 78° S (ether) 225 at 20° (Göbel) $[\alpha]_D = 13.8$ at 9.5° (independent of concentration) Forms the chief constituent of some kinds of Oriental bezoars (Göbel, *A* 39, 237, Etting, *A* 39, 242, Wöhler, *A* 41, 150, Heumann, *A* 41, 303, Malaguti & Sarzeau, *C R* 15, 518) The finely powdered bezoars are extracted with boiling alcohol, and the solution evaporated The crude acid then deposited is converted into sodium salt, and then into the Ba salt On recrystallising, barium lithobilate remains undissolved, and the solution of barium lithofellate is then decomposed by HCl (Roster, *G* 9, 364) Minute hexagonal crystals (containing aq) (Hoppe Seyler, *Vuchow's Arch.* 25, 528, Grattarola, *J* 1880, 831) Insol water Dextrorotatory Its salts are also dextro-rotatory, somewhat bitter in taste On distillation it gives off aromatic fumes

Reactions — 1 When heated with sugar and H_2SO_4 it gives a crimson colour (Pettenkofer's reaction) (Strecker, *A* 67, 53) — 2 Boiling HClA resinifies it — 3 Hot nitric acid gives a yellow acid, $\text{C}_{20}\text{H}_{30}(\text{NO}_2)_8$, which may be crystallised from HOAc

Salts — The sodium salt forms a pale-yellow gummy mass, exceedingly sol water and alcohol $[\alpha]_D = +18.16$ at 14.5° — BaA'_{10} aq large prisms, sol boiling water and alcohol $[\alpha]_D = +19.7$ at 15° — AgA' flocculent pp

LITHOSPERMUM ERYTHRORHIZON (M Kuhara, *C J* 35, 22) The Japanese prepare a dye called Shikon or Tokio purple from the root of this plant The colouring matter may be extracted by exhausting successively with water and alcohol Lead subacetate is added to the latter extract, and the purple pp washed and decomposed by aqueous H_2S The dye is extracted from the dried pp by means of alcohol It is resinous with green lustre Its composition may be represented by $\text{C}_{20}\text{H}_{30}\text{O}_{10}$ Its alcoholic solution is purple, and shows an absorption spectrum similar to that of alkanet Alkalis turn the solution blue, acids turn it red Baryta gives a purple pp $\text{C}_{20}\text{H}_{30}\text{BaO}_{10}$ The dye is readily oxidised to a brown scaly substance, $\text{C}_{20}\text{H}_{30}\text{O}_{11}$ Bromine forms a product of substitution, $\text{C}_{20}\text{H}_{30}\text{Br}_2\text{O}_{10}$ With PCl_5 it forms a black resinous body, $\text{C}_{20}\text{H}_{30}\text{Cl}_2\text{O}_{10}$

LITHURIC ACID $\text{C}_{15}\text{H}_{15}\text{NO}_8$ (?) [205?] Occurs as magnesium salt in some urinary

calculi of oxen (Roster, *A* 165, 104) Slender, silky needles M sol boiling water and alcohol — MgA_2 , minute monoclinic prisms, m sol boiling water, v sl sol cold water, insoluble in alcohol

LITMUS Obtained chiefly from various species of *Roccella*, *Varolaria*, and *Lecanora*, the same lichens that yield archil The blue colouring matter appears to be developed by fermentation after the mass has been treated with alkaline carbonate (Géhis, *Rev Scient* 6, 50, *J Ph* 24, 277)

When 2 pts of *Roccella tinctoria* and 1 pt of K_2CO_3 are repeatedly moistened with a solution of ammonium carbonate, the mass acquires a fine blue colour in forty days The mass is then mixed with chalk and gypsum

According to Kane (*T* 1840, 298), litmus contains azolitmin, spaniolitmin, erythrolein, and erythrolitmin Azolitmin is a reddish brown amorphous powder, which dissolves in ammonia with blue colour, and forms blue and violet lakes Spaniolitmin was not isolated by Kane, but appeared to be light red According to Kane, erythrolein is a red viscid mass, forming a purple solution in ammonia, while erythrolitmin forms deep red crystalline grains, forming a purple solution in ammonia The method employed by Kane to isolate these substances is as follows Litmus is exhausted with boiling water, the residue is acidified by HCl, again washed, and then boiled with alcohol The alcoholic solution is evaporated to dryness, and the residue extracted with ether, which leaves erythrolitmin undissolved The ether leaves erythrolein on evaporation The reddish brown powder left after boiling with alcohol is impure azolitmin

De Luynes (*C R* 59, 49), by heating orcein with aqueous NH_3 and Na_2CO_3 at 70° , obtained a blue substance, which he regarded as the blue colouring matter of litmus

According to Wartha (*B* 9, 217), cold alcohol extracts from litmus a red substance, which is not affected by acids, and the residue yields to water the blue colouring matter If the aqueous extract be evaporated, and the residue treated with absolute alcohol and some HOAc, a scarlet dye, turned purple by NH_3 , is extracted, while the residue is the pure litmus blue, left as a brown powder

A blue ethereal extract of litmus shows an absorption-band at D, a red ethereal extract shows an absorption-band extending to E (Vogel)

An aqueous solution of litmus, kept in a closed vessel, gradually becomes decolourised This is due to the action of a micrococcus, which reduces the colouring matter to a leuco-derivative, which is readily re-oxidised by air (Dubois, *Bl* [2] 49, 963, cf Bellamy, *J Ph* [5] 18, 483)

According to Förster (*Fr* 28, 428), litmus is best purified by exhausting with alcohol in the cold, digesting with water, filtering, and evaporating The residue is dissolved in water, filtered, and ppd by a mixture of alcohol and HOAc The pp is washed with alcohol, and the solution, ppn, and washing repeated as long as any reddish-violet substance is removed thereby The pp is then dried, dissolved in water, the solution filtered and ppd by feebly ammoniacal

alcohol The pp is finally washed with alcohol and dried

LIVER OF SULPHUR A name formerly applied to a mixture of the polysulphides of potassium, obtained by heating together K_2CO_3 and S in a closed vessel (v POTASSIUM, SULPHIDES or)

LIXIVIATION The application of water to solid mixtures, for the purpose of extracting the soluble parts

LOBARIC ACID $C_{17}H_{16}O_5$ A resinoid acid obtained by extracting the lichen *Lobaria adusta* with ether (Knop, *C C* 1872, 172) Warty masses, made up of thin plates Insol water and baryta water In aqueous or alcoholic NH_3 it forms colourless solutions, which turn rose-red on exposure to air KOHAq forms a yellow solution, turning brown on evaporation.

LOBELINE An alkaloid existing in *Lobelia inflata* (Bastick a Procter, *Ph* 10, 270, 456) Obtained by extracting the leaves with dilute acetic acid, and ppg with magnesia Thick, oily mass, decomposed by heat V sol water, alcohol, and ether Narcotic Forms crystalline salts with HCl, HNO_3 , H_2SO_4 , and oxalic acid Its solutions are ppd by tannin

Lobelia nicotianifolia contains also a second alkaloid, dissolved by HCl, from solutions made alkaline by NH_3 It resembles lobeline in physiological action, and, like it, exhibits no characteristic colour reactions (Dragendorff a von Rosen, *C C* 1886, 873)

LOGANIN $C_{25}H_{34}O_{14}$ [215°] A glucoside found by Dunstan and Short (*Ph* [3] 14, 1025) in the pulp in which the seeds of *Strychnos nuxvomica* are embedded Extracted by alcohol chloroform, and recrystallised from alcohol Prisms V sol water and alcohol, less sol ether, $CHCl_3$, and benzene The aqueous solution is not ppd by reagents for alkaloids, nor by lead acetate or $AgNO_3$ Not coloured by $FeCl_3$, HNO_3 , or H_2SO_4 , and $K_2Cr_2O_7$ Conc H_2SO_4 gives a red colour on warming, changing to purple Loganin does not reduce Fehling's solution Boiling dilute H_2SO_4 splits it up into glucose and loganetin Loganetin is sol water and alcohol, less sol ether and $CHCl_3$ It also gives a purple with H_2SO_4

LOKAOIC ACID $C_{22}H_{18}O_{27}$ (K) or $C_{26}H_{22}O_{31}$ (C a G) *Lokain* The colouring matter of Chinese green, obtained from the berries of the buckthorn (*Rhamnus utilis*), contains lokaonic acid (Kayer, *B* 18, 3417, cf Cloez a Guignet, *J* 1872, 1068) Lokao, the commercial article, consists of calcium and aluminum lokaonates It is decomposed by boiling with ammonium carbonate, and on adding alcohol to the filtrate ammonium lokaonate is ppd This is decomposed by oxalic acid Lokaonic acid is a deep blue or bluish black mass, which exhibits metallic lustre when rubbed It is insol water, alcohol, ether, and chloroform In aqueous alkalis it forms a blue solution, changed to red by mild reducing agents, such as H_2S Boiling dilute H_2SO_4 splits it up into a sugar, lokaose, and lokaonic acid

Salts — NH_4A (at 100°) deep blue pp, with bronze lustre when dry, sol water, insol alcohol — $(NH_4)_2A$ similar to the preceding. Begins to lose NH_3 at 40° — K_2A dark blue

powder — BaA" (at 100°) deep blue powder, insol water — PbA" (at 100°) blue black pp

Lokanic acid $C_{18}H_{16}O_{12}$ (K) or $C_{18}H_{16}O_{12}$ (C a G) *Lokatin*. Obtained by boiling lokaonic acid with dilute H_2SO_4 , dissolving the pp in NH_4Aq and ppg with oxalic acid. Violet black crystalline powder, which exhibits a bronze lustre when rubbed. Insol water, alcohol, ether, and chloroform. Alkalis form a violet solution. At 120° it becomes $C_{18}H_{14}O_{20}$. Conc H_2SO_4 dissolves it in the cold, and on adding water a reddish brown powder, $C_{18}H_{16}O_{16}$, is ppd. This dissolves in NH_4Aq , and the solution gives, with $BaCl_2$, a reddish brown pp, $BaC_{18}H_{16}O_{16}$. Lokaonic acid is decomposed by boiling conc KOH into phloroglucin and delokanic acid. Hot dilute nitric acid forms nitro phloroglucin.

Salts — NH_4A (dried at 100°) bluish black powder, sol water insol alcohol. Dyes cotton, silk, and wool deep violet without mordant. Mixed with $Na_2S_2O_3$, it dyes cotton a permanent sky blue (C a G) — BaA" (dried at 100°) blue black powder, insol water and alcohol. PbA" deep blue powder, insol water and alcohol.

Delokanic acid $C_{15}H_{10}O_8$. Formed as above. Brown powder, insol water, sol alcohol. Its solution in alkalis is cherry red. It reduces Fehling's solution with difficulty in the cold.

Lokaose $C_{18}H_{16}O_8$. A sugar formed by hydrolysis of lokaonic acid. It reduces chloride of gold and Fehling's solution in the cold. It reduces half as much CuO as glucose.

LOPHINE v vol 1 p 474

LOTURINE [234°] Occurs to the extent of 24 p.c., together with 02 p.c. of colloturine and 06 p.c. of loturidine, in lotur bark from *Symplocos racemosa*, growing in India (Hesse, B 11, 1042). The alkaloids are extracted from the bark by hot alcohol, and are converted into acetates. Loturine and colloturine are ppd from the neutral solution by potassium sulphocyanide, leaving loturidine in solution. The crystalline pp is decomposed by Na_2CO_3 , and the alkaloids are extracted with ether and recrystallised from alcohol. The efflorescent crystals of loturine are separated mechanically from the non efflorescent crystals of colloturine.

Properties — Long prisms. May be sublimed. Insol water, sol alcohol, acetone, ether, and chloroform. Its acid solutions exhibit violet fluorescence. It gives no colour with $FeCl_3$, conc H_2SO_4 , HNO_3 , or bleaching powder followed by ammonia. Its hydrochloride forms white prisms. The hydrochloride, nitrate, and chromate crystallise in needles. The platinochloride is a yellow pp.

Colloturine. Long prisms, may be sublimed. Its solutions in aqueous HCl and H_2SO_4 fluoresce violet. Its aurochloride is a yellow amorphous pp.

Loturidine. Extracted from the filtrate from the ppd sulphocyanides of loturine and colloturine by adding NH_3 and shaking with ether. Yellowish brown amorphous mass. Forms amorphous salts. Its solutions in dilute mineral acids fluoresce violet.

LOXOPTERYGINE $C_{22}H_{22}N_2O_2$ [81°] Occurs, together with another alkaloid and tannin, in red Quebracho bark from *Loxopterygium Lorentzii* (Hesse, A 211, 274). Extracted by alcohol,

the alcoholic solution being evaporated, the residue treated with aqueous $NaOH$ and the alkaloids extracted with ether. The acetic acid solution of the alkaloids is mixed with potassium sulphocyanide, which ppts one alkaloid and leaves the loxopterygine in solution, whence it is ppd by ammonia. White amorphous mass, sol cold water, v sol alcohol, ether, chloroform, and benzene. Its solutions exhibit alkaline reaction. Tastes intensely bitter. Conc H_2SO_4 and a little $K_2Cr_2O_7$ gives a violet colour.

LUPANINE $C_{14}H_{12}N_2O$. Occurs in the seeds of the blue lupine (*Lupinus angustifolius*), from which it is extracted by alcohol containing HCl (Hagen, A 230, 370). Viscid yellow liquid with green fluorescence, characteristic smell, and very bitter taste. Sol water, but separates on warming, m sol cold alcohol, v sol ether and chloroform. Not volatile with steam. Strongly alkaline, fuming with HCl .

Salts — $BHCl$ 2aq [127°] M sol water and alcohol, insol ether. The base is set free from this salt by KOH but not by NH_3 — BHl 1½aq yellow crystals (from hot water), sol. CS, insol alcohol and ether — $BHCyS$ ½aq pale yellow crystals, v sol hot alcohol, sl sol water, insol ether — $BH PtCl_3$ 3½aq — $BH AuCl_4$ splendid yellow needles, sl sol ether, insol water and alcohol.

Methyl iodide $BMeI$ [215°] Crystals, sl sol water, insol alcohol and ether. The methyl hydroxide is formed by treatment with Ag_2O but not with KOH .

Methyl chloride $BMeCl$ 2aq [128°] Extremely deliquescent crystals, insol ether — $BMeHPTCl_3$ ½aq red crystals, insol ether, sl sol alcohol — $BMeAuCl_4$ ½ $AuCl_3$ lemon yellow needles.

LUPETIDINE v DI METHYL PYRIDINE HEXAHYDRIDE

LUPINIDINE v LUPININE

LUPININ $C_{25}H_{32}O_{16}$ *Lupinin*. A glucoside in the buds of the yellow lupine (*Lupinus luteus*). Extracted by 50 p.c. alcohol (Schulze a Barbieri, B 11, 2200). Slender yellowish white needles (containing 7aq). Sl sol water and alcohol. Alkalis form a deep yellow solution. Its ammoniacal solution gives a lemon yellow pp with lead acetate. Boiling with water or dilute acids splits it up into glucose and lupigenin.

Lupigenin $C_8H_{12}O_6$. Formed as above. Minute yellow needles, insol water, sl sol alcohol. Forms a deep yellow solution in aqueous NH_3 — NH_4A ¼aq lemon yellow powder.

LUPININE $C_{24}H_{30}N_2O_2$ [68°] (256°) An alkaloid in the seeds of the yellow lupine (*Lupinus luteus*), and extracted by alcohol containing HCl . The extract is evaporated to a syrup, treated with KOH , and shaken with light petroleum. The petroleum is shaken with aqueous HCl , the solution treated with KOH , and the alkaloid extracted by ether, from which it is recrystallised (Baumert, L V 27, 15, cf Beyer, L V 14, 161, v also lupinidine, *infra*).

Properties — Trimetric crystals, may be distilled in a current of hydrogen. Tastes bitter. V sol cold water and alcohol, less sol hot water, v sol benzene, chloroform, and CS_2 . Strong base, liberating NH_3 from its salts and fuming with HCl . In its solutions tannin gives a flocculent pp, phospho-molybdic and phos-

photungstic acids give yellow pps. Iodine gives a dirty brownish red pp.

Reactions — 1 HI gives $C_{10}H_{16}N_2O_2I_2$ — 2 With fuming HCl at 200° it forms anhydro lupinine $C_{10}H_{14}N_2O$ and finally di-anhydro lupinine $C_{10}H_{12}N_2$ (Baumert, A 214, 366) — 3 P_2O_5 heated with the hydrochloride at 180° forms oxylupinine, which with platinum chloride forms $C_{10}H_{10}N_2O_2.H_2PtCl_6$, crystallising in yellow plates. If the mother liquor be heated with P_2O_5 to a higher temperature anhydro lupinine is formed (Baumert, A 214, 360) — 4 Sodium dissolves in fused lupinine, but the product is decomposed by water into NaOH and lupinine (Baumert, B 15, 631)

Salts — $B''H_2Cl_2$ large trimetric crystals — $B''HNO_3$ crystals, v sol water and alcohol — $B''H_2SO_4$ deliquescent prisms — $B''(H_2AuCl_4)_2$ needles, v sl sol water — $B''H_2PtCl_6$ crystals, sol water

Acetyl derivative $C_{10}H_{14}Ac_2N_2O_2$ Obtained by heating lupinine with $AcCl$ or with Ac_2O and $NaOAc$ (Baumert, A 224, 313) Oil — $B''H_2PtCl_6$ orange trimetric plates

Methylo-iodide $B''Me_2I_2$ white hexagonal plates, sl sol alcohol (Baumert, B 14, 1221)

Methylo-chloride $B''Me_2Cl_2$ pearly plates — $B''Me_2PtCl_6$ orange red needles — $B''Me_2Au_2Cl_6$ yellow pp

Ethyl-iodide $B''Et_2I_2$ hexagonal plates (Baumert, B 14, 1321) Decomposed by AgO, but not by KOH. From it may be obtained $B''Et_2PtCl_6$ and $B''(Et_2AuCl_4)_2$ [70°]

Anhydro-lupinine $C_{10}H_{14}N_2O$ Formed by heating lupinine with fuming HCl at 150° – 200° (Liebscher, B 14, 1880) Liquid which cannot be distilled. Oxidised by air. Smells like conine. Forms crystalline salts — $B''H_2PtCl_6$ red tables, v sol water

Di-anhydro-lupinine $C_{10}H_{12}N_2$ (220°) From lupinine and conc $HClAq$ at 200° (Liebscher, A Baumert, A 214, 371) Oil. Readily oxidised by air — $B''H_2PtCl_6$ dark red crystals, v sol warm water

Lupidine $C_{10}H_{16}N_2$

Preparation — Lupine seeds are extracted with dilute alcohol acidified with H_2SO_4 , and, after as much fat as possible has been removed, the sulphate solution is evaporated to a syrup and the residue triturated with absolute alcohol, when it solidifies to a mass of crystalline plates. This mass is again triturated with absolute alcohol, when a white crystalline meal of acid lupidine sulphate separates. The mother liquors are treated again in the same manner, until the residue, on trituration with alcohol, either remains liquid or at least redissolves on washing with absolute alcohol, in this case it consists for the most part of lupinine sulphate. The mother liquors containing the lupinine sulphate are freed from alcohol, dissolved in water, and ppd by $BaCl_2$. The filtrate now contains chiefly lupinine chloride, from which any lupidine salt present can be ppd as platinum chloride (G Baumert, A 225, 365) Lupidine is obtained by decomposing the acid sulphate by one of the stronger bases, shaking with ether, and distilling in a current of hydrogen. The fact that this alkaloid distils over in a stream of hydrogen, between the wide limits of 250° – 320° ,

is explained on the assumption that the lupidine got from the seeds of the yellow lupine is a mixture of crystallisable hydrate (see below) and a liquid anhydride. The formula $C_{10}H_{16}N_2$ is calculated from analyses of its salts

Properties — Thick yellow oil of intensely bitter taste and disagreeable hemlock like smell, but this smell is probably due to a decomposition product. Strong base. Very easily oxidisable by the air when in contact with acids or with KOHAq, but not with NH_4Aq . Gives no acetyl derivative or ethylo iodide (Baumert, A 225, 365) Lupidine is a feeble poison, acting on frogs like curare (Kobert, A 227, 219)

Salts — $B''HCl$ yellow deliquescent crystals — $B''HI$ rather large crystals resembling alum, v sol hot water, m sol alcohol — $B''H(?)$ Formed by heating lupidine with EtI . Thus lupidine resembles berberine in giving a hydroiodide when heated with EtI (Baumert, A 227, 207) — $B''H_2SO_4$ minute crystals, v sol water, v sl sol alcohol — $B''H_2PtCl_6$ 2aq trimetric crystals, $a b c = 885 \ 1 \ 1171$ (Lueddecke, Z K 12, 297), sl sol water, m sol $HClAq$, v sol dilute alcohol (Baumert, A 225, 365)

Hydrate $C_{10}H_{16}N_2aq?$ The liquid and crystalline portions of the lupine alkaloid are to be considered as modifications of the same base (the hydrate being crystalline), since both yield the same double salt with $PtCl_6$. The existence of a hydrate, stable only in the cold, may also explain why cold aqueous solutions of lupidine, lupinine, and conine become milky on heating. A further proof of the correctness of this view is that the yellow oil only (probably the pure anhydride $C_{10}H_{14}N_2$) was got on treating the lupidine with P_2O_5 (Baumert)

Lupinine A crystalline alkaloid, called by this name, was obtained by Betteli (G 11, 240) by extracting white lupine (*Lupinus albus*) with alcohol. It forms white needles, v sol water. An alcoholic extract of the plant has been used with some success in cases of intermittent malarial fever. Solutions of the alkaloid give white pps with tannin, $HgCl_2$, and $AgNO_3$, in the last case reduction soon takes place. Platine and gold chlorides and picric acid give canary yellow pps. $HIAq$ containing iodine gives a red amorphous pp. It does not reduce Fehling's solution. Boiling KOH gives off ammonia

Lupinine A liquid alkaloid, also called lupinine, was obtained by Campani (G 11, 237) from the white lupine. It boiled between 210° and 218° and was poisonous

The young shoots of the yellow lupine contain an alkaloid called ARGEMINE ($q v$). The danger to sheep from eating lupines appears to be due to another substance 'ictiogen,' perhaps produced by a parasite growing on the lupines (Baumert, Ar Ph [3] 24, 49, Kobert, A Liebscher, J 1886, 1696, Kühn, *ibid*)

LUPULIN The yellow granular aromatic powder situated at the base of the cones of the hop and forming from 8 to 18 p.c. of the cones. It contains hop oil, a resin, a nitrogenous substance, a gummy substance, and a bitter principle. Lerner (D P J 179, 54) extracted the bitter principle by ether, shook the ether with aqueous KOH, ppd by $CuSO_4$, decomposed the copper compound with H_2S , and recrystallised from ether. He describes

the 'lupulin' so obtained as large white prisms, insol water, v sol alcohol, ether, and chloroform. The alcoholic solution, diluted with water, had a bitter taste and acid reaction. Lerner also obtained a compound $[110^\circ]$ which likewise formed a copper salt. Lupulic acid, as described by Bungener, appears to be Lerner's lupulin Isleib (*Ar Ph* [3] 16, 345) found a 'pseudo glucoside' $C_{10}H_{16}O_{10}$ which he extracted by cold water. The extract was treated with animal charcoal, the charcoal dried and exhausted with 90 p c alcohol, and the yellow solution partially evaporated. A brown resin $C_{10}H_{16}O_2$ is deposited while a bitter uncrystallisable substance remains in solution. From the aqueous solution of this bitter substance ether extracts only the bitter principle (which amounts to 0.04 of the hops) and leaves in the water $C_{10}H_{16}O_2$, a tasteless product of the oxidation of oil of hops $C_{10}H_{16}O$. The bitter principle, according to Isleib, forms a yellow solution in alkalis, and is split up by boiling dilute H_2SO_4 into lupuluretin $C_{10}H_{16}O_2$ and lupulic acid $C_{10}H_{16}O_4$. This lupulic acid forms a crystalline barium salt $BaC_{10}H_{16}O_4 \cdot 5aq$. According to Bissell (*Ph* [3] 8, 508) lupulin does not contain all the active principle of the hop. The bitter principle may usually, but not always, be ppd from an infusion of hops by lead acetate (Allen *Ar* 13, 43). According to Hayduck (*U C* 1887, 694), when hops are exhausted with ether, and after evaporating the ether, the residue is treated with alcohol, a white wax is left behind. The alcoholic solution gives a yellow pp with lead acetate, and the filtrate contains two resins, one soluble, and the other insoluble, in light petroleum.

Lupulic acid $C_{10}H_{16}O_4$ [93°]

Preparation—By extracting hop flour with light petroleum spirit, distilling off the solvent, cooling, filtering off the black liquid from the crystals formed, and recrystallising the crude acid from alcohol and petroleum spirit.

Properties—Colourless prismatic crystals, rapidly altered by exposure to the air. V sol alcohol, ether, benzene, $CHCl_3$, CS_2 , and the ethereal oil of the hop, sol petroleum spirit, insol water. Easily reduces ammoniacal $AgNO_3$. Exposed to the air is transformed into a yellowish resin. This product of oxidation is sol water, to which it imparts an intense bitter taste, and this resin is the bitter principle of the hop—CuA'. A crystalline powder (Bungener, *Bl* [2] 45, 487).

LUPULINE v HOPPING

LUTEIC ACID $C_{10}H_{16}O_{12}$ (?) [274°] S

0.09 in the cold, 0.3 at 100° S (alcohol) 4.2 in the cold S (ether) 37. A yellow colouring matter prepared from the flowers of *Euphorbia Cyparissias* (Hohn, *Ar Ph* [2] 140, 218). The fresh flowers are exhausted with alcohol of 60 p c, the greater part of the alcohol distilled off, and the residual liquid filtered and ppd with basic lead acetate. The pp is suspended in water, decomposed by H_2S , and filtered. The filtrate is evaporated over H_2SO_4 , and the yellowish crusts that separate are recrystallised from ether-alcohol, and then from hot water containing 4 p c of alcohol. Slender yellow needles. Has no smell and a bitter taste. Sublimes at 220° . Its solutions exhibit acid reaction. Lutein somewhat resembles luteolin and,

like the latter, gives protocathechuic acid by potash fusion. It dissolves in caustic and carbonated alkalis forming yellow solutions. It reduces $AgNO_3$, mercurous nitrate, and Fehling's solution on heating. $FeCl_3$ gives a green colour, changing to reddish brown on further addition of $FeCl_3$. Conc H_2SO_4 dissolves luteic acid, but water reprecipitates unaltered. Boiling dilute H_2SO_4 does not attack it.

LUTEIN This name is given by Thudichum (*Pr* 17, 253) to the yellow substance obtained by Piccoli and Lieben (*Z* 1868, 645) from the ovary of the cow, and called by them 'hæmolutein' (*cf* Holm, *Z* 1867, 779). It occurs also in the human ovary. The same or a similar substance occurs in butter, yolk of egg, serum of blood, in some diseased swellings, in carrots, &c. It occurs also in the retina of the eyes of fowls (Capranica, *J Th* 1877, 317, Kuhne, *J Th* 1877, 317). According to Maly (*C C* 1881, 485, *M* 2, 359), the lutein in yolk of egg of hens and shrimps is a mixture of vitello lutein and vitello rubin. If the yolk is extracted with alcohol, and the extract treated with hot baryta water, vitellorubin is ppd while vitellolutein remains in solution. Or the yolk extract may be boiled with a little acid, and the ppd albumen treated with petroleum which extracts vitellolutein, while CS_2 extracts vitellorubin from the residue. Vitellorubin contains no nitrogen. It is red, and forms a Mg compound sol ether, chloroform, and CS_2 , but ppd by alcohol in red flakes. An alcoholic solution of vitellorubin shows a broad but weak absorption band including the line F. Vitello lutein forms a yellow alcoholic solution which exhibits two narrow absorption bands, one including F, the other between F and G. It contains no nitrogen.

LUTECHROMIUM SALTS, v vol II p 160

LUTECOBALTIC SALTS, v vol II p 228

LUTEOLIN $C_{15}H_{18}O_4$ (Moldenhauer, *A* 100, 180) or $(C_{15}H_{18}O_4)_2$ 3aq (Schutzenberger a Paraf, *A Suppl* 1, 256) [320°] S 0.07 in the cold, 0.2 at 100° S (alcohol) 2.7 S (ether) 16. The yellow colouring matter of weld (*Reseda luteola*) (Chevreul, *J Chim Méd* 6, 157). Obtained by boiling weld with water (16 pts) mixed with alcohol (1 pt), filtering, evaporating, dissolving the yellow flakes in alcohol, and pouring into water. It is recrystallised from a mixture of water and glycerin (Rochleder, *J pr* 99, 433). Yellow, four sided needles in radiate groups. May be sublimed. Partially decomposed on fusing. Has a slightly bitter, astringent taste. Reddens litmus slightly. Dissolves with deep yellow colour in caustic and carbonated alkalis. Conc H_2SO_4 forms a yellow solution, whence water reprecipitates it. Potash-fusion gives CO_2 , phloroglucin, and protocathechuic acid. Very dilute $FeCl_3$ gives a green pp, excess of $FeCl_3$ forms a brownish red solution. Alcoholic lead acetate gives a pp $C_{15}H_{18}PbO_4$ aq.

(β -LUTIDINE $C_{15}H_{18}N$ Ethyl pyridine (?) Mol w 107 (166°) SG ρ 956 (W), 959 (C) VD 3.79 (calc 3.70) S 4 (W). Obtained, together with homologues, by distilling cinchonine with KOH, a mixture of some ten bases is obtained and these are separated by fractional distillation (Greville Williams, *J*

1855, 549, 1864, 487, *Pr* 18, 305, Oechsner de Coninck, *C R* 92, 413, *Bl* [2] 85, 296, *R T C* 1, 132, *A Ch* [5] 27, 462, 488) The same base appears to be formed by distilling with zinc dust the syrupy acid formed by oxidising cinchonine with chromic acid mixture (Weidel & Hazura, *M* 3, 780) This, or an isomeric ethyl pyridine, is formed by distilling brucine with KOH (O de Coninck, *Bl* [2] 42, 100) Liquid, sl sol water, sol alcohol and ether The aqueous solution does not become turbid on warming, the base being more soluble in hot than in cold water It appears to form an unstable hydrate $C_8H_9NH_2O$ Smells somewhat like nicotine Physiologically it is a violent poison and stops tetanus produced by strychnine (Greville Williams & Waters, *Pr* 32, 162) By chromic acid mixture or by $KMnO_4$, it is oxidised to pyridine carboxylic (nicotinic) acid [231°] When heated with sodium it appears to form di (β) lutidine $C_{16}H_{18}N_2$ (Greville Williams, *Pr* 33, 159) Chlorine passed into (β) lutidine containing dissolved iodine appears to form tri-chloro (β)-lutidine Chloroacetic acid gives $C_8H_7NClCH_2CO_2H$ (163°) forming ($C_8H_7NO_2Cl$), $PtCl_2$, 2aq (Pictet, *J* 1882, 1079)

Salts — $BHCl$ Very deliquescent crystals — B_2HBr — $B_2H_2PtCl_4$, orange red leaflets Its ppp is not retarded by presence of excess of HCl (difference from bone oil lutidine) Hot water decomposes it forming B_2PtCl_4 crystallising in pale yellow leaflets — B_2PtCl_4 From platinous chloride (1 pt) and lutidine (1 pt) combination takes place with rise in temperature (of 70°), the product being a hard brittle mass (Williams) — B^+HAuCl_4 Yellow pp On boiling with water it forms yellow $B^+HAu_2Cl_4$ and ultimately B^+AuCl_4 , a red crystalline powder — $B_2H_2PdCl_4$, garnet red prisms, obtained by mixing solutions of the hydrochloride and of palladium chloride (W) Decomposed at 100° , giving off HCl and leaving B_2PdCl_4 , sl sol water — $B_2H_2Cl_4U_2O_2Cl_4$ Formed from uranyl chloride and (β) lutidine hydrochloride Yellow — $B_2H_2SO_4(U_2O_2)(SO_4)_2$ From uranyl sulphate and (β) lutidine sulphate Small yellow needles —Picrate $B^+C_6H_7(NO_2)_3OH$ Yellow needles (Williams, *Pr* 33, 159)

Combinations — B_2CuSO_4 , 4aq When (β) lutidine is added to a solution of cupric sulphate a copious pale green pp is formed, which dissolves in excess, forming a rich blue liquid, which deposits blue prisms of B_2CuSO_4 , 4aq — B_2AgNO_3 Formed by ppg silver nitrate solution with (β)-lutidine and recrystallising from alcohol

Di (β) lutidine When sodium is warmed with (β) lutidine dissolved in toluene a product is obtained from which it is possible to get a platinochloride containing a percentage of platinum corresponding to the formula $C_{16}H_{18}N_2H_2PtCl_4$ (Williams, *C N* 44, 308)

Tetra- (β)-lutidine When sodium is warmed with (β)-lutidine a violent action takes place, and from the product a platinochloride may be obtained containing a percentage of platinum corresponding to the formula $C_{32}H_{36}N_4H_2PtCl_4$ (Williams)

Hydride of (β)-lutidine C_8H_7N Greville Williams found that sodium amalgam had no

action on (β) lutidine According to Oechsner de Coninck (*Bl* [2] 42, 121), however a hexahydride (155° – 160°) may be formed by the action of sodium on an alcoholic solution of the base (cf Wyschnegradsky, *B* 13, 2401) It combines with MeI and the product, when distilled with KOH , yields C_8H_7N (160°) It also combines with EtI , and the product treated with potash yields a tertiary base boiling at 175°

Lutidine C_8H_7N (173°) Obtained by distilling strychnine with zinc dust (Seichlani & Magnanini, *G* 12, 445) Yellow liquid, insol water, sol alcohol and ether Smells like pyridine Gives with sodium phosphomolybdate a dark yellow pp, sol ammonia Potassium mercuric iodide gives a yellow amorphous pp Mercuric chloride gives a white pp Iodine in $KIAq$ gives a crimson pp, insol dilute $HClAq$

Coal tar lutidines v **DI METHYL PYRIDINE**

Isomerides v **ETHYL PYRIDINE** and **DI METHYL-PYRIDINE**

LUTIDINE CARBOXYLIC ACIDS v **DI METHYL PYRIDINE CARBOXYLIC ACID**

LUTIDINIC ACID v **PYRIDINE-DI CARBOXYLIC ACID**

LUTIDONE v **OXY DI METHYL PYRIDINE**

LUTIDO-STYRIL v **OXY DI METHYL PYRIDINE**

LUTIDYL-QUINOLINE v **DI METHYL PYRIDYL QUINOLINE**

LYCACONITINE v **ACONITE ALKALOIDS**

LYCINE The base from *Lycium barbarum* to which this name was applied has been shown to be identical with betaine

LYCOPODINE $C_{11}H_{12}N_2O$, [115°] Occurs in *Lycopodium complanatum* (club moss), from which it is obtained by extracting the root with alcohol, evaporating the extract, dissolving the residue in water, adding lead subacetate, removing excess of lead by H_2S , adding $NaOH$, and shaking with ether When the ether is evaporated the alkaloid is left, and may be purified by dissolving in $HClAq$ and ppg with conc $NaOHAq$ (Bodeker, *A* 208, 363) Monoclinic prisms, v sol most menstrua Taste: bitter — $B^+H_2Cl_4$, aq glassy hexagonal crystals (Solling, *J* 1884, 463) — $B^+(HAuCl_4)_2$, aq small glistening yellow needles

LYCOPODIUM BITTER The alcoholic and aqueous extracts of *Lycopodium clavatum* are evaporated, and the residues mixed and extracted with water The solution is treated with lead acetate and subacetate, filtered, freed from lead by H_2S and evaporated The residue is then washed with alcohol and dissolved in water From this solution lead subacetate ppt the bitter principle, and the pp is then suspended in water and decomposed by H_2S The filtrate is concentrated, freed from sugar by careful fermentation with yeast, dried, and extracted with absolute alcohol, which leaves the bitter principle on evaporation (Kamp, *A* 100, 300) Slender needles (from water) v e sol water, alcohol, and ether Extremely bitter Neutral to litmus Contains no nitrogen Colours tincture of iodine orange red Does not reduce Fehling's solution, but after boiling with dilute H_2SO_4 , it does so

LYCOPODIUM OIL The oil from the fresh spores of club moss contains phytosterin $C_{27}H_{48}O$, together with glyceryl ethers of lycopodine

$C_{15}H_{31}O_2$, oleic, arachic, stearic, and palmitic acids (Bukowski, *Chem Zeit* 1889, 174, cf Langer, *Ar Ph* [3] 27, 625)

LYCOPESIN $C_{21}H_{40}O$ [170°] Extracted from *Lycopodium chamæcyparissus* by alcohol, and remaining in the mother liquor after lyco stearone has separated (Kamp, *A* 100, 300) Minute crystals, almost insol boiling water, v sol alcohol and ether, v sol cold alkalis

LYCOSTEARONE $C_{21}H_{40}O_2$ [75°–100°] An amorphous tasteless substance, which may be extracted by alcohol from *Lycopodium chamæcyparissus* (Kamp, *A* 100, 300)

parissus (Kamp, *A* 100, 300) Insol cold water, sol hot water forming a jelly on cooling SL sol cold alcohol and ether, v sol alkalis

LYDINE A violet dye obtained by pouring a solution of aniline (100 g) in fuming $HClAq$ (100 g) diluted with water (120 cc) into a solution of K_2FeCy_6 (90 g) in water (850 cc), and heating the mixture to boiling. The product is dissolved in dilute oxalic acid and ppt by alkalis Violet powder, sol alcohol, sl sol ether and benzene, insol water Poisonous (Guyot, *C R* 69, 829) Possibly identical with mauvine.

M

MACLEYIN $C_{10}H_{15}NO_3$ (?) [201°] S ($CHCl_3$) 7 S (ether) 1 An alkaloid in *Macleya cordata* the Japanese *Tachibaku* (Eijkman, *R T C* 3, 182, *Ph* [3] 13, 87) Extracted from the root by alcohol and dilute H_2SO_4 , and separated from sanguinarin by ether in which it is very slightly soluble Prisms Almost insol water and alkalis, v sl sol hot alcohol Its salts have a bitter taste Its chemical properties resemble those of protopine Conc H_2SO_4 gives a faint yellow changing through violet to green Fumes of nitric acid give an ultramarine colour H_2SO_4 and a little $K_2Cr_2O_7$ give a blue colour

Salts — $B'HCl$ Prisms S 7 — $B'H.PtCl_2$ 2aq yellowish whitepp — $B'H Cr_2O_7$ orange prisms — Hydroiodide crystalline sphres — Normal sulphate colourless needles Thiocyanate needles — Acetate needles — Acid oxalate crystals — Acid tartrate needles — Benzoate [166°] long needles

MACLURIN $C_{11}H_{19}O_6$ (H a P), $C_{11}H_{19}O_6$ (L) *Morintarnic acid* [200°] (Wagner) S 52 at 14° Occurs, together with morin (q v), in old fustic, the wood of *Morus tinctoria*, from which it is obtained by extracting with water The evaporated extract deposits morin, and from the filtrate maclurin may be pptd by HCl (Hlasiwetz a Pfander, *A* 127, 352, *J pr* 94, 65, Löwe, *Fr* 14, 117, Benedikt, *A* 185, 114) Almost pure maclurin is also found in patches in the wood (Wagner, *J* 1850, 529)

Properties — Yellow, crystalline powder (containing aq), v sol alcohol and ether, sl sol water Above 270° it gives CO_2 and pyrocatechin Its aqueous solution is pptd by alkaloids, gelatin, and albumen $FeSO_4$ gives a greenish black pp Its alkaline solution turns brown in air It dyes mordanted cotton pale brownish-yellow Boiling conc KOH aq gives phloroglucin and protocatechuic acid $C_{11}H_{19}O_6 + H_2O = C_6H_6O_3 + C_5H_8O_3$ Dilute H_2SO_4 at 120° does the same Bromine gives tri bromo maclurin $C_{11}H_5Br_3O_6$ aq minute needles Conc H_2SO_4 at 190° gives brown $C_{10}H_{11}O_5$ (Hlasiwetz, *A* 143, 308) Reduction with zinc and H_2SO_4 gives phloroglucin and machronin When its solution is heated with sodium amalgam and the product acidified, ether extracts phloroglucin and amorphous $C_{11}H_{19}O_6$, which is sol water and alcohol, is pptd by lead acetate, gives a grass green colour with $FeCl_3$, and reduces $AgNO_3$ and Fehling's solution.

Salts — $PbC_{11}H_{19}O_6$ aq yellow crystals — $C_{11}H_{19}O_6$ 3 PbO (L)

Acetyl derivative $C_{11}H_{17}AcO_6$ 14aq viscid oil

Macronin $C_{11}H_{19}O_6$ 3aq (?) Formed by the action of zinc and dilute H_2SO_4 on maclurin, and separated from phloroglucin by pptn with lead acetate The pp is decomposed by H_2S Colourless spangles composed of tufts of minute needles (from dilute alcohol) V sl sol water and alcohol, m sol ether Turns dark blue when exposed to air or when heated The hot aqueous solution turns violet blue in air, and then yields an indigo blue pp with HCl $FeCl_3$ gives, in a very dilute alcoholic solution, a violet-red colour, changing to blue Alkaline solutions also turn blue on exposure to air $AgNO_3$ gives a violet colour with reduction of silver Reduces hot Fehling's solution Conc H_2SO_4 gives an orange solution, turning emerald green on warming or diluting

Rufumoric acid $C_{10}H_{17}O_6$ (?) Obtained by boiling maclurin with dilute HCl (Wagner, *J* 1851, 420) Dark red mass, v sol alcohol, m sol water, v sl sol ether Forms a crimson solution in alkalis

MACROCARPINE v THALICTRINE

MADDER v OXY ANTHRACINONES

MAGDALA RED $C_{10}H_7N_3Cl$ Formed by heating naphthalene azo (a) naphthylamine with (a) naphthylamine hydrochloride (Hofmann, *B* 2, 374, 412) Appears to be a compound of amidonaphthazine $C_{10}H_5(NH) \begin{smallmatrix} \diagup N \\ | \\ \diagdown N \end{smallmatrix} C_{10}H_5$ allied to saf-franine, viz

$C_{10}H_5(NH_2) \begin{smallmatrix} \diagup N \\ | \\ \diagdown NCl(C_{10}H_5NH_2) \end{smallmatrix} C_{10}H_5$ (Julius, *B* 19, 1365)

MAGNESIA *Magnesium oxide* MgO (q v)

MAGNESIA ALBA A basic carbonate of Mg, varying somewhat in composition, v Carbonates vol 1 p 699

MAGNESIA USTA. A name given in pharmaceutical nomenclature to MgO prepared by calcining *magnesia alba*

MAGNESIUM At w 24 Mol w probably same as at w (v infra) [c 450°] (Devillea Caron, *A* 101, 359), [c 500°] (Ditte, *B* 73, 108), [between 700° and 800°] (V Meyer, *B* 20, 497) Scarcely volatilises in CO at red to white heat (V. M. Lc.). S G. 1.69 to 1.71 at 17° (Kopp);

1 77 at 0° (Wurtz, *Am Ch* 1876 (March)) S H 20° to 51° 245 (Kopp, *T* 155, [1] 71) C E at 40° 00002694 (Fizeau, *C R* 68, 1125) T C (Ag=100) 84.3 (Lorenz, *W* 18, 422) E C (Hg at 0°=1) 22.84 at 0°, 16.34 at 100° (Lorenz, *W* 18, 422, 582) S V S c 141 Characteristic lines in emission spectrum 5527.4, 5183, 5172, 5166.7 (Thalén, Upsala, 1868, v also Liveing & Dewar, *Pr* 44, 241)

Occurrence—The metal does not occur native, but compounds of Mg are very widely distributed, and are found in some places in large quantities, they usually accompany Ca compounds *Magnesite* $MgCO_3$, *dolomite* a mixture of the isomorphous carbonates of Mg and Ca, *kieserite* $MgSO_4 \cdot H_2O$, *carrollite* $MgCl_2 \cdot KCl \cdot 6H_2O$, various silicates e.g. *asbestos* $(MgCa)SiO_3$, *spinnelle* $MgAl_2O_3$, &c, occur as minerals. Borate, phosphate, sulphate, chloride, nitrate, and hydroxide, of Mg are found in mineral waters, and some of them in sea water. Mg compounds, chiefly phosphate and carbonate and compounds of organic acids, are found in plants and animals.

Magnesia alba (a basic carbonate of Mg) was introduced as a medicine in the beginning of the 18th century, the name is said to have been given merely in contradistinction to *magnesia nigra*, a term applied to the mineral pyrolusite because of its colour and supposed magnetic properties. Magnesia was long supposed to be the same, or nearly the same, as lime. Hoffmann, Bergmann, Margraff, and especially Black (1755) established the characteristic properties of magnesia. Black showed *magnesia alba* to be a compound of fixed air (CO_2) with an earthy base, to this base he gave the name magnesia. In 1800 Davy obtained a small quantity of a metal by reducing magnesia at white heat by vapour of K, in 1830 Bussy obtained the metal in greater quantity and purity, by heating together $MgCl_2$ and K.

Formation—1 By heating together $MgCl_2$ and K in a Pt crucible, and washing out KCl by water (Bussy, *Buff*, *P* 18, 140, Liebig, *P* 19, 137)—2 By electrolysis fused $MgCl_2$ (Bunsen, *A* 82, 137), or a mixture of $MgCl_2$ and KCl with a little NH_4Cl (Matthiessen, *C J* 8, 107)—3 By reducing a molten mixture of $MgCl_2$, CaF_2 , KCl, and NaCl, by Na (Deville & Caron, *C C* 1863 993)—4 By reducing MgF_2 , NaF by Na (Tissier, *C R* 56, 848)—5 By electrolysis a mixture of $MgSO_4$ and $(NH_4)_2SO_4$ dissolved in water (Gerhard, *English Patents*, No 16,691, 1884)—6 Puettner patented a method for reducing $MgCO_3$ by heating with Fe_2O_3 and C (*German Patents*, No 31,319)—7 Lauterborn reduces a double Mg Na cyanide by Zn (*German Patents*, No 39,915).

Preparation—1 The upper part of a porcelain crucible is divided vertically into two halves by a thin porcelain plate, the crucible is closed by a porcelain cover, through which pass the carbon electrodes of a battery of 8-10 Bunsen cells, one electrode on each side of the dividing vertical plate, indentations are cut in the negative electrode, the crucible is heated to redness, then filled with fused $MgCl_2$, and the current is passed through the molten mass. The Mg lodges in the cavities of the negative electrode, and is thus prevented from floating to the surface,

the dividing plate serves to prevent contact between the Cl₂ evolved at the positive electrode, and the Mg (Bunsen, *A* 82, 137)—2 A mixture of $MgCl_2$ and KCl, prepared by evaporating a mixture of the salts, in the ratio 4 $MgCl_2$, 3 KCl, to dryness, is mixed with a little NH_4Cl , and is electrolysed when fused in the apparatus described in 1, as Mg is heavier than the molten mixture of salts it is not necessary to form serrations in the negative electrode (Matthiessen, *C J* 8, 107)—3 A mixture of 6 pts dry $MgCl_2$, 1 pt of a mixture of 7 pts NaCl and 9 pts KCl, 1 pt powdered CaF_2 , and 1 pt Na carefully freed from oil, is heated to bright redness in a covered earthen crucible. The Mg, which separates in globules, is heated nearly to whiteness in a boat of compact charcoal, placed in an inclined tube of the same material through which is passed a stream of dry H, the Mg condenses on the upper part of the tube, it is then melted into large globules with a flux of $MgCl_2$, NaCl, and CaF_2 . To free the Mg from C, Si, and N, it may be distilled in a current of H at a high temperature (Deville & Caron, *A* 83, 137, for an apparatus in which the distillation may be conveniently conducted v Sonstadt, *J pr* 90, 307). In a later trial D & C used 7 pts $MgCl_2$, 4.8 pts CaF_2 , and 2.3 pts Na (*C C* 1863 993)—4 Wöhler (*A* 101, 362) reduces a fused mixture of $MgCl_2$, NaCl with excess of NaCl by Na, the mixture is obtained by adding excess of NaCl to $MgCl_2$ aq and evaporating to dryness. The amount of $MgCl_2$ in the mixture must be known in order that the quantity of Na to be used may be calculated, for every 1 pt $MgCl_2$, about $\frac{1}{2}$ pt Na should be used.

Properties—A silver white, very lustrous, moderately hard, metal, malleable, not very tenacious, may be filed and polished, S G c 1.75 Mg occludes c $\frac{2}{3}$ its volume of H, by heating *in vacuo* the H is suddenly evolved. Obtained in lustrous, white, crystals, hexagonal, $a:c=1.1689$, isomorphous with Zn (Des Cloix, *zeaux*, *C R* 90, 1101). Unchanged in dry air, superficially oxidised in ordinary air. Heated in air, burns to MgO giving out a brilliant white light, very rich in actinic rays. Burns when heated in steam, also in CO_2 , and in SO_2 . Combines directly with the halogens, S, P, As, N. Hot water is slowly decomposed by Mg. Dissolves readily in dilute acids.

Mg is a distinctly metallic element, it forms one series of salts MgX_2 , where $X=Cl, NO_3, \frac{1}{2}SO_4, \frac{1}{2}PO_4$, &c, a few basic salts, and many double salts, are known. In its chemical relations, Mg is analogous to the alkaline earth metals Ca, Sr, and Ba, and also to the metals Be, Zn, Cd, and Hg. MgO and $MgO \cdot H_2O$ are alkaline, $MgO \cdot H_2O$ is formed by the reaction of H_2O and MgO , and is dehydrated to MgO by heat. As the V.D. of no binary compound of Mg with H or one of the halogens has been determined, the valency of the atom of Mg in gaseous molecules is not known with certainty, but from the close similarities between Mg, Zn, and Cd, there is little doubt that the atom of Mg is divalent in gaseous molecules (v MAGNESIUM GROUP OF ELEMENTS, p 163, cf. ALKALINE EARTHS, METALS OF THE, vol 1 p 112).

The atomic weight of Mg has been determined (1) by determining SO_4 in $MgSO_4 \cdot 7H_2O$

(Gay Lussac, *A Ch* [2] 13, 308, Scheerer, *P* 69, 555), (2) by converting MgO into MgSO_4 (Berzelius, *P* 8, 188, Svanberg a Nordenfeldt, *J pr* 45, 474, Jacquelin, *A Ch* [3] 32, 202, Marignac, *A Ch* [6] 1, 289), (3) by converting $\text{MgCO}_3 \cdot 2\text{H}_2\text{O}$ into MgO (S a N 1c), (4) by converting MgCO_3 into MgO (Marchand a Scheerer, *J pr* 50, 385), (5) by determining Cl in MgCl_2 by ppn with Ag (Dumas, *A Ch* [3] 55, 189), (6) by determining S H of Mg

Molecular weight of magnesium—Ramsay (*C J* 55, 521) has determined the lowering of vapour pressure of Hg produced by dissolving Mg in Hg, the results make it probable that the molecular weight of Mg is the same as the atomic weight. This conclusion assumes the accuracy of Van't Hoff's law, that equal volumes of dilute solutions contain equal numbers of molecules of the dissolved substances, and it also supposes the molecular weight of liquid Hg to be the same as the atomic weight of Hg

Reactions—1 Slowly oxidised in moist air—2 Decomposes warm water slowly, heated in steam burns to MgO —3 Heated in air or oxygen to above its M P burns to MgO , with brilliant white light, rich in actinic rays—4 Heated in chlorine, bromine, or iodine forms MgCl_2 , MgBr_2 , or MgI_2 —5 Heated in sulphur forms MgS —6 Heated in nitrogen forms Mg_3N_2 , with phosphorus forms Mg_3P_2 , and with arsenic forms Mg_3As_2 —7 Decomposes carbon monoxide and dioxide, also sulphur dioxide, when heated in these gases, forming MgO —8 Reduces ferric chloride and sulphate solutions to 1eCl, and FeSO_4 (Pfeiffer, *J pr* 61, 441)—9 Reduces nitric acid to HNO_2 (Jenzsch, *N T für Minerale* 1853 535)—10 Reduces carbonates of the alkali metals except Cs, also oxides of copper, silver, and gold, when heated with them in solid form (Winkler, *B* 23, 44), also reduces oxides of Ca, Sr, Ba, Zn, Cd, and Hg (W B 23, 120)—11 Dissolves in dilute acids with formation of salts MgX_2 , X = Cl, $\frac{1}{2}\text{SO}_4$, &c., and evolution of H, or N oxides from HNO_3 . Conc hot H_2SO_4 forms MgSO_4 and SO_2 —12 Dissolves in hydrogen peroxide solution with formation of MgO_2H_2 . From solutions of salts of copper, iron, manganese, and cobalt Mg ppts the metals, with evolution of H, from some metallic salts Mg ppts hydrated oxides, e.g. AlO_2H_2 , from solutions of Al salts, from solutions of salts of arsenic or antimony, Mg evolves AsH_3 or SbH_3 —13 Mg forms alloys with most of the metals (v *Magnesium, alloys of*)

Detection and estimation—Solutions of Mg salts are not pptd by HClAq , $\text{H}_2\text{SO}_4\text{Aq}$, H_2S , or NH_4HS . Fixed alkalis and their carbonates ppt MgO_2H_2 or MgCO_3 , insoluble in excess, but soluble in NH_4 salts, especially in NH_4ClAq , hence Mg salts are not pptd by NH_4Aq in presence of NH_4Cl . $\text{Na}_2\text{HPO}_4\text{Aq}$, added to Mg salt solutions, in presence of NH_4Aq , ppts MgNH_4PO_4 .

Mg is usually estimated by ppn as MgNH_4PO_4 , by addition of NH_4Aq and $\text{Na}_2\text{HPO}_4\text{Aq}$, when strongly heated, the pp is changed to $\text{Mg P}_2\text{O}_7$. For details, and methods of separation of Mg from other metals, *Manuals of Analysis* must be consulted.

Magnesium, alloys of Mg forms alloys with most of the metals. By reason of the easy

oxidation of Mg, the preparation of alloys of this metal is somewhat difficult. They may be prepared by melting the metals in H, also by melting the other metal and rapidly immersing the Mg in it. According to Parkinson (*J pr* 101, 375), Mg forms alloys with the alkali metals, Al, Sb, Bi, Cd, Cu, Au, Pb, Hg, Pt, Ag, Tl, Sn, Zn, also with Cu and Ni together, but not with Co, Fe, or Ni alone. The alloys of Mg are generally very brittle.

Magnesium, aluminate of, MgAl_2O_4 , v ALUMINATES, vol 1 p 141

Magnesium, antimonate of, $\text{Mg}(\text{SbO}_3)_2 \cdot 12\text{H}_2\text{O}$, v ANTIMONATES, vol 1 pp 285, 286

Magnesium, arsenates of, v ARSENATES, vol 1 p 308

Magnesium, arsenide of, Mg_3As_2 . Brown mass. Formed by heating a mixture of As and Mg turnings in H (Parkinson, *J pr* 101, 375, cf ARSENIC, *Combinations* No 9, vol 1 p 303).

Magnesium, arsenites of, v ARSENITES, vol 1 p 306

Magnesium, borates of, v BORATES, vol 1 pp 529, 530

Magnesium, boride of. According to Phipson (*C N* 9, 219), a greenish black mass, which is probably a boride of Mg, is produced by heating Mg with boric acid.

Magnesium, borofluoride of, $\text{Mg}(\text{BF}_4)_2$ (Berzelius, *P* 2, 113), v BOROFLOURIDES, vol 1 p 526

Magnesium, bromide of, MgBr_2 . Mol w unknown, as V D has not been determined. Occurs in small quantity in sea water, and in some mineral springs, also in the Stassfurt salt deposits. Obtained by passing Br vapour over melted Mg. The reaction is very violent (Lerch, *J pr* [2] 28, 338). A white, very deliquescent, fusible, wax like, crystalline mass. By concentrating a solution of MgO in HBrAq , white deliquescent crystals of the hydrated bromide, $\text{MgBr} \cdot 6\text{H}_2\text{O}$, are obtained (Rammelsberg, *P* 55, 239), the same hydrate is formed, along with an insoluble oxybromide, by the reaction between Mg and Br under water (Lerch, 1c). Evaporation of MgBr_2Aq to dryness, or heating the crystals of $\text{MgBr} \cdot 6\text{H}_2\text{O}$, is attended with evolution of HBr and formation of an oxybromide. Heated in a stream of O, MgBr_2 is entirely decomposed to MgO (Schulze, *J pr* [2] 21, 407). Kremers (*P* 108 118) and Gerlach (*Fr* 8, 285) give the following data for S G, and percentage composition, of MgBr_2Aq —

Pct MgBr_2	S G of solution	Pct MgBr_2	S G of solution
5	1 043	30	1 310
10	1 087	35	1 377
15	1 137	40	1 451
20	1 191	45	1 535
25	1 247	50	1 625

The double compounds $\text{MgBr} \cdot \text{KBr} \cdot 6\text{H}_2\text{O}$ and $\text{MgBr}_2 \cdot \text{NH}_4\text{Br} \cdot 6\text{H}_2\text{O}$ are obtained by evaporating mixed solutions of the constituent salts (Lerch, *J pr* [2] 28, 338).

Magnesium, chloride of, MgCl_2 . Mol w unknown, as V D has not been determined.

Occurrence—In sea water and in several mineral springs, also in the salt deposits of Stassfurt.

Formation—1 The mother liquor, after re-

moving KCl from the Stassfurt salts, is evaporated to SG 1375 and allowed to crystallise. The crystals of $\text{MgCl}_2 \cdot 6\text{H}_2\text{O}$ are heated to melting, whereby they are partially dehydrated, but at the same time some oxychloride is formed. By exposing the crystals to a temperature of $c 100^\circ$ under greatly reduced pressure almost pure MgCl_2 is obtained—2 MgCl_2 is obtained as a by-product in Weldon's process for making Cl, and in Solway's soda ammonia process when MgO is used to decompose NH_4Cl —3 By heating MgO with NH_4Cl , adding some NH_4Cl from time to time

Preparation—1 Powdered MgO , or *magnesia alba*, is dissolved in HClAq , NH_4Cl is added in quantity sufficient to prevent pptn of $\text{MgO} \cdot \text{H}_2\text{O}$, on addition of NH_4Aq , the solution is made slightly alkaline by NH_4Aq , any silica which separates is filtered off, the liquid is neutralised by HClAq , and evaporated to dryness. The double salt $\text{NH}_4\text{Cl} \cdot \text{MgCl}_2 \cdot x\text{H}_2\text{O}$ is thus obtained, it is heated to melting ($c 460^\circ$) in a Hessian crucible, in the bottom of which is placed a smaller Pt dish (Bunsen, *A* 82, 137), until NH_4Cl is entirely removed, the MgCl_2 collects in the Pt dish at the bottom of the Hessian crucible (*cf* Doberner, *S* 28, 90)—2 By heating $\text{MgCl}_2 \cdot 6\text{H}_2\text{O}$ in a current of HCl (Hempel, *B* 21, 897)

The hydrated chloride $\text{MgCl}_2 \cdot 6\text{H}_2\text{O}$ is obtained by evaporating the solution of MgO in HClAq to the crystallising point

Properties—A white, deliquescent, solid, consisting of a mass of small pearly crystals, SG 2177 (Playfair & Joule, *C S Mem* 2, 401). Can be distilled in H at red heat. Dissolves readily in water with production of much heat, $[\text{MgCl}_2 \cdot \text{Aq}] = 85,920$ (*Th* 3, 243) Thomsen (*lc*) gives the thermal data $[\text{Mg}, \text{Cl}^\circ] = 151,010$, $[\text{Mg}, \text{Cl}^\circ \text{Aq}] = 186,930$. Sol in alcohol, also in amyl alcohol Gerlach (*J* 1859 43) gives the following table—

P ct. MgCl_2	SG $\text{MgCl}_2 \cdot \text{Aq}$	P ct. MgCl_2	SG $\text{MgCl}_2 \cdot \text{Aq}$
1	1.0084	19	1.1686
2	1.0169	20	1.1780
3	1.0253	21	1.1879
4	1.0338	22	1.1977
5	1.0422	23	1.2076
6	1.0510	24	1.2175
7	1.0597	25	1.2274
8	1.0684	26	1.2373
9	1.0772	27	1.2482
10	1.0859	28	1.2586
11	1.0949	29	1.2690
12	1.1040	30	1.2794
13	1.1130	31	1.2903
14	1.1220	32	1.3012
15	1.1311	33	1.3121
16	1.1404	34	1.3230
17	1.1498	35	1.3340
18	1.1592		

The hydrate $\text{MgCl}_2 \cdot 6\text{H}_2\text{O}$ crystallises in monoclinic forms, SG 1558 (Fihol, *A Ch* [3] 21, 415) *S* 167 cold water, 367 hot water (Cassacca, *C R* 37, 350) *S* 20 in alcohol 19 SG, 50 in alcohol 817 SG

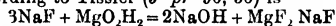
Reactions—1 When the crystals of $\text{MgCl}_2 \cdot 6\text{H}_2\text{O}$ are heated in air, water is given off and at the same time partial decomposition occurs with evolution of HCl and formation of

oxychlorides—2 MgCl_2 is partly decomposed by much water to MgO and HCl

Combinations—1 With *magnesia* to form oxychlorides (*v* *Magnesium, oxychlorides of*, p 162)—2 With *potassium chloride* to form $\text{MgCl}_2 \cdot \text{KCl} \cdot 6\text{H}_2\text{O}$, occurs native as *carminite*, crystallising in rhombic forms according to Marignac (*C R* 155, 650), in hexagonal forms according to Rammelsberg (*Hand der Krystall Chemie*, 204). The double salt is prepared by mixing solutions of the constituents in the proper ratio, evaporating, and crystallising—3 With *ammonium chloride*, and *calcium chloride*, to form $\text{MgCl}_2 \cdot 11\text{H}_2\text{O} \cdot \text{CaCl}_2 \cdot 6\text{H}_2\text{O}$ (*v* Pfaff & Hantz, *A* 66, 250), and $\text{MgCl}_2 \cdot \text{CaCl}_2 \cdot 12\text{H}_2\text{O}$, respectively—4 With *sodium chloride* to form $\text{MgCl}_2 \cdot \text{NaCl} \cdot \text{H}_2\text{O}$ (Poggiale, *C R* 20, 1180)—5 With *ferrie chloride*, and *chromic chloride*, to form $2\text{MgCl}_2 \cdot \text{Fe}_2\text{Cl}_3 \cdot 2\text{H}_2\text{O}$, and $2\text{MgCl}_2 \cdot \text{Cr}_2\text{Cl}_3 \cdot 2\text{H}_2\text{O}$, respectively (Neumann, *A* 244, 328)—6 With *phosphorus oxychloride* to form $\text{MgCl}_2 \cdot \text{POCl}_3$ ($9\text{MgCl}_2 \cdot 10\text{POCl}_3$ according to Cronander, *B* 4, 753), a deliquescent solid, decomposed by hot water (Casselmann, *A* 98, 223) 7 With *iodine trichloride* to form deliquescent, easily decomposed, crystals of $\text{MgCl}_2 \cdot 2\text{ICl}_3 \cdot 5\text{H}_2\text{O}$, obtained by dissolving $\text{Mg}(\text{IO}_3)_2$ in conc HClAq , leading HCl into the liquid to saturation, and surrounding by a freezing mixture (Fihol, *J Ph* 2, 442)—8 With *ammonia* to form $\text{MgCl}_2 \cdot 4\text{NH}_3$, a white sublimate by heating MgCl_2 in NH_3 (Clark, *A* 78, 369)

Magnesium, chromate of, *v* CHROMATES, vol II p 155

Magnesium, fluoride of MgF_2 . Mol w unknown as VD has not been determined. Occurs native as *sellaite*. Formed by dissolving *magnesia alba* in HF Aq , evaporating to dryness, and heating the residue, also by ppg a Mg salt in solution by a soluble fluoride. Obtained in crystals, either by melting the salt, prepared as described, with NaCl or KCl (Cossa, *Z K* [2] 1, 207), or by melting together 5 parts MgCl_2 , 4 parts NaF, and 4 parts NaCl (Röder). Four sided, very hard, white needles SG 2856 at 12° (Cossa, *B* 10, 295) Guntz (*A Ch* [6] 3, 5) gives the thermal data $[\text{MgO} \cdot \text{H}^\circ; 2\text{HF}] = 28,400$. Forms a double compound with NaF, viz $\text{MgF}_2 \cdot \text{NaF}$. Obtained by adding $\text{MgO} \cdot \text{H}$ to NaFAq and evaporating, the decomposition according to Tissier (*J pr* 90, 50) is



Magnesium, haloid compounds of Mg forms a single compound, MgX_2 , with each of the halogens. None of these compounds has been gasified, hence the molecular weight of none is known with certainty, but the great similarities between Mg, Zn, and Cd, make it extremely probable that the formula MgX_2 expresses the composition of the molecules of the compounds in question MgCl_2 , MgBr_2 , and MgI_2 crystallise with $6\text{H}_2\text{O}$, when these crystals are heated they are partly decomposed to MgO and $\text{HX} (\text{X} = \text{Cl}, \text{Br}, \text{or I})$

Magnesium, hydrosulphide of $\text{MgS} \cdot \text{H}_2$. A solution of this compound is obtained by passing H_2S through MgO suspended in water, the MgO dissolves and leaves a nearly colourless solution, which evolves H_2S when exposed to air or heated. The solution thus prepared contains 16 p c $\text{MgS} \cdot \text{H}_2$, and has SG 1.118 at 15°

(Divers & Shimidzu, *C J* 45, 699) $\text{MgS.H}_2\text{Aq}$ is also formed by treating MgS with water. A solution of MgS.H_2 is a convenient source of H_2S , a steady stream of the gas is obtained by heating to $60^\circ\text{--}65^\circ$ (D & S *lc*). Thomsen gives the thermal data $[\text{Mg}, \text{S}^2, \text{H}^2, \text{Aq}] = 114,880$ (*Th* 3, 243).

Magnesium hydroxide of MgO.H_2 . Occurs native as *brucite*, forming white, pearly, crystalline tablets (hexagonal rhombohedra), SG 2 35 to 2 46. Prepared by ppg a Mg salt by NaOHAq , washing, and drying at 100° , also by direct combination of water with MgO , and drying at 100° , SG 2 36 at 15° (Schulten, *C R* 101, 72). MgO which has been heated to whiteness does not combine with water (*v* Deville, *C R* 61, 975, Knapp, *D P J* 202, 513, Schwarz, *D P J* 186, 25, Ditte, *C R* 73, 111, 191, 270) $[\text{Mg}, \text{O}^2, \text{H}^2] = 217,320$, $[\text{Mg}, \text{O}, \text{H}, \text{O}] = 148,960$ (*Th* 3, 243). Soluble c 100,000 parts water, $\text{MgO.H}_2\text{Aq}$ is feebly alkaline to litmus, with acids MgO.H_2 reacts as a strong base, forming salts MgX_2 , $\text{X} = \text{Cl}, \text{NO}_3, \frac{1}{2}\text{SO}_4, \frac{1}{2}\text{PO}_4$, &c., $[\text{MgO}^+\text{H}^2, \text{H}^2\text{SO}_4\text{Aq}] = 31,220$, $[\text{MgO}^+\text{H}^2, 2\text{HNO}_3\text{Aq}] = 27,520$, $[\text{MgO}^+\text{H}^2, 2\text{HClAq}] = 27,690$ (*Th* 3, 243) (*cf* *Magnesium, oxide of*).

Magnesium iodide of MgI_2 . Mol w unknown as VD has not been determined. Obtained by throwing *I*, in small quantities at a time, on to molten Mg (Lerch, *J pr* [2] 28, 338), reaction is very violent. A white, waxy solid, very deliquescent. A hydrate, $\text{MgI}_2.8\text{H}_2\text{O}$ according to Lerch (*lc*), is formed by the reaction between *I* and Hg under water (*l*), or by dissolving *magnesia alba* in HIAq and evaporating over H_2SO_4 , very deliquescent. MgI_2 is decomposed by heating in air, giving off *I*, and forming MgO , heated in *O* it is entirely changed to MgO (Schulze, *J pr* [2] 21, 407). MgI forms double compounds with *KI* and NH_4I (Lerch, *lc*). The following table is given by Kremers (*P* 108, 118) —

Pc MgI_2	SG MgI_2Aq at 19°	Pc MgI_2	SG MgI_2Aq at 19°
5	1 043	35	1 395
10	1 088	40	1 474
15	1 139	45	1 568
20	1 194	50	1 668
25	1 254	55	1 780
30	1 320	60	1 915

Magnesium nitride of Mg_3N_2 . Mol w unknown, as VD has not been determined. Obtained in small transparent crystals by Deville & Caron in making Mg . Briegleb & Geuther (*A* 123, 236) prepared Mg_3N_2 by heating Mg turnings in a stream of dry pure *N*. Mg_3N_2 is also formed, according to Mallet (*C N* 88, 39), when Mg is heated in a small quantity of air. Prepared by heating Mg in *N*, Mg_3N_2 is an amorphous, greenish yellow, powder, becomes brown on heating. Heated in air it is slowly oxidised to MgO , the oxidation is rapid when *O* is substituted for air. With water, forms MgO and NH_3 . With dilute acids, forms Mg salts and NH_3 salts. Heated in HCl , MgCl_2 and NH_4Cl are formed. Heated in dry H_2S , MgS and $(\text{NH}_4)_2\text{S}$ are slowly formed. Heated to redness in CO or CO_2 , MgO , *C*, and C_2N_2 are produced. PCl_5 vapour reacts to form MgCl_2 and *P* nitride.

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Magnesium, oxide of. MgO (*Magnesia*)

Mol w unknown, as compound has not been gasified. SG 3 642 at 12° (Cossa, *B* 10, 1747), SG crystallised 3 636 (Ebelmen, *J* 4, 15), for SG of MgO heated to temperature from 350° to bright redness *v* Ditte, *C R* 73, 111, 191, 270.

A new medicine was introduced in Rome in the beginning of the eighteenth century, the name *magnesia alba* given to it is supposed to have been suggested as a distinction from *magnesia nigra*, a term then applied to *pyrolustite*, MnO . The preparation of the new medicine was kept secret. In 1709 Slevogt showed that *magnesia alba* could be obtained by ppg the mother liquor from the preparation of saltpetre by K_2CO_3 . *Magnesia alba* was supposed to be the same as lime, but Hoffmann found that some of its reactions differed from those of this compound. In 1755 Black demonstrated *magnesia alba* to be a compound of CO_2 with a new earthy base, to which he gave the name *magnesia*.

Occurrence — MgO occurs as *periclase*, crystallised in regular octahedra, SG 3 67, usually coloured green from presence of a little FeO .

Formation — 1 By adding NaOHAq or KOHAc to solution of a Mg salt, washing, and strongly heating the pp of MgO.H_2 . — 2 By calcining *magnesia alba*, a basic Mg carbonate obtained by ppg a Mg salt by an alkaline carbonate (*v* CARBONATES, vol 1 p 699). — 3 By burning Mg in *O*. — 4 By decomposing MgCl_2Aq by calcined dolomite, $\text{MgCl}_2\text{Aq} + \text{CaO.MgO} = 2\text{MgO} + \text{CaCl}_2\text{Aq}$. — 5 In crystals same as *periclase*, by strongly heating powdered MgO in a slow stream of HCl gas (Dewille, *A* 120, 183), also by heating powdered MgO in a porcelain oven (*H* Rose).

Preparation — 1 A boiling solution of pure $\text{MgSO}_4.7\text{H}_2\text{O}$ is pptd by $\text{Na}_2\text{CO}_3\text{Aq}$, after boiling for a little the pp is washed and dried, and the product $(4\text{MgCO}_3.\text{MgO.H}_2\text{O})$ is calcined in a Pt vessel until every trace of CO_2 and H_2O is removed. — 2 *Magnesia alba* is treated with enough dilute HNO_3Aq to dissolve nearly but not quite all, after standing for some time in the air with frequent stirring, to ppt $\text{Fe}_2\text{O}_3.x\text{H}_2\text{O}$, the liquid is filtered, the filtrate is mixed with a little pure MgSO_4 , alcohol is added, and, after digesting for some time with pure CaSO_4 , the liquid is evaporated to crystallisation, and agar filtered, finally the filtrate is evaporated to dryness, the $\text{Mg}(\text{NO}_3)_2$ thus obtained is calcined, the residue is very thoroughly washed, and again calcined (Wuriz). (For preparation of pure MgO from *magnesite v* Caron, *C R* 66, 840).

Properties — A white powder, more or less flocculent according to the temperature and length of time it has been heated. MgO absorbs water and CO_2 from the air, slowly forming $\text{MgCO}_3.x\text{MgO.H}_2.y\text{H}_2\text{O}$. It is slightly soluble in water, 1 part requires 55,368 cold or hot water for solution according to Fresenius, 100,000 to 200,000 according to Bineau. The solution of MgO is feebly alkaline to litmus. When very strongly heated MgO does not melt but gives out a very clear white light. Made into a paste with a little water, MgO sets to a hard, white mass (Dewille, *C R* 61, 975), MgO which has been heated to whiteness does not set with water (*v* Knapp, *D P J* 202, 513, Schwarz, *D P J* 186, 25). MgO reacts with acids as a strong base.

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forming salts MgX_2 , $X = Cl, NO_3, \frac{1}{2}SO_4, \frac{1}{2}PO_4$, &c

Reactions—1 With water to form the hydroxide $MgO.H_2$.—2 Heated with ammonium chloride NH_4Cl is evolved and $MgCl_2$ formed.—3 With acids MgO reacts as a strong base forming salts MgX_2 , $X = Cl, \frac{1}{2}SO_4$, &c.—4 With salts of heavy metals MgO reacts similarly to CaO, BaO , and SrO , ppg hydrated oxides.—5 With carbon dioxide and water, basic carbonates $MgCO_3.xMgO.H_2.yH_2O$ are formed.—6 Suspended in water through which is passed a current of hydrogen sulphide, a solution of $MgS.H_2$ is obtained (v *Magnesium, hydrosulphide of*, p 160).—7 Heated in carbon disulphide vapour, MgO MgS is formed (Reichel, *J pr* [2] 12, 55).

Magnesium, oxychlorides of When freshly heated MgO is moistened with $MgCl_2$ Aq, SG 1.16 to 1.26, the whole sets to a firm, hard mass, which probably consists of a mixture of oxychlorides $xMgO.yMgCl_2$ (v *Sorel, C R* 65, 102). By warming 30 parts MgO with 1500 parts $MgCl_2$ in solution, in an atmosphere free from CO_2 , needle shaped crystals slowly form, when washed free from $MgCl_2$, pressed, and dried over soda lime, the crystals are $10MgO.MgCl_2.18H_2O$, dried at 110° they lose $4H_2O$ (Krause, *A* 165, 38).

Magnesium, oxysulphide of $Mg_2OS (= MgO.MgS)$ An amorphous, reddish, solid, obtained, along with COS , by passing CS_2 vapour over heated MgO (Reichel, *J pr* [2] 12, 55).

Magnesium, phosphide of Mg_3P_2 A hard, brittle, steel grey solid, obtained by heating molten P with Mg in H (Parkinson, *C J* [2] 5, 127, 309), or in a sealed glass tube (Emmerling, *B* 12, 152). Decomposed by water, evolving PH_3 , and forming MgO , with acids gives Mg salts and PH_3 .

Magnesium, salts of. Compounds obtained by replacing H of acids by Mg The salts of Mg belong to one series, MgX_2 , where $X = Cl, NO_3, \frac{1}{2}SO_4, \frac{1}{2}CO_3, \frac{1}{2}PO_4$, &c. A few basic salts, and many double salts, of Mg are known. $MgCO_3$, the basic carbonates and phosphates, and a few other salts are insoluble in water, but most of the Mg salts readily dissolve in water. The salts are usually prepared by dissolving $MgO, MgO.H_2$, or $MgCO_3$, in the various acids, and evaporating, some are prepared by double decomposition from the soluble salts, generally from $MgSO_4$. As none of the salts of Mg has been gasified the formulae given are not necessarily molecular, they are based on the results of analysis and considerations of the similarities between the salts of Mg, Zn, and Cd (v **MAGNESIUM GROUP OF ELEMENTS**). The Mg salts are numerous, and as a class very definite and stable bodies, the following are the chief salts derived from oxyacids—bromate, carbonates, chlorate, dithionate, hypochlorite, hypophosphite, iodate, nitrate and -ite, perchlorate, iodates, phosphates and -ite, selenate and -ite, silicates, sulphate and -ite, tellurate and -ite, thiosulphate (v **CARBONATES, NITRATES, SULPHATES**, &c).

Magnesium, selenide of A flesh coloured pp, probably a selenide of Mg, is obtained by adding K_2Se to solution of a Mg salt, the p.p. is decomposed by heat, and by acids, with separation of Se (Berzelius).

Magnesium, silicides of A compound of Mg and Si is prepared by placing a layer of fused and powdered NaCl in the bottom of a Hessian crucible, adding one half of a mixture of $2\frac{1}{2}$ parts NaCl and 7 parts Na_2SiF_6 , then $2\frac{1}{2}$ parts Mg, and then the other half of the mixture of NaCl and Na_2SiF_6 , covering the whole with NaCl, and heating in a wind furnace, when the reaction is completed the contents of the crucible are stirred with a porcelain rod, and allowed to cool. The regulus formed at the bottom of the crucible is a mixture of Mg and Mg silicide, by washing with NH_4Cl Aq the Mg is dissolved out (Geuther, *J pr* 95, 424). As thus prepared the silicide has the composition Mg_2Si . It forms metal like, lustrous crystals. Reacts with HCl Aq to form $MgCl_2, C_2H_4, H_2SiO_3$, and H. This substance may be a compound, or a mixture, of Mg_2Si and $MgSi$ ($Mg_2Si = 2Mg.Si + MgSi$). A silicide Mg_2Si is obtained by pulverising, in a warm mortar, 40 parts fused $MgCl_2$, 35 parts Na_2SiF_6 , 10 parts NaCl, adding 20 parts Na in small pieces, and projecting the mixture into a hot Hessian crucible. After fusion and cooling, the greyish black mass in the crucible is found to contain metal like tablets and globules, these consist of Si and Mg_2Si , by treating with HCl Aq Si remains and SiH_4 is evolved (Wohler, *A* 107, 113, cf *Martius, A* 107, 112).

Magnesium, silicofluoride of $MgSiF_6$, said to be obtained as a gum like, very soluble substance, by dissolving MgO in H_2SiF_6 Aq (Berzelius).

Magnesium, sulphhydrate of, v *Magnesium, hydrosulphide of*, p 160

Magnesium, sulphides of MgS has been isolated, polysulphides, probably MgS_2, MgS_3 , and MgS_4 , seem to exist in the solution obtained by warming MgS with S and H_2O .

MAGNESIUM MONOSULPHIDE, MgS Produced by passing H laden with S vapour over red hot Mg (Reichel, *J pr* [2] 12, 55, cf *Parkinson, C J* [2] 5, 125, 309). According to Fremy, MgS is formed, along with CO , when MgO is strongly heated in dry H_2S , but Reichel (*l.c.*) says that the products of this reaction are MgO, MgS and COS . MgS is also said to be formed, along with $(NH_4)_2S$, by heating $Mg.N_2$ in dry H_2S (Briegleb & Geuther, *A* 123, 236). Fremy (*A Ch* [5] 38, 324) gives $[Mg.S] = 39,800$. MgS is formed, but only in very small quantity, by strongly heating $MgSO_4$ with C. Alkaline sulphides do not ppt MgS , but $MgO.H_2$, from solutions of Mg salts.

MgS , formed by heating Mg and S, is described as a yellowish grey amorphous, or as a reddish brown crystalline mass, analysis always gives rather more S than required by MgS . With water, MgS is decomposed to $MgO.H_2$ and $MgS.H_2$ Aq, the latter being afterwards converted into $MgO.H_2$ and H_2S .

MAGNESIUM POLYSULPHIDES By warming, not boiling, MgS and S with water, a dark yellow solution is obtained, containing Mg and S in a ratio varying from Mg 3S to Mg 5S (Reichel, *J pr* [2] 12, 55). The solution decomposes in air, giving off H_2S and ppg $MgO.H_2$ and S.

Magnesium, sulphocyanide of, $Mg(SCN)_2.4H_2O$, v vol in p 350

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MAGNESIUM ETHIDE $Mg(C_2H_5)_2$, Magnesium flings act rapidly on ethyl iodide, gas

being given off. If the product be heated in a sealed tube at 125° the contents solidify to a white mass which on distillation gives off MgEt , and leaves a residue of MgI , (Cahours, *A* 114, 227, 354, Hallwachs a Schafarik, *A* 109, 215). These statements are contradicted by Löhr (*A* 161, 48), who finds that EtOAc must be added to the mixture of Mg and EtI , and that the resulting magnesium ethyl is a non volatile solid.

MAGNESIUM GROUP OF ELEMENTS—

Beryllium, Magnesium, Zinc, Cadmium, Mercury Of these five metals, only Zn and Hg are found native, and these in small quantities. Be occurs chiefly as silicate in combination with Al silicate, not widely distributed, Mg salts, especially MgCO_3 and MgSO_4 , are found in large quantities, the chief ores of Zn are ZnCO_3 and ZnS , they occur fairly widely distributed and in considerable quantities, small quantities of CdS and CdCO_3 accompany many Zn ores, the chief ore of Hg is HgS , which is found only in

a few localities. Be is classed with the rare elements, approximately pure Be was not prepared until 1885. Mg was obtained by Davy, in small quantity and impure, in 1800, in 1830 Bussy prepared the metal in a state of approximate purity. Although brass has been made for very many centuries, yet it was not till the middle of the eighteenth century that the art of preparing metallic Zn was known in Europe. Cd was discovered and prepared in 1817 by Stromeyer. Hg has been known certainly from the beginning of the Christian era.

Be and Mg are prepared by reducing the chlorides MCl by Na or K , ZnO and CdO are reduced by heating with C , Hg is prepared from HgS by heating with CaO , whereby CaS and CaSO_4 are formed and Hg is vapourised, or by heating with scrap iron, when FeS , SO_2 , and Hg are produced.

The following table presents some of the prominent physical and chemical properties of the magnesium metals —

	BERYLLIUM	MAGNESIUM	ZINC	CADMIUM	MERCURY
Atomic weights	9.08	24	64.9	111.7	199.8
	One or more compounds of each element, except Mg , have been gasified. Specific heats have been directly determined. Molecular weights of Zn , Cd , and Hg , and probably also of Mg , are the same as the atomic weights.				
Melting points (approx)	—	700°–800°	430°	320°	–39°
Sp gr (approx)	1.85	1.7	7.2	8.7	13.5
Sp heats	62	245	0.94	0.56	0.33
At w	4.92	14.1	9	12.8	14.8
Sp gr (approx)					
Occurrence and preparation	Double silicate of Be and Al , and a compound of BeO and Al_2O_3 , occur in a few rocks in small quantities. Prepared by reduction of molten BeCl_2 by Na , not by electrolysis of BeCl_2 .	MgCO_3 , MgSO_4 , MgCl_2 , and Mg silicates occur in large quantities and widely distributed. Prepared by reducing molten MgCl_2 by Na , also by electrolysis of MgCl_2 .	ZnO , ZnS , and ZnCO_3 are fairly widely distributed, Zn found native in small quantities. Prepared by reducing ZnO by C at moderately high temperature.	Compounds frequently found accompanying Zn ores. Prepared by reducing CdO by C .	Hg found native in small quantities. Chief ore is HgS , found in a few localities in fair quantity. Prepared by heating HgS in air, or with Fe , or CaO , and condensing Hg .
Physical properties	Steel-coloured, hard, hexagonal crystals, lustrous.	Silver-white, very lustrous, moderately hard, malleable, not very tenacious. Crystallises in hexagonal forms, isomorphous with Zn . Melts at temp much higher than M.P. of Cd or Zn , has not been gasified at white heat.	White, with tinge of blue, crystallises in hexagonal forms, soft; brittle at ordinary temperatures, malleable at 100°–150°, brittle again at 200°. Flexibility varies with temp. Melts at c 430°, can readily be distilled.	White, with slight tinge of blue, crystallises easily in regular forms, soft, but harder than Zn , very malleable, ductile, and flexible. Vapour is yellow. Melts at moderate temperature, c 320°, distils at lower temp than Zn .	Liquid metal at temps above –39°, white, showing faint tinge of blue when compared with Ag , very lustrous. Crystallises in regular octahedra. Boils at c 356°. Volatilises even at very low temperatures.

TABLE—cont

	BERYLLIUM	MAGNESIUM	ZINC	CADMIUM	MERCURY
<i>Chemical properties</i>	Scarcely changed by heating in air or O, burns in O H flame to BeO. Scarcely acted on by S at red heat. Burns in Cl to BeCl ₂ . Dissolves slowly in acids to form salts. Only one series of salts known, BeX ₂ , where X = Cl, $\frac{1}{2}$ SO ₄ , $\frac{1}{2}$ PO ₄ , &c. Dissolves slowly in alkali solutions, evolving H. Distinctly positive metal, does not show alloy tropy, BeO has no acidic properties. Atom of Be is divalent in gaseous molecules.	Unchanged in dry air, heated in air, or O, burns to MgO. Decomposes hot water slowly, burns when heated in steam, forming MgO. Decomposes CO ₂ at red heat. Combines directly with halogens S, P, As, and N. Dissolves in acids to form salts. Only one series of salts is known, MgX ₂ , X = Cl, $\frac{1}{2}$ SO ₄ , &c. MgO.H ₂ is a strong base, compounds of Mg do not show any acidic properties. Valency of atom of Mg not certainly known, as no binary haloid compounds have been gasified. Mol w probably same as at w.	Unchanged in air, heated strongly in air, or O, burns to ZnO. Decomposes steam at red heat. Combines directly with halogens, but not with S. Slowly soluble in alkali solution, with evolution of H. Dissolves in acids to form salts. One series of salts known, ZnX ₂ , X = Cl, &c. ZnO.H ₂ dissolves in alkalis, but is ppt again on evaporation. Over H ₂ SO ₄ <i>in vacuo</i> . There are indications of existence of a hydride of Zn. Atom of Zn is divalent in gaseous molecules. Mol w same as at w.	Slowly oxidised on surface in air, heated in air or O, burns to CdO. Cd vapour and steam heated together form H and CdO. Combines directly with halogens, also with S and P. Dissolves in acids to form salts. Only one series of salts known, CdX ₂ , X = Cl, &c. CdO.H ₂ is not soluble in alkalis. No compound of Cd shows any acidic properties. Atom of Cd is divalent in gaseous molecules. Mol w same as at w.	Unchanged in air, heated nearly to B.P. is slowly oxidised to HgO. Does not decompose water or steam. Combines directly with halogens and S. Dissolves in acids to form salts. Two series of salts* are known, HgX and HgX ₂ , X = Cl, &c., salts of both series are numerous and stable. HgO dissolves in molten KOH to form K ₂ O.2HgO, HgS dissolves in Na or K sulphide solution to form compounds xNa ₂ S.yHgS. Hg forms very many ammonio-compounds. Atom of Hg is monovalent and divalent in gaseous molecules. Mol w same as at w.

General formulæ and character of compounds

Oxides —MO, also in case of Hg, Hg₂O.
 Hydroxides —MO.H₂. Sulphides —MS, also in case of Hg, Hg₂S. Haloid compounds —MX₂, also in case of Hg, HgX. Salts —MX₂, also in case of Hg, MX, X = NO₃, ClO₄, $\frac{1}{2}$ CO₃, $\frac{1}{2}$ SO₄, $\frac{1}{2}$ PO₄, &c.

The oxides MO are basic, they react with acids to form salts MX₂, X = NO₃, $\frac{1}{2}$ SO₄, &c. MgO reacts with water to form MgO.H₂, the other hydroxides MO.H₂ are formed by ppg solutions of salts by alkalis. HgO shows feebly marked acidic properties, it dissolves in molten KOH, and the compound 2HgO.K₂O is obtained on cooling. ZnO.H₂ is sol KOH aq, but is reppd on evaporation *in vacuo* over H₂SO₄. Mercurous oxide Hg₂O is the only representative of the class M₂O, it is very easily resolved into Hg and HgO, it reacts with dilute acids in the cold, as a basic oxide, forming salts HgX. Hg₂O in some respects resembles Cu₂O and Ag₂O, and exhibits the relations of Hg to the copper group (v COPPER GROUP OF ELEMENTS, vol II p 250).

The hydroxides MO.H₂ are all decomposed

by heat to MO and H₂O, HgO.H₂ if it exists is extremely easily separated into its oxide and H₂O, the only hydroxide obtained by direct reaction between MO and H₂O is MgO.H₂. The hydroxides are basic, MgO.H₂ is slightly alkaline.

The sulphides MS are basic. The existence of BeS is extremely doubtful, MgS is readily decomposed by water, giving MgS.H₂Aq, and finally MgO.H₂ and H₂S. ZnS, CdS, and HgS are more stable than MgS, HgS shows slightly acidic properties, it dissolves in K₂SAq or Na₂SAq to form compounds xK₂S.yHgS. Mercurous sulphide Hg₂S is very easily decomposed to HgS and Hg.

The haloid compounds MX₂ may be obtained by the direct union of the elements, the mercurous compounds HgX are also obtained in this way. MX₂ and HgX are also prepared by dissolving the corresponding oxides in HXAq and evaporating, but MgCl₂ cannot be thus obtained, as evaporation towards dryness of MgCl₂Aq results in formation of HCl and MgO mixed with oxychlorides xMgO.yMgCl₂. The compounds

BeCl_2 , BeBr_2 , ZnCl_2 , CdBr_2 , HgCl_2 , HgI_2 , and probably HgCl , have been vapourised without decomposition. The haloid compounds form many double compounds. They all combine with NH_3 , various ammonio derivatives of HgX_2 , and HgX are known. Numerous oxyhaloid compounds, $\text{xMO} \cdot \gamma\text{MX}_2$, are known.

The salts MX_2 are numerous and well marked, they are obtained by dissolving the corresponding oxides or carbonates in acids, and evaporating. Mercurous salts HgX are prepared by digesting excess of Hg with the various acids in the cold. Many basic salts of Be , Zn , and Hg have been prepared, most of the salts of Mg are normal.

The five metals Be , Mg , Zn , Cd and Hg belong to Group II, as the elements are arranged in accordance with the periodic law. The following table shows their positions in the Group —

Even Series				
2	4	6	8	
$\text{Be} = 9$	$\text{Ca} = 40$	$\text{Sr} = 87$	$\text{Ba} = 137$	
Odd Series				
3	5	7	9	11
$\text{Mg} = 24$	$\text{Zn} = 65$	$\text{Cd} = 112$	—	$\text{Hg} = 200$

The properties of Be , the first member of the group, suggest those both of the even and odd series members. Physically it approaches Mg , Zn , and Cd more than Ca , Sr , and Ba , in the conditions under which it is oxidised, the basic but non alkaline character of its hydroxide, the existence of many basic salts, &c. Be shows closer resemblances to Zn , Cd , and Hg than to Ca , Sr , and Ba . The solubility of $\text{BeO} \cdot \text{H}_2$ in KOH aq. also suggests the solubility of $\text{ZnO} \cdot \text{H}_2$ in KOH aq., and of HgO in molten KOH , the hydroxides of Ca , Sr , Ba , and Mg are quite insoluble in alkalis. Mg is closely related in many respects to the metals of the alkaline earths, Ca , Sr , and Ba , but its much greater stability in air, its non decomposition of cold water, the easy resolution of its hydroxide into MgO and H_2O by heat, &c., exhibit the analogies between Mg and Zn , Cd , Hg . Sulphides of Ca , Sr , and Ba are readily formed, but MgS is only produced by heating Mg to redness in H_2 laden with S vapour, the difficulty of forming MgS recalls the non-isolation of any compound of Be and S . Alkali carbonate solutions ppt MCO , from solutions of salts of Ca , Sr , and Ba , but the pps obtained from Mg salts are basic carbonates $\text{xMgCO}_3 \cdot \gamma\text{MgO} \cdot \text{H}_2\text{O}$. The sulphates of Ca , Sr , and Ba are reduced to MS by heating with C , but MgSO_4 is scarcely, if at all, reduced in this way. Hg is distinguished from the other members of the magnesium family by its physical properties, the formation of two series of salts, the formation of a large number of ammonio-derivatives of both series of Hg salts, and in other ways.

In the composition of their salts, and sometimes in the isomorphism of salts, the metals of the Mg family resemble those of the Fe family, viz Mn , Fe , Ni , Co (*v* IRON GROUP OF ELEMENTS, p 65). For a further treatment of the elements of Group II *v* CLASSIFICATION, vol II pp 204-207, *cf* also ALKALINE EARTHS, METALS OF THE, vol I p 112.

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MAGNETIC ROTATORY POWERS of liquid compounds, *v* PHYSICAL METHODS, section Optical methods.

MAIROGALLOL $\text{C}_{18}\text{H}_{18}\text{Cl}_2\text{O}_{10}$ (?) [190°] (*From malpw*, I glitter). Obtained, together with leucogallol, by passing chlorine through pyrogallol (15 g) dissolved in 60 p.c. acetic acid (45 g) until the liquid is brown, and allowing the liquid to stand (Groves, *A* 179, 237, Stenhouse & Groves, *C J* 28, 706, Webster, *C J* 45, 208). Tri-metric prisms, insol water, *v* sol hot alcohol, extremely sol ether, insol CS_2 , and petroleum ether. Decomposed by boiling with water yielding CO , oxalic acid, and tri-chloro pyrogallol. Reduced by zinc and H_2SO_4 to tri-chloro pyrogallol (Webster, *C J Proc* 3, 130, Hantzsch & Schniter, *B* 20, 2033). Boiling with dilute H_2SO_4 gives a condensation product, and also a substance that is pptd from aqueous solution by gelatin. Aqueous NaHSO_3 and zinc dust reduce it to tri-chloro pyrogallol and an amorphous substance which yields furfuraldehyde when distilled with dilute H_2SO_4 . Mairogallol gives a crimson colour with solution of sodium sulphite (Cross & Bevan, *C J* 43, 18). Mairogallol when gradually heated gives a sublimate of tri-chloro quinone and tetra-chloro quinone. These reactions indicate that mairogallol is derived by condensation from $\text{CO} < \begin{smallmatrix} \text{C}(\text{OH})_2 & \text{C}(\text{OH})_2 \\ \text{CCl} & - & \text{CCl} \end{smallmatrix} > \text{CCl}_2$ (Webster).

MALACHITE GREEN *v* TETRA METHYL-DIAMIDO TRI PHENYL-CARBINOL.

MALAMIC ACID *v* MALEIC ACID.

MALEIC ACID $\text{C}_4\text{H}_2\text{O}_4$, *vc*

$\text{CO}_2\text{H} \cdot \text{CH} \cdot \text{CH} \cdot \text{CO}_2\text{H}$. Mol w 116 (confirmed by Raoult's method, Paternò, *B* 21, 2158) [130°] R_D 38.6 (in a 6.9 p.c. aqueous solution, Kanonnikoff, *J pr* [2] 31, 347, *cf* Knop, *A* 248, 175) H C v 331,702 (Lougounine, *C R* 106, 1290) H C v 326,900 (Stohmann, Kleber & Langbein, *J pr* [2] 40, 217) H C p 326,300 (S K & A) H F 187,700 (S K & A) Heat of solution -4,438 (Gal & Werner, *Bl* [2] 47, 158) Heat of neutralisation 26,648 (G & W).

Formation—1 Together with fumaric acid (*q v*) by the distillation of maleic acid (Lassaigne, *A Ch.* [2] 11, 93, Pelouze, *A Ch.* [2] 56, 72, Liebig, *A* 11, 276, Buchner, *A* 49, 57, Kekulé, *A Suppl* 1, 129, 2, 85, Von Richter, *Z* [2] 5, 453)—2 By heating silver succinate at 180° (Bourgoin, *C R* 72, 52)—3 By saponification of the ether obtained from di-chloro acetic ether by heating with reduced silver at 220°, or by the action of sodium upon its ethereal solution in the cold (Tanatar, *B* 12, 1563)—4 By the action of KCN on α -di-bromo propionic acid (Tanatar, *B* 13, 159)—5 From δ -bromo pyromucic acid and dilute HNO_3 (Hill & Sanger, *A* 232, 55)—6 By the action of baryta on tri-chloro phenomalic acid ($\begin{smallmatrix} \text{CH} & \text{C}(\text{OH}) & \text{CCl}_2 \\ || & & \\ \text{CH} & \text{CO} & \end{smallmatrix}$).

Preparation—Maleic acid is rapidly distilled until the residue thickens, the flame being then withdrawn, the distillation then continues by itself until the residue is solid (fumaric acid). The distillate consists of maleic acid and maleic anhydride, and may be wholly converted into maleic anhydride by AcCl . The maleic anhydride may be purified by crystallisation from

HOAc and then converted by hot water into maleic acid (Anschutz, *B* 12, 2281, cf Perkin, *B* 14, 2547)

Properties—Monoclinic prisms ∇ sol water and alcohol, m sol ether. Its aqueous solution reddens litmus strongly. Begins to boil at 160°, being then restored into maleic anhydride and water, which partially re unite in the receiver. An aqueous solution of maleic acid is ppd by baryta (difference from fumaric acid). Unlike fumaric acid, maleic acid is not ppd by adding mineral acids to solutions of its salts.

Reactions—1 Reduced by water and sodium amalgam to succinic acid. Zinc added to an aqueous solution of maleic acid yields zinc maleate and succinate. 2 Calcium maleate fermented in contact with cheese also changes to succinate (Dessaignes). 3 *Electrolysis* gives acetylene, succinic acid, and a small quantity of fumaric acid (Kekulé, *A* 180, 1). 4 Combines readily with bromine in the cold, forming iso dibromo succinic acid and some di bromo succinic acid derived from fumaric acid into which a portion of the maleic acid is at the same time transformed (Kekulé, Petri, *A* 195, 59). *Chlorine* gives a di chloro succinic acid (Petrifff, *Bl* [2] 41, 309). Bromine in presence of water forms iso di-bromo-succinic, bromo fumaric, bromo succinic, and fumaric acids (Wilschusen, *A* 246, 58). 5 Dissolves easily in aqueous HBr, the solution even at 0° soon depositing fumaric acid and bromo succinic acid in equimolecular proportions. The bromo succinic acid may easily be extracted from the product by water (Fittig & Dorn, *B* 9, 1191). 6 Converted into fumaric acid by boiling conc HIAq, HBrAq, dilute HNO₃, or HClAq (Kekulé, *A* Suppl 1, 184, 2, 93, Kekulé a Strecker, *A* 223, 186). When HIAq is used the ultimate product is succinic acid. Dry HCl does not convert maleic acid into fumaric. Maleic acid is converted into fumaric acid by heating its aqueous solution at 100°–180° (Semenoff, *Bl* [2] 46, 816). An aqueous solution of sodium maleate is not affected by being heated for 10 hours in a sealed tube at 100° to 185°. 7 With hypochlorous acid it forms a compound which reduced by sodium amalgam yields inactive malic acid, oxymaleic acid (?), and a liquid which breaks up on heating into CO₂ and C₆H₁₀O₆ (Petrifff). 8 KMnO₄ oxidises it to inactive tartaric acid. 9 Alcoholic NH₃ at 150° slowly forms aspartic acid (Engel, *C R* 104, 1805). 10 The acids of the *maleic series* readily split off H₂O from their acid aniline salts, when the aqueous solution of the latter is left to stand for several days or boiled, a crystalline pp being formed of the acid anilide $X'' < \begin{smallmatrix} \text{CONHPh} \\ \text{CO}_2\text{H} \end{smallmatrix}$ or of

an isomeride thereof. Under the same conditions the acid aniline salts of the acids of the *fumaric series* are quite stable. This constitutes a ready method of determining to which of the two isomeric series a given acid belongs. The acid is half neutralised with aniline and either left to stand or heated to boiling, if a pp is obtained insoluble in dilute HCl the acid belongs to the *maleic series* (Michael, *B* 19, 1872).

Salts—K₂A'' (dried at 100°). Radiating crystals, ∇ sol water, insol alcohol. —KHA'' $\frac{1}{2}$ aq small crystals, ∇ sol water. The solution reddens litmus. —Na₂A'' needles. Ppd as a

granular crystalline powder by adding alcohol to its aqueous solution. Is not deliquescent. —NaHA'' $\frac{3}{4}$ aq Trichinic crystals (Bodewig, *J* 1881, 716). Sl sol cold water. —KNaA'' aq deliquescent crystalline powder (Büchner, *A* 49, 60). —BaA'' $\frac{2}{3}$ aq gelatinous pp, changing to scales *S* 11 at 20° (Regnault). —BaA'' aq (Anschutz) —BaH₂A'' $\frac{5}{6}$ aq indistinct crystals, which redden litmus, ∇ sol water (Büchner). —CaA'' aq (dried at 100°) small needles, sol water. —CaH₂A'' $\frac{5}{6}$ aq long prisms, ∇ sol water, insol alcohol. —SrA'' $\frac{5}{6}$ aq silky needles. —SrH₂A'' $\frac{8}{9}$ aq rect angular prisms. —MgA'' $\frac{3}{4}$ aq —MgH₂A'' $\frac{6}{7}$ aq hard crystals, ∇ sol water, insol alcohol. —ZnA'' $\frac{2}{3}$ aq crystals, ∇ sol water, insol alcohol. —NiA'' aq apple green crystals. —CuA'' light blue crystals, sl sol boiling water, ∇ sol NH₃Aq, forming a solution whence alcohol ppts an azure blue crystalline powder. —PbA'' $\frac{3}{4}$ aq amorphous pp, changing to needles. —Pb₂A''O (dried at 100°, Otto, *A* 127, 178). —AgA'' white pp, changing to lustrous needles. —AgHA'' (dried at 100°) needles.

Acid aniline salt C₆H₅NH₃HA'' $\frac{1}{2}$ aq prisms, sol water. On allowing the solution to stand for some days it forms phenyl aspartic acid C₆H₅NH C₆H₄(CO₂H)₂ [132°] (Anschutz a Wirtz, *A* 239, 140, cf Michael, *B* 19, 1373, *Am* 9, 333, 197). The neutral aniline maleate when boiled in aqueous solution deposits the phenylimide of phenyl aspartic acid C₆H₅NH C₆H₄ < $\begin{smallmatrix} \text{CO} \\ \text{CO} \end{smallmatrix}$ > NC₆H₅, a body [212°] which was formerly described as the di anilide of maleic acid C₆H₄(CONHPh)₂.

Phenyl-imide C₆H₅ < $\begin{smallmatrix} \text{CO} \\ \text{CO} \end{smallmatrix}$ > NC₆H₅ [91°] (162° at 12 mm). Formed by the dry distillation of acid aniline maleate (Anschutz a Wirtz, *Am* 9, 238, cf Michael a Wing, *Am* 7, 280). Long yellow needles, ∇ sol benzene, CHCl₃, ether, and alcohol, sl sol CS₂, and ligrom. Its solution in chloroform takes up bromine, forming the phenyl imide of di bromo succinic acid [159°]. Baryta water at 40° gives the acid anilide of fumaric acid.

Mono methyl ether HMeA'' Probably exists in the solution of maleic anhydride in MeOH. Such a solution becomes hot when mixed with NaOMe, probably forming NaMeA'', but the solution presently deposits a bulky white, partly crystalline, pp, which forms an alkaline aqueous solution, becoming neutral on boiling, the neutral solution contains the methyl derivative of sodium methyl maleate

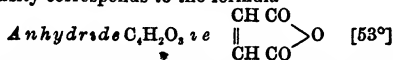
C₆H₅(OMe)(CO₂Na)(CO₂Me) (Purdie, *C J* 47, 873)

Di methyl ether Me₂A'' (205° i ∇) *S G* 14 1158 H C 669,570 (Ossipoff, *C R* 109, 312). From silver maleate and MeI, excess of iodine being avoided, as this would convert it into di methyl fumarate (Anschutz, *B* 12, 2282). Bromine effects the same change.

Mono-ethyl ether HEtA'' From the anhydride and boiling alcohol. Also from HAgA' and EtI (Ossipoff, *J R* 20, 254). Not identical with the mono ethyl ether of fumaric acid. Forms a salt NaEtA'' By treating HEtA'' with NaOEt and boiling the product with water C₆H₅(OEt)(CO₂Et)(CO₂Na) is obtained, identical with that derived from fumaric acid (Purdie)

Di-ethyl ether Et₂A" (225° i V) From Ag₂A" and EtI (Anschütz, *B* 11, 1644, 12, 2281) Vapour density corresponds to the formula (O) Colourless liquid Converted by iodine into the fumaric ether Bromine gives di bromo succinic ether Boiling with NaOMe forms a product whence boiling water produces C₂H₃(OMe)(CO₂Et), (Purdie, *C J* 47, 868)

Di-isopropyl ether Pr₂A" Vapour density corresponds to the formula



(Anschütz), [57°] (Pelouze, *A Ch* [2] 56, 72), [60°] (Fittig, *A* 188, 87) (202° i V) (A) V D 48 (H=1) (calc 49) (Hubner & Schreiber, *Z* [2] 7, 712) H C 336,920 (Ossipoff, *C R* 109, 311) Appears to be the sole anhydride of both maleic acid and fumaric acids

Formation—1 By rapidly distilling maleic or fumaric acid, and frequently rectifying the product (Pelouze, *A* 11, 263, Kekulé, *A Suppl* 2, 87) *In vacuo*, maleic acid splits up into water and anhydride even at 100° (Reicher, *R T C* 2, 312) —2 By heating fumaric acid with AcCl and HOAc at 100° —3 From silver fumarate and fumaryl chloride (Perkin, *B* 14, 2645) —4 By distilling chloro or bromo succinic anhydride (Anschütz & Bennert, *B* 15, 643) —5 By distilling maleic anhydride

Preparation—1 Maleic acid is treated with an excess of acetyl chloride, and the product (which probably chiefly consists of acetyl maleic anhydride) is distilled, when it splits up into acetic acid and maleic anhydride, the yield is 45 p c (Perkin, *C J* 39, 562, *B* 14, 2547) —2 By dry distillation of maleic acid and treatment of the residue and distillate with AcCl, the yield is 56 p c of the theoretical (Anschütz, *B* 12, 2281)

Properties—Trimetric crystals, *abc* = 6408 1 4807 (Bodewig, *B* 14, 2788) In contact with water it is converted into maleic acid, which is conveniently prepared in this way The abnormal rapidity of the etherification of maleic acid by heating with alcohols appears to be due to an intermediate formation of anhydride (Reicher, *R T C* 2, 308) Maleic anhydride heated in a sealed tube at 260° gives a carbonaceous residue and gases (Semenoff, *Bl* [2] 46, 816)

Reactions—1 Unites with bromine at 100°, forming iso-di bromo succinic anhydride, which is resolved at 180° into HBr and bromo maleic anhydride —2 PCl₅ gives the chloride of fumaric acid (Perkin, *B* 14, 2648), but maleyl chloride (71° at 11 mm) appears to be first formed (Anschütz & Wirtz) —3 With a saturated solution of HCl in HOAc it forms at 100° chloro-succinic acid —4 When heated with phenylhydrazine (1 mol) at 150° it forms the phenyl hydrazide C₆H₅·C₂O₂·N₂·HPh, which crystallises in needles [259°], and turns brown at 180° (Hötte, *J pr* [2] 35, 295) —5 When heated with phenols it produces fluorescent compounds analogous to the phthaleins, which may therefore be called 'maleins' The resorcin malein has a green fluorescence, (a) naphthol malein has a greenish-red fluorescence (Burckhardt, *B* 18, 2864)

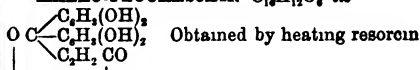
Constitution—The constitution of fumaric

and maleic acids is discussed under FUMARIC ACID, vol ii p 585 The anomalous formula CO₂H C CH₂ CO₂H for maleic acid, based on the formula CO₂H CBr CH₂ CO₂H, is rendered improbable by the observation that bromo-fumaric and not pyruvic acid is the product formed, together with CO₂, when barium iso-di bromo succinate is treated with moist Ag₂O in the dark (Demuth & V Meyer, *B* 21, 264) Racemic acid is formed when silver iso-di bromo succinate is boiled with water The representation of maleic and fumaric acids by formulae in space has been discussed by Wislizenus in his treatise *Ueber die räumliche Anordnung der Atome in organischen Moleculen*

References—AMIDO, BROMO, and CHLORO MALEIC ACIDS

Allo maleic acid v FUMARIC ACID

MALEO-FLUORESCÉIN C₁₂H₁₂O₄, s.s



with maleic anhydride (Lunge & Burckhardt, *B* 17, 1598) Small needles Sol alcohol, sl sol water Dissolves in alkalis to a red solution, with a strong green fluorescence With KOH, MeI, and MeOH it gives a dimethyl derivative, C₁₂H₁₀Me₂O₄, crystallising in red needles AcCl in HOAc gives C₁₂H₁₀Ac₂O₄, crystallising from HOAc in yellowish needles, insol water, CHCl₃, and benzene, sl sol alcohol (Burckhardt, *B* 18, 2864)

MALEYL CHLORIDE v Chloride of FUMARIC ACID

MALIC ACID C₄H₄O₄, s.s

CO₂H CH₂ CH(OH) CO₂H Mol w 134 [100°] S G 2 156 (Schröder, *B* 12, 1611) S G of solutions (Schneider, *A* 207, 262) *Heat of solution*—3148 *Heat of neutralisation* 24,919 (Gal & Werner, *Bl* [2] 46, 803) [α]_D = -3 in dilute solutions, as the liquid becomes more concentrated it approaches +5.9

Occurrence—Maleic acid was discovered by Scheele in 1785, but its composition was first correctly determined by Liebig (*A* 26, 166) It is very widely diffused in the vegetable kingdom, being contained in plants sometimes in the free state, sometimes in the form of potassium, calcium, or magnesium salt It is found in abundance, together with citric acid, in unripe apples, in the fruits of the barberry, sloe, elder, and mountain ash, and in gooseberries, cherries, bilberries, strawberries, raspberries, and many other fruits It is likewise found in the roots of marsh mallow, angelica, aristolochia, bryony, liquorice, primrose, and madder, in carrots and potatoes, in the leaves and stems of aconite, belladonna, hemp, celandine, holy thistle, lettuce, tobacco, poppy, rue, sage, house-leek, tansy, thyme, valerian, and melilot, in the flowers of chamomile, elder, and mullein, in pine apples and grapes, in the seeds of caraway, cummin, parsley, anise, flax, and pepper, in asafetida, opoponax, and myrrh It also occurs in the leaves of the common ash (Gintl, *Z* [2] 5, 377), and in the fruit of the sumach (Reinsch, *Z* 1866, 221) Yolk, the sweat of sheep, contains potassium malate to the extent of 2.5 p c. of the solid residue (Buisine, *C R* 106, 1426)

Formation—1. Maleic acid, with the same

optical properties as those with which it exists in plants, may be formed from asparagine or active aspartic acid by treatment with nitrous acid (Piria, *A Ch* [3] 22, 160)—2 Malic acid, with a rotatory power equal and opposite to that contained in plants, may be obtained from ordinary tartaric acid by the action of phosphorus, iodine, and water (Dessaignes, *A* 117, 134, Bremer, *Bl* [2] 25, 6, *B* 8, 861, 1594)

Preparation—1 The juice of mountain-ash berries, not quite ripe, after being pressed, boiled up, and filtered, is partly neutralised with carbonate of potassium, leaving, however, a sufficient excess of acid to redden litmus pretty strongly, then precipitated by nitrate of lead, set aside for a few days, till the curdy precipitate is completely converted into small needles, these crystals freed from the admixed mucous or flocculent compound of lead oxide and colouring matter by careful elutriation with cold water, are, lastly, well washed with water. The needles of impure malate of lead are boiled with a slight excess of dilute sulphuric acid, the filtrate divided into two equal portions, one portion exactly neutralised with ammonia, the other portion then added, and the reddish liquid evaporated and cooled, it then yields nearly colourless crystals of acid malate of ammonia, which may be rendered quite colourless by recrystallisation. These crystals are then precipitated by acetate of lead, and the precipitate, after thorough washing, is decomposed by sulphydric or sulphuric acid (Liebig)—2 The juice is boiled for some time with milk of lime in quantity not quite sufficient for neutralisation, and the pp poured into a boiling mixture of HNO_3 (1 pt) and water (10 pts) The acid calcium malate which crystallises out on cooling is recrystallised from water, dissolved in water, ppd by lead acetate, and the lead malate decomposed by H_2S (Liebig, *A* 38, 259) By similar processes malic acid may be obtained from house-leek, from cherries or barberries, from the berries of *Rhus coriaria*, from the stems of rhubarb, from apples, and from tobacco

Properties—The aqueous solution, concentrated to a syrup and left to evaporate in a warm place, yields groups of colourless shining needles or prisms of 4 or 6 faces. They melt at 83° (Pelouze) or 100° (Pasteur), and do not suffer any loss of weight at 120° . They deliquesce in the air, and are very soluble in water and alcohol. Dilute solutions of malic acid are laevorotatory, but the rotation changes so rapidly with concentration that concentrated solutions are dextrorotatory. These dextrorotatory solutions are rendered laevorotatory by the addition of H_2SO_4 , or of HOAc $[\alpha]_D = 5.891 - 0.895q$ (where q = percentage of water in solution) (Schneider, *A* 207, 263). Solutions of sodium malate containing about 54 pts of the salt to 46 pts of water are inactive at 20° , stronger solutions are laevorotatory, while more dilute solutions are dextrorotatory (Thomsen, *J pr* [2] 35, 153). Malic acid kills algae (*Zygnemaceae*) (Loew & Bokorny, *J pr* [2] 36, 272). Malic acid prevents the ppn. of cupric and ferric salts by alkalis, 2 mols holding 1 mol of CuO in solution (Hofmeister, *A* 189, 27, cf Juette, *Fr* 7, 489). A dilute solution of malic acid or of a malate is not immediately ppd. by lime water or by CaCl_2 either in

the cold or on heating, but on the addition of alcohol a white pp of calcium malate separates (Braconnot, *A Ch* [2] 51, 331, H Rose, *P* 31, 210). However, a solution of malic acid (1 mol) mixed with pure milk of lime (2 mols) may solidify to a pasty mass (Iwig & Hecht, *A* 238, 171). A concentrated solution of an alkaline malate is ppd by CaCl_2 , but presence of NH_4Cl hinders the ppn, which, however, takes place on adding alcohol. Lead acetate gives a white pp, soluble in excess of malic acid and in ammonia, when the pp is heated in the mother liquid it melts to a semifluid translucent mass. Malates are not blackened by heating with fuming H_2SO_4 . Dilute HOAc at 60° dissolves lead malate, but not lead tartrate or citrate (Hartsen, *Fr* 14, 373, *Ar Ph* [3] 6, 11v). Malic acid may be separated from oxalic and tartaric acids by ppg the latter with CaCl_2 , filtering, and ppg calcium malate by adding alcohol to the filtrate (Barfoed, *Fr* 7, 403). Ammonium malate may be separated from ammonium citrate, tartrate, and oxalate by solution in alcohol (Barfoed)

Reactions—1 Heated for some hours at 140° it yields water and fumaric acid. At 180° it gives off water and maleic anhydride, while fumaric acid remains as a solid residue. If the malic acid be suddenly heated to 200° and kept at that temperature a comparatively large quantity of maleic anhydride is obtained. The maleic anhydride, combining with the water in the distillate, is partially converted into maleic acid. Malic acid is carbonised when suddenly exposed to a red heat—2 By treatment with potash and bromine, bromoform is obtained (Cahours, *A Ch* [3] 19, 507)—3 Nitric acid easily oxidises it to oxalic acid and CO_2 —4 Conc HIAq at 180° reduces it to succinic acid (Schmitt, *A* 114, 106). Succinic acid is also formed when calcium malate is fermented in contact with yeast (Piria, *A* 70, 102, Liebig, *A* 70, 104, 363). Fermentation of calcium malate by certain *Schizomycetes* produces succinic, acetic, butyric, and propionic acids and ethyl alcohol (Fitz, *B* 11, 1896, 12, 481, cf Dessaignes, *C R* 28, 16, Liebig, *A* 70, 104, 363, Kohl, *A* 78, 252, Baer, *Ar Ph* [2] 69, 147, Winkler, *Jahrb pr Pharm* 22, 300, Rebling, *Ar Ph* [2] 67, 300, Béchamp, *C R* 70, 999)—5 By slow oxidation in the cold with $\text{K}_2\text{Cr}_2\text{O}_7$ it is converted into malonic acid (Dessaignes, *A* 107, 251)—6 When boiled in dilute aqueous solution with MnO_2 it yields a distillate containing aldehyde (Liebig, *A* 113, 14)—7 Heated with KOH it yields acetic and oxalic acid—8 When calcium malate (1 pt) is heated with PCl_5 (4 pts) the chloride of fumaric acid passes over (Perkin & Duppa, *A* 112, 24, Liébs Bodart, *A* 100, 327)—9 Boiling aqueous H_2SO_4 at 135° gives aldehyde, CO_2 , water, and CO (Weith, *B* 10, 1744)—10 The electrolysis of potassium malate gives aldehyde, CO_2 , and some acetic acid (Bourgois, *Bl* [2] 9, 427)—11 Water containing a few drops of H_2SO_4 at 160° gives fumaric acid (Markownikoff, *A* 182, 351)—12 When heated with phenol and H_2SO_4 it gives

CO and C_2H_4 $\begin{array}{c} \text{O} - \text{CO} \\ \diagup \quad \diagdown \\ \text{CH} \quad \text{CH} \end{array}$ (Pechmann, *B* 17, 929, 1649)—13 Long boiling with conc HClAq gives fumaric acid—14 Conc HBr at 100° gives bromo-succinic acid and fumaric acid (Kekulé,

A. 180, 11) —15 Chloral at 125° forms
 $\text{CCl}_3\text{CH} \begin{array}{l} \diagup \text{O CH CH}_2\text{CO}_2\text{H} \\ \diagdown \text{O CO} \end{array}$ [140°] This forms

large crystals, sl sol cold water (Wallach, A 193, 42) With PCl_5 it gives the oily chloride $\text{C}_4\text{H}_5\text{Cl}_5\text{O}_2$, whence alcohol forms $\text{C}_4\text{H}_5\text{Cl}_4(\text{OEt})\text{O}_2$ [46°], and MeOH gives $\text{C}_4\text{H}_5\text{Cl}_4(\text{OMe})\text{O}_2$ [85°] — 16 Phenyl hydrazine at 120° forms $\text{C}_{16}\text{H}_{18}\text{N}_4\text{O}_8$, & $\text{C}_2\text{H}_5\text{O}(\text{CO N}_2\text{H}_5\text{Ph})_2$ [223°], which crystallises from dilute alcohol in plates (Bulow, A 236, 195, cf Fischer a Passmore, B 22, 2734) — 17 Malic acid (3 pis) heated with *m* amido-benzoic acid at 150° forms a product whence by washing with hot water and alcohol, dissolving in aqueous NH_3 and ppg by HCl , there is obtained a white powder $\text{C}_8\text{H}_8\text{O}(\text{CO NH C}_6\text{H}_4\text{CO}_2\text{H})_2$. The ammonium salt of this acid forms with cupric acetate a pp of $\text{Cu}(\text{C}_8\text{H}_8\text{N}_2\text{O}_8)_2$. Ac_2O decomposes it, forming NH_4Ac , $\text{C}_6\text{H}_4\text{CO}_2\text{H}$ (Schiff, G 16, 28, A 232, 166)

Salts—Malic acid has a tendency to form acid salts. At 200° the salts give off water and are converted into fumarates— $\text{NH}_4\text{HA}''$ transparent trimetric prisms. Not hemihedral, but becomes so after fusion and recrystallisation SG 125 155 μ S 32 at 157° [α] = -6° On dry distillation it forms fumarimide, fumaric, and maleic acids, and active and inactive malic acid. Combines with acid ammonium tartrate. Normal ammonium malate $(\text{NH}_4)_2\text{A}''$ is v e sol water, and its solution gives off NH_3 to the air, becoming acid. In a solution containing 63 p.c. of the salt [α]_D = -4.2 In a 17 p.c. solution [α]_D = -7.2 (Schneider, A 207, 276) — KHA'' crystals. Begins to decompose at 100° Sol water, insol alcohol [α]_D = -5 in a 21 p.c. solution, = -5.7 in a 9.4 p.c. solution at 20° — KA'' Uncrystallisable and deliquescent mass [α]_D = -2.2 in a 62 p.c. solution, = -6.8 in a 9.4 p.c. solution at 20° (Schneider) — NaA'' Uncrystallisable [α]_D = +4.7 in a 65.5 p.c. solution, = -8.4 in a 5.3 p.c. solution — LiHA'' Uncrystallisable [α]_D = -4.6 in a 50 p.c. solution, = -8.4 in a 10 p.c. solution — $\text{Li}_2\text{A}''$ Uncrystallisable [α]_D = -4.1 in a 39 p.c. solution, = -12 in a 6 p.c. solution at 20° (Schneider, A 207, 273) — BaA'' aq thin plates [α]_D = +8.2 in a 9.4 p.c. solution, = -2.6 in a 2 p.c. solution at 20° — $\text{BaH}_2\text{A}''_2$ Uncrystallisable and more soluble than the neutral salt — SrA'' aq (dried at 100°) Crystalline mass, v sol water — $\text{SrH}_2\text{A}''$ Crystalline pp formed by adding malic acid to a solution of the preceding Sl sol cold, m sol hot, water — CaA'' 3aq S 839 at 15°, 711 at 45°, 566 at 58°, 600 at 65°, 663 at 72°, 737 at 86° (Iwig a Hecht, A 233, 170) Granular crystalline mass. When a solution of malic acid is added to lime water a pasty mass of $\text{A}''\text{Ca}_2(\text{OH})_2$ 9aq (?) is first formed, but when the theoretical quantity of malic acid has been added this dissolves up again, and the clear liquid deposits nodules of CaA'' 3aq in the course of 24 hours — $\text{CaH}_2\text{A}''$ 6aq (Hagen, A 38, 263) Occurs in stems of *Geranium zonale*, the berries of *Rhus glabra* (Rogers, Am S 27, 294), and in tobacco. Separates in well formed trimetric

octahedra from a solution of the neutral salt in warm dilute nitric acid S (of $\text{CaH}_2\text{A}''$) 1287 at 15°, 8514 at 45°, 32236 at 57°, 13127 at 68°, 7437 at 78° (I a H) [α]_D = +5 (Bremer, R T C 3, 164) When the solution obtained by neutralising malic acid with CaCO_3 is boiled a nearly insoluble granular pp is got, which consists of CaA'' aq (Richardson a Meundorf, A 26, 135), or of CaA'' (Hagen, A 38, 257) — MgA'' 6aq Prisms, which separate from a highly concentrated solution (Liebig, A 5, 148) Alcohol ppts MgA'' — $\text{MgH}_2\text{A}''$ 4aq (or 3aq) Flattened prisms — ZnA'' 3aq Monoclinic crystals (Handl, J 1859, 289) Slowly deposited in the cold from the product of solution of zinc carbonate in aqueous malic acid at 30° When these substances are boiled together a basic salt $\text{ZnO}(\text{ZnA}'')_2$ 4aq is deposited as a jelly, changing to a sandy powder, while the filtrate deposits ZnA'' 3aq as hard, four sided prisms S 17 at 20° — $\text{ZnH}_2\text{A}''$ 2aq Elongated octahedra S 4 in the cold — The neutral manganese salt is very soluble and uncrystallisable, by adding malic acid to its solution the acid salt is p.p.d. as a white powder, S 25 From boiling water it separates in rose coloured crystals — The neutral and acid ferric salts are gummy masses, v sol water and alcohol — CuA'' aq green gummy mass, v sol water — $\text{CuH}_2\text{A}''$ 2aq blue crystals, obtained by evaporation at 40° — $\text{CuO}(\text{CuA}'')_2$ 4aq green insoluble powder obtained by boiling aqueous malic acid with cupric carbonate. If evaporated at 45° dark green crystals of $\text{CuO}(\text{CuA}'')_2$ 6aq may be got H Schulze (Ar Ph [2] 57, 273) obtained green crystals of a double salt of cupric malate and ammonium sulphate — PbA'' 3aq Obtained by ppg neutral lead acetate with a solution of calcium or potassium malate, as a white curdy pp, which slowly changes to radiating four sided needles. Melts in boiling water V sl sol cold, m sol boiling, water, from which it is deposited in needles. Dissolves easily in nitric acid, acetic and malic acids do not dissolve it more readily than water. When dry it does not melt at 170° but at 220° it forms lead fumarate — PbA'' aq (dried at 100°) (Otto, A 127, 175) — $\text{PbO}(\text{PbA}')$ — Obtained by digesting the neutral salt with ammonia, or by dropping a solution of a neutral malate into a boiling solution of lead subacetate. Amorphous pp, not becoming crystalline. Does not melt under water. Melts under hot dilute HOAc , evidently changing to the neutral salt. Nearly insol water, to which, however, it imparts an alkaline reaction — AgA'' White granular pp, formed by adding silver nitrate to a solution of neutral or acid ammonium malate — Aniline malate $\text{C}_6\text{H}_5\text{NH}_2\text{HA}''$ [144°] White prisms (from alcohol) On dry distillation it gives water, aniline, the phenyl imide of phenyl aspartic acid [211°], and the phenyl-imide of maleic acid [91°] (Anschutz a Wirtz, Am 9, 237, A 239, 140)

Acetyl derivative $\text{C}_4\text{H}_5(\text{OAc})(\text{CO}_2\text{H})$, [132°] From the acetyl derivative of the anhydride and water (Anschutz, A 254, 165)

Di-methyl ether $\text{Me}_2\text{A}''$ (122 at 10 mm)

Δ g $\frac{24.6}{24.6}$ 12386 Formed, together with MeHA'' , by passing HCl into a cooled solution of malic acid in MeOH (Demondésir, A 80, 301; C. R

23, 227, Anschütz, *B* 14, 2790, 18, 1953) — $\text{Ca}(\text{MeA})_2$, is sol alcohol

Acetyl derivative of the di methyl ether $\text{C}_2\text{H}_5(\text{OAc})(\text{CO}_2\text{Me})_2$ (129° at 11 mm)

From the ether and AcCl

Di-ethyl ether Et₂A (128° at 10 mm)

Formed in like manner Can only be distilled *in vacuo* — $\text{Ca}(\text{EtA})_2$, is sol alcohol

Acetyl derivative of the di ethylether $\text{C}_2\text{H}_5(\text{OAc})(\text{CO}_2\text{Et})_2$ (137° at 12 mm)

Nitroxyl derivative of the di-ethyl ether $\text{C}_2\text{H}_5(\text{ONO})(\text{CO}_2\text{Et})_2$ SG 1^a 202 From the ether, fuming HNO_3 , and conc H_2SO_4 (Henry, *B* 3, 532) Thick oil

Di-n-propyl ether $\text{C}_2\text{H}_5(\text{OH})(\text{CO}_2\text{Pr})_2$ (151° at 10 mm) (Anschutz)

Acetyl derivative of the di-n propyl ether $\text{C}_2\text{H}_5(\text{OAc})(\text{CO}_2\text{Pr})_2$ (157° at 12 mm)

Mono isoamyl ether $\text{C}_2\text{H}_5(\text{OH})(\text{CO}_2\text{H})(\text{CO}_2\text{C}_4\text{H}_9)$ From malic acid and isoamyl alcohol at 120° (Breunlin, *A* 91, 323) — NH_4A needles — CaA , aq plates

Acetyl derivative of the anhydride $\text{CH}(\text{OAc})\text{CO} \begin{array}{l} | \\ \text{CH}_2 - \text{CO} \end{array} \text{O}$ (54°) (161° at 14 mm)

Formed, together with maleic anhydride, by heating malic acid with AcCl (Anschutz, *B* 14, 2791) Decomposed by distillation under atmospheric pressure into HOAc and maleic anhydride

Amide $\text{C}_2\text{H}_5(\text{OH})(\text{CONH}_2)_2$ **Malamide** When gaseous NH_3 is passed into an alcoholic solution of malic ether crystals of malamide are deposited Separates from water in well defined crystals Resolved by hydrolysis into NH_3 and malic acid

Amic ether $\text{C}_2\text{H}_5(\text{OH})(\text{CO}_2\text{Et})(\text{CONH}_2)$ **Malamic ether** Formed as a crystalline mass when gaseous NH_3 is passed into malic ether (Pasteur, *J* 1853, 411)

Di-anilide $\text{C}_2\text{H}_5(\text{OH})(\text{CONHPh})_2$ [175°] Formed, together with the phenyl imide, by boiling aniline (2 mols) with malic acid (1½ mols) The phenyl imide is extracted by boiling water, and the residue recrystallised from alcohol, using animal charcoal (Arppe, *A* 96, 106) Colourless scales Nearly insol water, dilute acids, and alkalis

Mono-anilide $\text{C}_2\text{H}_5(\text{OH})(\text{CO}_2\text{H})(\text{CONHPh})$ **Malamic acid** [145°] Obtained by boiling the phenyl-imide with aqueous ammonia, the product is ppd by baryta, and the Ba salt decomposed by H_2SO_4 , avoiding any excess White granules composed of very minute needles (from alcohol), ν sol water, m sol alcohol, sl sol ether Reddens litmus Hot dilute H_2SO_4 resolves it into water and the phenyl imide Its salts are soluble in water — AgA white pp, crystallising from water in shining scales

Phenyl-imide $\text{C}_2\text{H}_5(\text{OH}) \begin{array}{l} \text{CO} \\ \diagup \quad \diagdown \\ \text{CO} \end{array} \text{NPh}$ [170°] Formed by heating malic acid with aniline and crystallising from water Groups of delicate needles, ν sol water, alcohol, and ether

Inactive malic acid $\text{CO}_2\text{HCH}_2\text{CH}(\text{OH})\text{CO}_2\text{H}$ [c 108°] (Pictet, *B* 14, 2648), [c 114°] (Kekulé), [135°] (Pasteur, Loydl, Bischoff) Occurs as calcium salt in the leaves of the common ash (*Fragaria vesicaria*) (Gintl, *J* 1868, 800, Garot, *J* 1853, 409)

Formation — 1 By mixing equal parts of laevo- and dextro malic acids (Van't Hoff, jun, *B* 18, 2170, *R T C* 4, 130) — 2 By reducing racemic acid with HIAq (Bremer, *Bl* [2] 25, 6) — 3 By the action of nitrous acid on inactive aspartic acid obtained from fumarimide (Pasteur, *A Ch* [3] 34, 46, *A* 82, 330) — 4 By the action of moist Ag_2C on bromo succinic acid (Kekulé, *A* 117, 126, 130, 24) — 5 By heating fumaric acid with a large quantity of water at 150°–200° (Jungfleisch, *Bl* [2] 30, 147) — 6 By heating fumaric acid (1 pt) with NaOH (4 pts) and water (40 pts) for 100 hours at 100° (Linne mann & Loydl, *A* 192, 80, *B* 9, 925) — 7 To gether with fumaric acid, by treating di β chloro propionic ether with a weak alcoholic solution of KCy , and boiling the product with potash (Wergo & Tanatar, *A* 174, 367) Also in like manner from di β bromo propionic acid (Tanatar, *B* 13, 160) — 8 From chloro ethanetri carboxylic ether $\text{CO}_2\text{EtCH}_2\text{CCl}(\text{CO}_2\text{Et})_2$, and dilute alcoholic KOH (Bischoff, *A* 214, 49)

Properties — Crystallises more readily than active malic acid, being less soluble in water and not deliquescent, or at any rate less deliquescent than the active acid ν e sol water, ν sol alcohol, ν sl sol ether When heated at 200° it yields fumaric acid

Salts — NH_4HA trimetric crystals (Van't Hoff) — NH_4HA aq monoclinic crystals *abc* = 5856 1 5377, β = 68° 12' — CaA granular pp, sl sol water — CaA aq — CaA 2 aq nodules of transparent crystals (Pasteur) The acid calcium salt resembles that of the active acid except that its crystals exhibit hemihedral faces The lead salt melts under water, but crystallises less easily than the inactive salt The lead salt obtained by Tanatar from di bromo propionic acid did not melt under water — AgA

By crystallising the cinchonine salt Bremer (*B* 13, 352) was able to separate it into salts of laevo and dextro malic acid

Ethyl derivative $\text{C}_2\text{H}_5(\text{OEt})(\text{CO}_2\text{H})_2$ [86°] Obtained by saponifying its di ethyl ether (Purdie, *C J* 39, 348) Transparent crystals (from ether), sol water Not ppd by lead subacetate, but in neutral and feebly acid solution it is ppd by lead nitrate and, more slowly, by neutral lead acetate Fuming HI at 120° reduces it to succinic acid — CaA insoluble — AgA sl sol water — BaA aq hygroscopic, ν sol water

Di ethyl ether of the ethyl derivative $\text{C}_2\text{H}_5(\text{OEt})(\text{CO}_2\text{Et})_2$ (135°–200°) at 250 mm From fumaric ether and NaOEt , the product being neutralised (Purdie)

Isobutyl derivative $\text{C}_2\text{H}_5(\text{OC}_4\text{H}_9)(\text{CO}_2\text{H})_2$ Crystalline, deliquescent mass (Purdie) — CaA insol water — AgA flocculent, insol water

The following compounds are probably also derivatives of the same inactive malic acid —

Amic acid $\text{CO}_2\text{HCH}_2\text{CH}(\text{OH})\text{CONH}_2$ **Malamic acid** [146°] Obtained, together with its ether, by boiling the amide of diazo succinic ether with water (Curtius, *J pr* [2] 83, 479) Colourless prisms, ν sol water, alcohol, and ether

Methyl ether of the amic acid $\text{CO}_2\text{MeCH}_2\text{CH}(\text{OH})\text{CONH}_2$ [105°] Formed,

together with methyl fumaramate, by boiling methyl diazo succinamate

$\text{CO}_2\text{Me CH}_2\text{CH}(\text{CONH}_2)$ with slightly acidified water (Curtius, *J pr* [2] 88, 482) Silky plates, v e sol alcohol, ether, and water With benzoic acid at 150° it forms crystalline $\text{CO}_2\text{Me CH}_2\text{CH}(\text{OBz})\text{CONH}_2$ [80°], v sol ether

Benzoyl derivatives of the ethyl ether of the amic acid

$\text{CO}_2\text{Et CH}_2\text{CH}(\text{OBz})\text{CONH}_2$ [97°] Formed by heating ethyl diazo succinamate with HOBz , at 145° Colourless rhombohedra

Dextro-malic acid $\text{CO}_2\text{H CH}_2\text{CH}(\text{OH})\text{CO}_2\text{H}$ [c 100°] Formed from dextro asparagine (from sprouting vetch seeds) by treatment with nitrous acid (Pittet, *B* 19, 1693) The cinchonine salt is obtained by crystallisation from the cinchonine salt of inactive malic acid Resembles ordinary malic acid, except that it is dextrorotatory

Isomeride of malic acid $\text{C}_4\text{H}_6\text{O}_5$ [155° - 168°] Formed by heating acetylene dibromide (5 pts) with KCy (4 pts) and alcohol (28 pts) at 100° , and saponifying the resulting nitrile with potash (Sabanejeff, *A* 216, 275) Crystals, v sol water— $\text{Ag}_2\text{A}''$

Isomalic acid of Kammerer (*J pr* 88, 321, *A* 139, 257) is identical with citric acid (Ostwald, *B* 21, 3534)

Paramalic acid is described as **DIGLYCOLLIC ACID**

Iso-malic acid $\text{CH}_2\text{C}(\text{OH})(\text{CO}_2\text{H})_2$ [c 140°] From bromo iso succinic acid and moist Ag_2O (Schmoger, *J pr* [2] 14, 77, 19, 168, 24, 38) Crystals, v sol water, alcohol, and ether At 170° it splits up into CO_2 and lactic acid In active Its neutral solutions give no pp with CaCl_2 (unlike methyl tartaric acid, which gives a pp on warming)— BaA'' 2aq amorphous pp S l at 100° Loses its water of crystallisation at 130° (unlike methyl tartaric acid)— PbA'' Does not melt under water— $\text{Ag}_2\text{A}''$ needles

Methyl-tartaric acid $\text{CH}_3\text{C}(\text{OH})(\text{CO}_2\text{H})_2$ [178°] Obtained from pyruvic acid by addition of KCy and HCl , the product being boiled with baryta water (Bottlinger, *B* 14, 148, 17, 144) Rhombohedral crystals Decomposes on fusion, giving off CO Boiling conc HClAq splits it up into CO and lactic acid— BaA'' aq (dried at 130°) Heavy crystalline powder, ppd on boiling— ZnA'' 2aq crystalline powder— $\text{Ag}_2\text{A}''$ aq white crystalline powder

Reference—**BROMO MALIC ACID**

MALLOTOXIN $\text{C}_{11}\text{H}_{10}\text{O}_5$ or $\text{C}_{11}\text{H}_8\text{O}_5$ May be extracted by CS_2 from finely-divided kamala, a yellow dye got from the seeds of *Mallotus Philippensis* (A G Perkin & W H Perkin, jun, *B* 19, 8109) Flesh coloured needles, insol water, v sol hot alcohol Dissolves readily in alkalis, forming a yellowish red solution Ac_2O yields a di acetyl derivative, r ROTTLEIN

MALOBURIC ACID $\text{C}_3\text{H}_4\text{N}_2\text{O}_4$ v e

$\text{CO} \begin{smallmatrix} \text{NH CO} \\ \text{NH CO} \end{smallmatrix} \text{CH CO NH}_2$ (?) Formed, as ammonium salt, by heating barbituric acid (malonyl urea) with urea at 160° (Baeyer, *A* 135, 312) Formed also by boiling cyanuromalic acid with HClAq (Nencki, *B* 5, 888) Ppd by adding HCl to solutions of its salts as a granular pp, which, when quite pure, can be obtained in crystals Dissolves completely in bromine and

water, forming di bromo barbituric acid Nitric acid converts it into nitro barbituric (dilituric) acid— KA'' long needles S l in hot water

MALONAMIC ACID v **MALONIC ACID**

MALONAMIDE v *Amide of MALONIC ACID*

MALONANILIC ACID v *Mono-amide of*

MALONIC ACID

MALONIC ACID $\text{C}_3\text{H}_4\text{O}_4$ v e $\text{CH}_2(\text{CO}_2\text{H})_2$ Mol w 104 [184°] (Krafft & Noerdlinger, *B* 22, 816) S 109 at 1° , 138 at 16° (Miczynski, *M* 7, 258), 109 at 15° (Bourgoin, *Bl* [2] 33, 423) H F 212,700 (Stohmann, Kleber & Langbein, *J pr* [2] 40, 206) H C v 207,900 H C p 207,300 (S, K & L), 208,650 (Lougumme, *C R* 107, 597) Heat of solution = -4573 Heat of neutralisation by NaOH 27,120 (Gal & Werner, *Bl* [2] 46, 803), 26,650 (Massol), by KOH 27,800, by NH_3 25,040, by BaO.H , 30,135, by CaO.H , 27,090 (Massol, *C R* 107, 257, 393, 108, 813, 1060, 109, 27) S H (0° to 50°) 2832, (0° to 110°) 3262 (Hess, *P* [2] 35, 410)

Occurrence—In beet root (Lippmann, *B* 14, 1183)

Formation—1 Discovered by Dessaignes (*C R* 47, 76) as a product of the slow oxidation of malic acid by cold aqueous $\text{K}_2\text{Cr}_2\text{O}_7$.—2 By the action of alkalis or acids on cyano acetic acid (semi nitrile of malonic acid) or cyano acetic ether (Kolbe, *A* 131, 349, Hugo Muller, *C J* 17, 109)—3 By oxidation of allylene or propylene with cold alkaline KMnO_4 (Berthelot, *J* 1867, 335)—4 By boiling barbituric acid with potash (Baeyer, *A* 130, 143, Heintzel, *A* 139, 129)—5 By boiling mucobromic acid with baryta water (Jackson & Hill)—6 From di chloro acrylic ether by treatment with Ag_2O at 125° , and saponifying the product (Wallach & Hunaeus, *A* 193, 25)—7 By treating $\text{CBr}_3\text{CO}_2\text{CH}_2\text{CBr}_3$ with fuming HNO_3 (Demole, *B* 11, 1714)

Preparation—1 From ethylic, or better from potassic, chloro acetate Chloro acetic acid (100 g) dissolved in water (200 g) is neutralised by K_2CO_3 (75 g), mixed with KCy (70 g), and warmed The reaction heats the liquid to boiling and is soon over KOH (100 g) is now added, and the liquid boiled as long as NH_3 escapes The liquid is then acidified with HCl , evaporated to dryness, and the malonic acid extracted by ether (H v Miller, *J pr* [2] 19, 326, cf Finkelstein, *A* 133, 338)—2 Chloro acetic acid (100 pts) is dissolved in twice its weight of water, and neutralised with potassium carbonate (75 pts) To the solution is added 75 to 80 pts potassium cyanide (98 p.c), and the mixture heated for two hours, with continual renewal of the evaporated water The remaining mass is decomposed with concentrated potash When the odour of ammonia has disappeared the solution is neutralised with hydrochloric acid, and ppd as a calcium salt This is decomposed with the requisite quantity of oxalic acid, the residue extracted with ether, and the solution evaporated (Conrad, *A* 204, 121)—3 By dissolving chloro acetic acid (100 gms) in double its weight of water, saturating the solution with potassium bicarbonate (110 gms), adding potassium cyanide (75 gms), and warming on the water bath At the end of the reaction double the volume of concentrated hydrochloric acid is added and the precipitation of potassium chloride rendered complete by passing in a current of HCl gas The

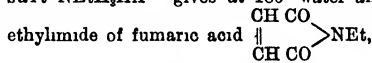
potassium and ammonium chlorides are filtered off, and the filtrate evaporated. The nearly dry residue is taken up with ether, which on evaporation yields pure malonic acid (70 gms) (Bourgoin, *A Ch* [5] 20, 271).—4 Chloro acetic ether is boiled with pure KCy dissolved in 70 p.c. alcohol for 4 hours. The alcohol is distilled off, and the residue mixed with dilute HCl and evaporated to dryness. The cyano acetic ether is extracted with ether and decomposed by fuming HClAq at 100° (Petrieff, *J R* 10, 64, *B* 7, 400).—5 In like manner from bromo acetic ether (Franchimont, *B* 7, 216).

Properties—Triclinic laminae or prisms. Decomposes a little over its melting point into CO₂ and acetic acid, but when heated under 10 mm pressure it may be sublimed. V sol water, alcohol, and ether. When heated with Ac₂O, malonic acid gives a reddish yellow liquid with greenish yellow fluorescence.—001 g is sufficient for this test (Kleemann, *B* 19, 2030). A small quantity of malonic acid treated with urea and POCl₃ yields a product which when evaporated with nitric acid leaves a residue which gives with ammonia the purple colour due to murexide (Grimaux, *C R* 88, 85).

Reactions—1 A solution of the free acid is hardly affected by the passage of an electric current. But the electrolysis of a concentrated alkaline solution gives off oxygen, CO, and CO₂ at the positive pole, without any hydrocarbon (von Miller, Bourgoin, *A Ch* [5] 20, 80, *C R* 90, 608).—2 Malonic acid (5g) heated with bromine (10 c.c.) and water (12 c.c.) for 18 hours at 120° to 145° gives bromoform, tri bromo acetic acid, CO₂, and HBr (Bourgoin, *Bl* [2] 34, 215).—3 HNO₃ (SG 1.53) in the cold gives off CO₂ (2 mols) (Franchimont, *R T C* 3, 422).—4 PCl₅ gives malonyl chloride and a compound C₂H₂ClO₂, crystallising in needles [122°], decomposed by water and alcohol with evolution of CO₂ (Béhal, *A Augur, Bl* [2] 50, 631).—5 Malonic acid (1 pt), NaOAc (1 pt), and Ac₂O (3 pts) at 100° forms a brownish yellow acid C₁₁H₂O₈. The sodium salt C₁₁H₂NaO₈ is v sol water, alkaline, and conc H₂SO₄ in sol HOAc. Its solutions exhibit fluorescence, but on heating CO₂ is evolved and the fluorescence disappears. An acetic acid solution of the salt C₁₁H₂NaO₈ boiled with a solution of phenyl hydrazine acetate forms CO₂, NaOAc, and C₁₀H₂O₈(N NHPh)₂, a substance insol water and ether, v sol alcohol and HOAc, forming a green solution in conc H₂SO₄, and melting with decomposition about 180° (Kleemann, *B* 19, 2030).—6 Benzoic aldehyde and HOAc forms benzylidene malonic acid C₆H₅CH(CO₂H)₂.—7 Propionic aldehyde and HOAc gives CH₃CH₂CH(CO₂H)₂ (Komnenos, *A* 218, 168).—8 *o*-Oxy benzoic aldehyde and HOAc at 100° forms coumarin-carboxylic acid (Vol 11 p 269).—9 Cinnamic aldehyde and HOAc at 100° gives phenyl-butene dicarboxylic acid C₆H₅CH=CHCH(CO₂H)₂ (Stuart, *C J* 49, 366).—10 Heated with phenyl thiocarbimide it gives acetanilide and the dianilide of malonic acid (Moine, *Ann Chem Farm* [4] 4, 201).

Salts—The salts, with the exception of those of the alkali metals, are sparingly soluble in water, more soluble in dilute malonic acid, HOAc, or nitric acid (Finkelstein, *A* 133, 338).—NaHA' aq: well defined crystals (F.).—

NaHA' aq trimetric prisms (Shadwell, *J* 1891, 699).—Na A' aq (Mulder, *Bl* [2] 29, 532, cf Massol, *C R* 107, 393).—KHA' aq prisms.—KHA' monoclinic prisms (Haushofer, *J* 1881, 699).—KA' 2aq deliquescent monoclinic crystals. Gives CO₂ and hydrogen on electrolysis (H von Miller).—K₂A' aq monoclinic prisms (H)—NH₄HA' very deliquescent crystals. Heat of formation from solid malonic acid and NH₄ 22,780 (Massol, *C R* 108, 1060).—(NH₄)₂A' very deliquescent needles. Heat of formation from solid malonic acid and NH₄ 41,015.—BaA' 2aq bulky flocculent pp gradually becoming crystalline. Crystallises from water in long slender needles. At 100° it loses aq, and when heated in a current of hydrogen at 140° it becomes anhydrous. BaA' aq is not dehydrated by heating in air at 150° (Pinna, Bischoff, *A* 179, 84). S 14 at 0°, 20 at 18°, 32 at 70° (Miczynsky, *M* 7, 261). Heat of solution —3830 (Massol, *C R* 109, 29).—CaA' 2aq monoclinic needles (from water). Ppd in gelatinous form on adding CaCl₂ to a solution of ammonium malonate. S 3 at 0°, 37 at 20°, 47 at 72° (Miczynsky). According to Massol (*C R* 108, 813) this salt separates in small brilliant scales on slow evaporation of its solution at 35°.—CaA' 4aq silky needles. Obtained by slow evaporation at 15° (Massol). V sol water. May be dehydrated in a current of hydrogen at 135°.—SrA' small brilliant scales (Ossipoff, *C R* 108, 815).—MgA' 2aq, MgA' aq, and MgA' 3aq are described by Finkelstein as crystalline powders.—ZnA' 2aq monoclinic crystals.—MnA' 2aq trimetric prisms.—CoA' 2aq monoclinic crystals (Haushofer).—NiA' 2aq bluish green powder.—CdA' deliquescent.—CdA' 4aq monoclinic crystals.—CdA' 12aq triclinic (Haushofer, *J* 1882, 362).—PbA' crystalline powder, sol HOAc.—CuA' 3aq blue triclinic crystals (H).—CuO A' bluish green pp.—AgA' crystalline pp, not blackened by boiling water.—Ethylamine salt NEt₃HA' gives at 180° water and the



whence concentrated caustic potash solution forms NHEt CO CH CH CO₂H [126°] (Piutti, *C C* 1888, 1529).

Methyl ether Me₂A' (181.5° cor) SG 1.1603, $\frac{d}{4}$ 1.1511 (Perkin, *C J* 45, 509), $\frac{d}{4}$ 1.1753 (W.) M M 5.28 at 17° SV 137.6 (Wiens, *A* 253, 297). From silver malonate and MeI (Osterland, *B* 7, 1286). Oil, sol alcohol and ether.

Mono-ethylether CO₂Et CH₂ CO₂H. When alcoholic KOH (1 mol) is added to alcoholic malonic ether the liquid forms a crystalline pulp of CO₂Et CH₂ CO₂K composed of needles (Van't Hoff, *B* 7, 1571, Freund, *B* 17, 780). PCl₅ converts it into CO₂Et CH₂ COCl (170°–180°).

Ethyl ether CH₃(CO₂Et)₂ (198° cor) (Perkin, *C J* 45, 508), $\frac{d}{4}$ 1.0761 (Wiens) SG 1.0610, $\frac{d}{4}$ 1.0525 M M 7.41 at 14° SH 439 at 0°, 45 between 10.6° and 82.2° (R Schiff, *Zent Phys Chem* 1, 376, *G* 16, 454) SV 185.1 (Wiens, *A* 253, 297).

Preparation.—1 Calcium malonate is boiled with H₂SO₄ and 4 times the theoretical quantity of alcohol for 24 hours (Conrad, *B* 12, 749).—

2 Chloro acetic acid (250 g) is dissolved in water (500 g) and K_2CO_3 (187 g), KOy (175 g) is added, and the whole heated on a sand bath until the reaction begins. The product is evaporated until its temperature is 135° , and is then allowed to cool. When cold it is treated with two-thirds of its weight of alcohol and gaseous HCl is passed in. The product is poured into iced water, extracted with ether, dried over $CaCl_2$, and distilled (Venable a Claisen, *A* 288, 131). Tricarballic ether (287°) is obtained as a by product (Daumichen, *C C* 1888, 1347).

Reactions—1 Water at 150° gives CO_2 and acetic ether (Hjelt, *B* 13, 1949).—2 Boiling with *m* amido benzoic acid and a little alcohol forms $CH_3(CO NH C_6H_4 CO_2 H)_2$, a powder which will not melt and is insol water, hardly sol boiling alcohol, sol dilute NH_4Ag , and dissolves without change in conc H_2SO_4 (Schiff, *A* 232, 143, *B* 17, 403, *G* 15, 534). The intermediate compound $CO_2 Et CH_2 CO NH C_6H_4 CO_2 H$ [173°] is also formed at the same time.—3 Chlorine forms chloro malonic ether.—4 Cautious treatment with nitric acid (5 pts of SG 1.5) gives nitro malonic ether (Franchimont a Klobbie, *R T C* 8, 283).—5 According to Lang (*B* 19, 2937) zinc methyl or zinc ethyl act in the cold, forming phloroglucin tricarboxylic ether, methane, and ethane.—6 When a mixture of malonic ether (1 mol) and EtI (2 mols) is heated with zinc there is formed ethane and ethyl-malonic ether (90 p.c. of the theoretical amount) (Damlar, *A* 20, 203). When malonic ether (20 g), EtI (100 g), and granulated zinc are heated with inverted condenser at 100° there is formed di ethyl malonic ether (Joulowsky, *J pr* [2] 39, 446).—7 With allyl iodide and zinc there is formed di-allyl malonic ether and propylene (Matvéeff, *J pr* 39, 452).—8 When malonic ether (2 mols) is treated with glyoxal (1 mol) and zinc chloride it forms di oxy butane tetracarboxylic ether $(CO_2 Et)_2 CH CH(OH) CH(OH) CH(CO_2 Et)_2$ (Polonowsky, *A* 246, 1).—9 With benzoic aldehyde and HCl it forms benzylidene malonic ether $C_6H_5 CH C(CO_2 Et)_2$ (Claisen a Cremer, *A* 218, 129).—10 Furfuraldehyde and acetic anhydride give furfuryl methenyl malonic ether $C_4H_3O CH C(CO_2 Et)_2$, which boils with slight decomposition at 293° , and gives on saponification the acid ether $C_4H_3O CH C(CO_2 H)(CO_2 Et)$ [102.5°], and finally the acid $C_4H_3O CH C(CO_2 H)_2$ [187°], which may be reduced by sodium amalgam to $C_4H_3O CH_2 CH(CO_2 H)_2$ [125°] (Marckwald, *B* 21, 1081).—11 Aldehyde forms $CH_2 C(CO_2 Et)_2$, and $CH_2 CH_2 C(CO_2 Et)_2$ (Komnenos, *A* 218, 145).—12 Diazobenzene chloride forms benzene azo malonic acid, identical with the phenyl hydrazide of mesoxalic acid (R Meyer, *B* 21, 118).

Sodium malonic ether $CHNa(CO_2 Et)_2$ is formed, with evolution of hydrogen, by dissolving sodium in malonic ether. The reaction is stopped by the crust of the sodium compound which forms on the surface of the metal, but this may be removed by adding alcohol (2 vols) which first forms $NaOEt$, and this then reacts with the malonic ether. The sodium malonic ether crystallises on cooling, and may be freed from alcohol by heating to 150° in a current of hydrogen (Conrad, *B* 12, 750). If, however, it be required merely for synthetical purposes the presence of alcohol is for the most part not injurious, and

the mixture of malonic ether and alcoholic $NaOEt$ may be used at once. This reacts upon organic halogen compounds thus—

$CHNa(CO_2 Et)_2 + XI = CHX(CO_2 Et)_2 + NaI$, the reaction being energetic at first but often requiring heat to finish it. If the substances are mixed in the right proportion the end of the reaction is recognised by the solution becoming neutral. Water is then added, and if X is a hydrocarbon radicle, $CHX(CO_2 Et)_2$, separates as an oil, which may be purified by fractional distillation. The resulting $CHX(CO_2 Et)_2$ still contains hydrogen displaceable by sodium, and if it be mixed with alcoholic $NaOEt$ and another halogen compound say YI , the reaction

$CNaX(CO_2 Et)_2 + YI = NaI + CXY(CO_2 Et)_2$, takes place (Conrad a Bischoff, *A* 204, 121). When the substituted malonic ethers are saponified acids are obtained, which at 150° or 160° give off CO_2 , and leave derivatives of acetic acid $CHX(CO_2 H)_2 = CO_2 + CH X CO_2 H$, $CXY(CO_2 H)_2 = CO_2 + CHXY CO_2 H$.

Hence malonic ether may be used, like acetoacetic ether, to effect the synthesis of organic acids of the general formula $CXYH CO_2 H$ where X and Y may be alkyls, groups like $CH_2 CO_2 Et$, or other radicles. When it is desired to prepare a di alkyl malonic ether it is not necessary to perform the operation in two stages, for the proper quantity of $NaOEt$ (2 mols) may be added all at once, and then the alkyl iodide (2 mols). By the action of $AcCl$ on sodio malonic ether in ethereal solution it may be converted into acetyl-malonic ether (120° at 17 mm), which is soluble in KOH aq., the yield is 55 p.c. of its weight. This ether forms an oxim and a phenyl hydrazide and therefore appears to be $CH_3 CO CH(CO_2 Et)_2$. When sodio aceto-acetic ether is acted upon by chloroformic ether $ClCO_2 Et$ the same ketonic ether $CH_3 CO CH(CO_2 Et)_2$ should be formed, but the product is found to be insol conc KOH aq., and to boil at a higher temperature (127° at 17 mm). It is decomposed by cold dilute KOH , and gives CO_2 and alcohol. The latter compound would therefore appear to be $CH_3 C(OCO_2 Et) CH CO_2 Et$ (Michael, *Am* 10, 158, *J pr* [2] 37, 473, cf Lange, *B* 20, 1325).

Reactions—1 EtI gives ethyl malonic ether. 2 Benzyl chloride gives mono and di benzyl malonic ether and regenerated malonic ether (Bischoff a Siebert, *A* 239, 94).—3 Malonic ether (32 g), sodium (9.2 g), dry alcohol (200 g) and chloroform (12 g) react thus (Conrad a Guthzeit, *A* 222, 250) $2CNa(CO_2 Et)_2 + CHCl_3 = 3NaCl + CNa(CO_2 Et)_2 + CH(CO_2 Et)_2$, forming sodium di carboxy glutaric ether (q.v.).—4 Trimethylene bromide reacts with formation of $(CO_2 Et)_2 C < \begin{smallmatrix} CH_2 \\ CH \end{smallmatrix} > CH_2$ (Perkin, jun, *C J* 51, 1, 702, 820).—5 $NaOEt$, allyl iodide, and iso butyl iodide gives an acid $C_8H_{14}O_4$, instead of allyl isobutyl-malonic acid.—6 Chloro-formic ether $ClCO_2 Et$ forms methane tricarboxylic ether $CH(CO_2 Et)_3$ (Conrad a Bischoff, *A* 214, 31, Claisen, *B* 21, 3567).—7 Tri bromo-di-nitro-benzene dissolved in ether reacts with formation of $C_6H_2Br(NO_2)_2 CH(CO_2 Et)_2$ [75°], although bromobenzene does not react (Jackson a Robinson, *B* 21, 2034).—8 Phthalyl chloride (1 mol) added to sodium-malonic ether (2 mols) dissolved in ether forms phthalyl malonic ether $C_8H_8O_4$,

[76°], phthalyl-dimalonic ether $C_{12}H_{10}O_8$ [48 5°], and 'phthaloxyl dimalonic' ether $C_{12}H_{10}O_8$, 116 5° (Wishnoen, *A* 242, 23) — 9 Treatment with the *chloride of mono-ethyl phthalate* $CO_2Et \cdot C_6H_4 \cdot COCl$ dissolved in benzene forms $(CO_2Et \cdot C_6H_4 \cdot CO)_2 \cdot C(CO_2Et)_2$ [above 180°] (Zelinsky, *B* 20, 1012) — 10 In alcoholic solution it is probably converted by carbonic acid gas into $(CO_2Et)_2CH \cdot CO_2Na$ (Michael, *J pr* [2] 35, 458) — 11 *Carbon disulphide* added to an alcoholic solution of sodium malonic ether gives yellow plates of $(CO_2Et)_2CNa \cdot CS_2Na$ — 12 SO_2 acts according to the equation $SO_2 + CHNa(CO_2Et)_2 + HOEt = CH_2(CO_2Et)_2 + NaO \cdot SO \cdot OEt$ — 13 *Resorcin* dissolved in alcoholic sodium malonic ether gives a greenish blue fluorescence, when after a few days the liquid is poured into water and acidified a small pp of needles of a condensation product $C_{11}H_{10}O_8$ [191°] is got. A little above its melting point it is split up into CO_2 and (β) methyl um belliferone of which it is probably the carboxylic acid (Michael, *J pr* [2] 37, 469) — 14 *Sulphur* dissolves in alcoholic sodium-malonic ether, and at 100° crystals separate. From these, acids liberate a stinking oil (thio tartaric ether?) — 15 With alcohol and *cinnamic ether* at 100° it forms $C_{11}H_{10}O_8$ (305°–310°). It is probably $Ph \cdot CH(CHNa \cdot CO_2Et) \cdot CH(CO_2Et) \cdot CO_2Et$, for on saponifying and heating the resulting acid, phenyl glutaric acid $CO_2H \cdot CH_2 \cdot CHPh \cdot CH_2 \cdot CO_2H$ is got (Michael, *J pr* [2] 35, 349) — 16 *Phenyl cyanate* $PhNCO$ acts violently upon an alcoholic solution of sodium malonic ether with formation of $(CO_2Et)_2CH \cdot CONHPh$ [124°] and the salt of an acid $C_{12}H_{10}NO_8$ [172°] (Michael, *J pr* [2] 35, 452) — 17 *Phenyl thio carbimide* forms crystalline $C_{11}H_{10}Na_2SNO_8$, whence acids separate $\cdot NHPh \cdot CS \cdot CH(CO_2Et)_2$ [60°] (Michael, *Am* 9, 124) — 18 *Urea* dissolves in malonic ether and rhombic prisms of $OHNa \cdot \begin{smallmatrix} CO-NH \\ CO-NH \end{smallmatrix} \cdot CO$ separate, whence acids liberate barbituric acid (Michael, *J pr* [2] 35, 456) — 19 *Thio urea* gives similarly flat plates of sodium thio barbiturate $CHNa \cdot \begin{smallmatrix} CO-NH \\ CO-NH \end{smallmatrix} \cdot CS$, whence acids liberate thio barbituric acid, which crystallises in six sided plates from water — 20 *Acetamide* forms crystalline $C_{10}O_4N_2Na_2H_8$, the reaction which occurs being $2CHNa(CO_2Et)_2 + 2CH_3 \cdot CO \cdot NH_2 = C_2O_4Na_2N_2H_8 + CH_3 \cdot C(CO_2Et)_2 + 2HOEt$ — 21 $CSCl_2$ forms $CS \cdot C(CO_2Et)_2$, which crystallises in flesh coloured needles [178°] (Bergreen, *B* 21, 337) — 22 *Iodine* forms $(CO_2Et)_2CH \cdot CH(CO_2Et)_2$ 23 *Di bromo maleic ether* yields 'di malonyl maleic' ether [75°], whence by saponification the corresponding acid $C_{10}H_8O_{12}$ [148°] may be obtained. This acid, the formula of which is $(CO_2H)_2CH \cdot C(CO_2H) \cdot C(CO_2H) \cdot CH(CO_2H)_2$, may be better called butylene hexa-carboxylic acid. It crystallises in plates, v sol water, and forms the salts, Na_2A^{10} 10aq and Ag_2A^{11} , and the ethers Me_2A^{12} [129°] and Et_2A^{13} [75°] (Pum, *M* 9, 450) When the acid is heated it splits up into CO_2 and butylene tetracarboxylic acid $CO_2H \cdot CH_2 \cdot C(CO_2H) \cdot C(CO_2H) \cdot CH_2 \cdot CO_2H$ [176°] 24 *Dry cyanogen chloride* forms cyano malonic ether (Haller, *Ch* [6] 16, 419)

Di-sodium-malonic ether

$CNa_2(CO_2Et)_2$. Obtained by ppg malonic ether (1 mol.) with $NaOEt$ (2 mols.) (Bischoff & Rach,

B 17, 2782) Very unstable Iodine converts it into $(CO_2Et)_2O \cdot C(CO_2Et)_2$

Ethyl propyl ether EtPrA" (211°) $SG \frac{1}{2}$ 10498 SV 207 8 (Wiens, *A* 253, 297)

Propylether PrA" (228°) $SG \frac{1}{2}$ 10271 SV 234 6 (Wiens, *A* 253, 297) SH (from 11 6° to 82 3°) 453 (R Schiff, *G* 17, 286)

Butyl ether (C₄H₉)₂A" (251 5°) $SG \frac{1}{2}$ 10049 SV 269 1 (Wiens, *A* 253, 297)

Chloride $CH_2(COCl)_2$ (58° at 27 mm) From malonic acid and $CSCl_2$ heated above 100° (Béhal & Auger, *Bl* [2] 50, 594) Liquid Smells slightly like chloral

Amide $CH_2(CONH_2)_2$ [170°] (Van't Hoff, *Ar Néerl* 10, 274) SG 83 at 8° (Henry, *Bl* [2] 43, 618) From malonic ether (50 cc) by shaking with strong aqueous NH_3 (150 cc) (Osterland, *B* 7, 1286, Freund, *B* 17, 133) The yield, in 2 days, is 75 p c Silky needles (from dilute alcohol), insol alcohol and ether Boiling aqueous NH_3 converts it into ammonium ma-

lonamate Salt $\begin{smallmatrix} CO \cdot NH \\ CO \cdot NH \end{smallmatrix} \cdot Hg$ white amorphous powder, insol alcohol and ether, sl sol hot water, v sol $HClAq$ (Freund)

Di-methyl amide $CH_3(CO \cdot NHMe)_2$ [128°] (F), [125°] (Henry), [136°] (Fianchimont, *R T C* 4, 199) Formed by the action of methylamine on malonic ether (Freund, *B* 17, 133) Small flat needles v e sol water and alcohol, sl sol ether Fuming HNO_3 converts it into $CH_2(CO \cdot N(NO_2)Me)_2$ [150°]

Di-ethyl amide $CH_2(CO \cdot NHEt)_2$ [149°] Six sided tables (Wallach & Kamenski, *B* 14, 170)

Ethylene diamide $CH_2 \cdot \begin{smallmatrix} CO \cdot NH \\ CO \cdot NH \end{smallmatrix} \cdot C_2H_4$

Formed by heating malonic ether or malonamide with ethylene diamine (Freund, *B* 17, 137) Crystalline solid v sol water, nearly insol alcohol

Amide-anilide $CONH_2 \cdot CH_2 \cdot CONHPh$ [163°] Obtained by heating malonamide with 1 mol of aniline for $\frac{1}{2}$ hour at 200°–220° (Freund, *B* 17, 135) Fine white felted needles Sol hot water and alcohol

Mono anilide $CO_2H \cdot CH_2 \cdot CO \cdot NHPh$ *Phenyl-malonamic acid* *Malonamic acid* *Malonphenylamic acid* [132°] Formed by boiling the amide anilide with milk of lime (Freund, *B* 17, 135) Formed also by heating sodiumacetyl phenyl carbamate $C_6H_5Nac \cdot CO_2Na$ for 5 or 6 hours at 130°–140° under pressure (Seifert, *B* 18, 1359), and by heating malonic acid (1 mol) with aniline (1 mol) at 105° (Rugheimer, *B* 17, 737) Large colourless monoclinic crystals, or slender needles At its melting point it breaks up quantitatively into CO_2 and acetanilide — $A^{14}Ag$ small white needles — $A^{15}Ca$ 4½aq large needles

Ethyl ether of the mono-anilide $CO_2Et \cdot CH_2 \cdot CONHPh$ [89°] From aniline and $CO_2Et \cdot CH_2 \cdot COCl$, both dissolved in benzene (Rugheimer & Hoffmann, *B* 17, 739) Crystals (from ether-ligroin) l sol water and ligroin, v e sol alcohol and benzene

Anilide $CH_2(CONHPh)_2$ [225°] Obtained by boiling malonic ether or malonamide with aniline (Freund, *B* 17, 134) White needles Insol water and ether, e sol hot alcohol

Tri-bromo anilide

$\text{CH}_2(\text{CO NH C}_6\text{H}_2\text{Br}_3)_2$ [148°] White silky needles, sl sol alcohol, insol water (Freund, *B* 17, 780)

Methyl-anilide $\text{CH}_2(\text{CO NMePh})_2$ [109°]

Obtained by boiling malonic ether with an excess of methyl aniline (Freund, *B* 17, 187) Colourless trimetric prisms *V* sol alcohol

Mono o-toluide

[2] $\text{CH}_2\text{C}_6\text{H}_4\text{NH CO CH}_2\text{CO}_2\text{H}$ *o*-Tolyl malonic acid Colourless needles, *v* sol water and alcohol Melts about 140°, giving off CO_2 — CaA'_2 3aq small needles, *m* sol water— BaA'_2 aq needles, *v* sol water— CuA'_2 2aq prisms (Rugheimer & Hoffmann, *B* 18, 2971)

Ethyl ether of the o-toluide EtA' [74°]

Long needles, sl sol ether and ligroin

Mono m-toluide

[8] $\text{CH}_2\text{C}_6\text{H}_4\text{NH CO CH}_2\text{CO}_2\text{H}$ [101°]

Mono-p-toluide

[4] $\text{CH}_2\text{C}_6\text{H}_4\text{NH CO CH}_2\text{CO}_2\text{H}$ Obtained by heating malonic acid with *p*-toluidine (Rugheimer & Hoffmann, *B* 17, 740, 18, 2971) Long colourless needles, sol water, alcohol, ether, and chloroform At 150° it gives off CO_2 PCl_5 converts it into tri chloro methyl quinoline [134°]— CaA'_2 4½aq long needles, *m* sol water— BaA'_2 5aq needles— AgA'_2 cheesy white pp or needles, sl sol water— CuA'_2 2aq minute needles, sl sol hot water— ZnA'_2 glistening plates, *m* sol water

Ethyl ether of the p-Toluide EtA' plates, *v* sol alcohol

Mono phenyl hydrazide

$\text{PhN}_2\text{H CO CH}_2\text{CO}_2\text{H}$ [154°] Formed from malonic acid and aqueous phenyl hydrazine acetate at 100° (Fischer & Passmore, *B* 22, 2734) Needles, *v* sol water Its phenyl hydrazine salt $\text{PhN}_2\text{H}_2\text{CO CH}_2\text{CO}_2\text{N}_2\text{H}_2\text{Ph}$ is converted at 200° into $\text{CH}_2\text{C}(\text{CO})\text{N}_2\text{HPh}$, crystallising in white needles, [128°], *v* sol alcohol

Di phenyl dihydrazide

$\text{CH}_2(\text{CO N}_2\text{H Ph})_2$ [187°] From malonic acid or the amide of malonic acid and phenyl hydrazine at 200° (Freund & Goldsmith, *B* 21, 1241) Plates (from dilute alcohol) With COCl_2 it gives $\text{C}_6\text{H}_5\text{N}_2\text{O}_2$, which crystallises from HOAc in laminae [205°]

Semi nitrile $\text{CO}_2\text{H CH}_2\text{CN}$ *v* CYANO

ACETIC ACID

Nitrile $\text{CH}_2(\text{CN})_2$ *Methylene cyanide* [30°] (219°) (*H*), (223°) (Berthelot & Petit, *A Ch* [6] 17, 131) *HF*—43200 Obtained by heating cyano acetamide $\text{CN CH}_2\text{CONH}_2$ with P_2O_5 (Henry, *C R* 102, 1394, 1481) White solid Appears to be polymerised by prolonged action of heat Sol water, *v* sol alcohol and ether Burns with a purple edged flame Conc HClAq dissolves it with evolution of heat and formation of malonic acid HClAq at 150° in sealed tubes forms CO_2 and chloro acetic acid With ammoniacal AgNO_3 it gives a white pp $\text{CAG}_2(\text{CN})_2$ (?), which explodes when heated

References—AMIDO-, BROMO-, CHLORO-, CYANO-, NITRO-, METHYL-, ETHYL-, PROPYL-, METHYL ETHYL-, and BENZYL MALONIC ACID

MALONYL-UREA *v* BARBITURIC ACID

Isomalonylurea $\text{CO} \begin{smallmatrix} \text{NH CH} \\ \text{NH CO} \end{smallmatrix} \text{CO OH}$ is formed, together with amido uracil, by reducing

nitro-uracil (Behrend, *A* 229, 89, *B* 21, 999)

It is converted by bromine into an acid isomeric with dialuric acid The acetyl derivative $\text{C}_6\text{H}_5\text{N}_2\text{O}_2\text{Ac}$ crystallises from hot water in prisms

MALONOXYL-AMIDO-BENZENE *v* CARB OXY PHENYL MALONAMIC ACID

MALTOBIONIC ACID $\text{C}_{12}\text{H}_{22}\text{O}_{12}$ Formed by oxidising maltose (1 pt) by bromine (1 pt) in water (7 pts) (E Fischer & Meyer, *B* 22, 1941) Almost colourless syrup *V* sol water, sl sol alcohol, insol ether It reduces Fehling's solution By heating with dilute sulphuric acid it is split up into dextrose and gluconic acid

Salt— CaA'_2 hard shining mass, *v* sol water

MALTONIC ACID Identical with GLUCONIC ACID (*q v*)

MALTOSE *v* SUGAR

MALYL UREIDE, so called, *v* URAMIDO

SUCCINIC ACID

MANDELAMIDINE $\text{C}_8\text{H}_{10}\text{N}_2\text{O}$ *se*

$\text{C}_6\text{H}_5\text{CH}(\text{OH})\text{C}(\text{NH}_2)\text{NH}$ [110°] From the hydrochloride of mandelic imido ether and alcoholic NH_3 (Beyer, *J pr* [2] 31, 387) Needles, *v* sol water and alcohol, sl sol ether Very unstable— B'HCl [214°] Prisms (from cold water)

MANDELAMIDOXIM $\text{C}_8\text{H}_9\text{N}_2\text{O}_2$ *se*

$\text{C}_6\text{H}_5\text{CH}(\text{OH})\text{C}(\text{NH}_2)\text{NOH}$ [159°] From the nitrile of mandelic acid and hydroxylamine (base) (Tiemann, *B* 17, 126) Crystals (from alcohol) Insol benzene, sl sol cold, *v* sol hot, water *V e* sol aqueous acids and alkalis FeCl_3 colours its aqueous solution blood red. It does not reduce Fehling's solution

Reactions—1 The hydrochloride, mixed with conc aqueous potassium cyanate forms $\text{C}_6\text{H}_5\text{CH}(\text{OH})\text{C}(\text{NOH})\text{NH CO NH}_2$ [127°]—2 Phenyl cyanate forms the corresponding $\text{Ph CH}(\text{OH})\text{C}(\text{NOH})\text{NH CO NHPh}$ [155°]—3 Excess of AcCl , or a mixture of Ac_2O and NaOAc forms $\text{C}_6\text{H}_5\text{CH}(\text{OAc})\text{C} \begin{smallmatrix} \text{N O} \\ \text{N} \end{smallmatrix} \text{C CH}_3$

[52°]—4 COCl_2 , added to its benzene solution, forms $(\text{C}_6\text{H}_5\text{CH}(\text{OH})\text{C}(\text{NH}_2)\text{N O})_2\text{CO}$ [131°]—5 Chloroformic ether, ClCCl_3Et , gives the compound $\text{C}_6\text{H}_5\text{CH}(\text{OH})\text{C}(\text{NH}_2)\text{N O CO Et}$ [107°] Salts— NaA' needles— HA'HCl (Gross, *B* 18, 1074)

Ethyl ether EtA' [89°] Slender needles, sl sol cold water With phenyl cyanate it forms $\text{Ph CH}(\text{OH})\text{C}(\text{NOEt})\text{NH CO NHPh}$ [119°]

Benzyl ether $\text{C}_6\text{H}_5\text{CH}_2\text{A}'$ [103°] From mandelamidoxim, NaOEt , and benzyl chloride (Gross, *B* 18, 1030) Needles

Acetyl derivative

$\text{C}_6\text{H}_5\text{CH}(\text{OH})\text{C}(\text{NH}_2)\text{NOAc}$ [140°] From mandelamidoxim and Ac_2O Crystals (from alcohol) Insol cold water, sol alcohol, ether, and benzene With water at 100° it forms

$\text{C}_6\text{H}_5\text{CH}(\text{OH})\text{C} \begin{smallmatrix} \text{N O} \\ \text{N} \end{smallmatrix} \text{C CH}_3$ [65°]

Di-acetyl derivative

$\text{C}_6\text{H}_5\text{CH}(\text{OAc})\text{C}(\text{NH}_2)\text{NOAc}$ [113°] From mandelamidoxim and a slight excess of AcCl (from alcohol)

Benzoyl derivative

$\text{C}_6\text{H}_5\text{CH}(\text{OH})\text{C}(\text{NH}_2)\text{NOBz}$ [149°] From mandelamidoxim and BaCl (1 mol.) With

AcCl it gives $C_6H_5CH(OAc)C(NH_2)NOBz$ [165°] (Gross)

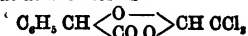
MANDELIC ACID $C_6H_5O_2$, *rac*
 $C_6H_5CH(OH)CO_2H$ *Phenyl-glycolic acid*
Oxy-phenyl acetic acid Mol w 152 [115°]
(Müller, *Ar Ph* [3] 2, 385), [118°] (Glaisen, *B* 10, 847, Lewkovitch, *B* 16, 1568) S G ± 1.361 (Schröder, *B* 12, 1612) S 16 at 20° *Heat of solution* -3100 *Heat of neutralisation* by NaOH +13860 (Berthelot, *A Ch* [6] 7, 185)

Formation -1 Discovered by Winckler (*A* 18, 310), who obtained it by heating bitter almond water with HCl, the benzoic aldehyde reacting with the HCy present (Liebig, *A* 18, 319) -2 By heating amygdalin with fuming HClAq (Wöhler, *A* 66, 238) -3 By boiling the compound of benzoic aldehyde with $KHSO_4$ for several hours with KCy and alcohol The nitrile $C_6H_5CH(OH)CN$ thus formed is saponified by dilute HClAq (O Müller, *B* 4, 980) -4 By reducing phenylglyoxylic acid $C_6H_5COCO_2H$ with sodium-amalgam (Schwebel, *B* 10, 2045) -5 From di-bromo acetophenone $C_6H_5COCHBr_2$ by boiling with dilute KOHAq (120), the compound $C_6H_5COCH(OH)_2$ being a theoretical intermediate product (Engler a Wöhler, *B* 20, 2202) -6 In small quantity, by boiling pseudo-phenyl hydantoin with baryta (Pinner, *B* 21, 2327) -7 By boiling a chloro-phenyl-acetic acid with alkalis (Spiegel, *B* 14, 239)

Preparation - Benzoic aldehyde (100 g), water (3500 c c), fuming HClAq (200 g), and 4 times the calculated quantity of hydrogen cyanide are boiled for 36 hours The product is evaporated at 100°, the residue extracted with ether, the extract evaporated, and the mandelic acid left recrystallised from water (Wallach, *A* 193, 38, cf Strecker, *A* 75, 27)

Properties - Large trimetric crystals, v sol water, alcohol, and ether Inactive to light By crystallisation of the cinchonine salt it can be separated into equal quantities of the dextro- and levorotatory acids If *Penicillium glaucum* is grown in it the levorotatory acid is destroyed, leaving the dextrorotatory (Lewkowsch, *B* 16, 1568)

Reactions -1 $KMnO_4$ and KOH convert it, in the cold, into phenyl glyoxylic acid (R Meyer a A Baur, *A* 220, 39) -2 Dry distillation yields benzoic aldehyde -3 Boiling with MnO_2 and H_2SO_4 yields benzoic aldehyde and CO_2 -4 Fuming $HBrAq$ converts it slowly in the cold, quickly at 125°, into a bromo-phenyl-acetic acid, whence alcoholic NaOEt forms the ethyl derivative of mandelic acid (Glaser a Radziszewsky, *Z* [2] 4, 140) -5 Fuming HClAq at 140° gives α -chloro-phenyl-acetic acid -6 Phosphorus and HI reduce it to phenyl acetic acid -7 When taken internally it passes unaltered into the urine (Schotten, *H* 8, 68) -8 Cannot be nitrated Conc NHO_3 forms benzoic aldehyde (Liebig, *A* 18, 321), dilute HNO_3 forms phenyl glyoxylic acid (Zincke a Hannaus, *B* 10, 1488) -9 Dilute H_2SO_4 in sealed tubes at 130° converts mandelic acid nearly quantitatively into benzoic aldehyde and formic acid (Biedermann, *B* 19, 638) 10 Chloral at 120° forms



which forms large transparent crystals [83°], insol water, sol alcohol and chloroform (Wal-

lach, *A* 193, 1) -11 *Phenyl-hydrazine* forms a compound [182°], crystallising in needles and almost insol boiling water (Reissert a Kayser, *B* 22, 2928)

Salts - The ammonium and potassium salts are very soluble and difficult to crystallise The Ba salt forms small needles S 8 at 24°, 16 at 100° (Zinn, *Z* 1868, 710) The lead salt is a crystalline powder, scarcely sol water - CuA' , (dried at 100°) - AgA' crystalline pp May be crystallised from water

Methylether MeA' [48°] Small laminae (from benzene-ligroin) (Zincke a Breuer, *B* 13, 636)

Ethyl ether $PhCH(OH)CO_2Et$ (254°) Formed by the action of water on the hydrochloride of mandelic imido ether (*qv*) Solidifies in a freezing mixture, but is liquid at ordinary temperatures (Beyer, *J pr* [2] 31, 389) But Naquet and Lugunin (*A* 139, 300), who prepared it from silver mandelate and EtI, say it melts at 75°

Methyl derivative $PhCH(OMe)CO_2H$ [72°] Formed from $PhCHClCO_2Me$, $MeOH$, and $NaOMe$ (R Meyer a H Boner, *A* 220, 44, *B* 14, 2392) Needles grouped concentrically (on solidifying) or thick tables (from light petroleum) V sol alcohol or ether, sl sol cold water or cold petroleum KOH and $KMnO_4$ forms phenyl glyoxylic acid

Salts - $NaA'2aq$ - $BaA'2aq$ - CaA' - $CuA'2aq$ - AgA'

Methylether of the methyl derivative $PhCH(OMe)CO_2Me$ (246° cor)

Ethyl derivative $C_6H_5CH(OEt)CO_2H$ From $C_6H_5CHBrCO_2H$ and alcoholic KOEt Viscid mass - AgA' pulverulent pp

Phenyl derivative $PhCH(OPh)CO_2H$ [108°] From methyl α -chloro phenyl acetate and sodium phenate (R Meyer a H Boner, *A* 220, 51) Radiating groups of slender needles (from water) V sl sol cold water, v e sol alcohol or ether KOH and $KMnO_4$ convert it into phenyl glyoxylic acid HNO_3 forms picric acid and benzoic aldehyde - $NaA'3aq$ - CuA' - AgA'

Acetyl derivative of the ethyl ether $C_6H_5CH(OAc)CO_2Et$ [74°] From mandelic acid by successive treatment with AcCl and alcohol (Naquet a Lugunin, *A* 139, 302) Slender needles (from ether) Insol water, v sol alcohol and ether

Amide $C_6H_5CH(OH)CONH_2$ [132°] S 3 at 24° S (boiling 93° c alcohol) 100 (Z) Formed, together with benzoic aldehyde, by heating the compound $(C_6H_5CHO)CNH$ with water or alcohol at 180° (Zinn, *Z* [2] 4, 709) Formed also by allowing a mixture of the nitrile with fuming HClAq to stand in the cold (Tiemann a Friedlander, *B* 14, 1967) Likewise obtained by the action of NH_3 on mandelic ether, and by heating mandelic imido-ether (C Beyer, *J pr* [2] 31, 386) Prismatic needles or plates Sol hot, sl sol cold, water, sl sol ether, v sol alcohol Decomposed by acids and alkalis with production of mandelic acid A polymeride (?) melts at 190°

Nitrile $C_6H_5CH(OH)CN$ [-10°] S G. 1.124 Obtained by adding fuming HClAq to a mixture of benzoic aldehyde and KCy (Spiegel, *B* 14, 239, Völkel, *A* 52, 861) Oil, sol alco-

hol and ether Dissolved in ether, mixed with alcohol (1 equivalent), and treated with HCl gas it forms $C_6H_5CH(OH)C(OEt)NH_2 \cdot HCl$ [125°] This is decomposed by water, forming NH_4Cl and mandelic ether (Beyer, *J pr* [2] 28, 190) At 170° mandelonitrile splits up into benzoic aldehyde and H_2C_4 Boiling aqueous HCl gives NH_4Cl and mandelic acid Fuming $HClAq$ forms the amide in the cold, but on heating it gives a chloro phenyl acetic acid NH_2 in the cold forms $C_6H_5CH(NH_2)CN$ Methylamine yields $C_6H_5CH(NHMe)CN$ Phenyl hydrazine produces the phenyl hydrazone of benzoic aldehyde (Reissert, *B* 17, 1451)

(*dextro*) Mandelic acid [133° cor] $[a]_D$ at 20° = +156 Prepared by converting inactive mandelic acid into the cinchonine salt and adding a crystal of cinchonine *dextro* mandelate to the aqueous solution when the *dextro* salt crystallises out, leaving the *laevo* salt in solution It can also be obtained by growing *Penicillium glaucum* in the inactive acid, which destroys the *laevorotatory* acid, leaving the *dextrorotatory* (Lewkowitsch, *B* 16, 1568) Resembles the *laevo*-acid, having the same solubility in water

(*laevo*) Mandelic acid [133°] *S* 864 at 20° $[a]_D$ at 20° = -158 Prepared by heating amygdalin with strong HCl for several hours on the water bath It can also be obtained from inactive mandelic acid which can be separated into equal quantities of the *laevo* and *dextrorotatory* acids by crystallisation of the cinchonine salt (Lewkowitsch, *B* 16, 1565, cf Wöhler, *A* 66, 240)

Nitro mandelic acid *v* Nitro oxy-phenyl-acetic acid

MANDELIC IMIDO ETHER

$PhCH(OH)C(NH)OEt$ [72°]

Preparation—Benzoic aldehyde (100 g) treated with KCN, dilute HCl and ether forms the cyanhydrin $PhCH(OH)CN$ which is dissolved by the ether If this is mixed with an equivalent of alcohol and dry HCl be passed in, the liquid being cooled, needles of the hydrochloride of mandelicimido ether (80g) are formed (C Beyer, *J pr* [2] 31, 3-4) $PhCH(OH)CN + EtOH + HCl = PhCH(OH)C(NH)OEt \cdot HCl$ These melt at [125°] The free ether is got by shaking these simultaneously with conc KOH and ether After evaporating the ether the residue is crystallised from ligroin

Properties—White needles Extremely soluble in ether, alcohol, and benzene

Reactions—1 At 140° the hydrochloride splits up thus $PhCH(OH)C(NH)OEt \cdot HCl = EtCl + PhCH(OH)CONH_2$, forming mandelic amide—2 Alcoholic NH_3 converts the hydrochloride into the hydrochloride of the amidine, $PhCH(OH)C(NH)NH_2 \cdot HCl$ This forms prisms [214°] Shaken with ether and potash, the free mandelic amidine, $PhCH(OH)C(NH)NH_2$, is dissolved by the ether It forms feathery needles of narcotic odour, melting at [110°]—3 Water quickly converts the hydrochloride into mandelic ether (*q v*) $C_6H_5CH(OH)C(NH)OEt \cdot HCl + H_2O = C_6H_5CH(OH)COOEt + NH_3 \cdot HCl$

MANDRAGORINE $C_8H_9NO_3$ [α 79°]

Extracted from powdered mandragora root by alcohol The extract is evaporated and the residue treated with very dilute acid The alkaloid is liberated from the acid solution by

adding K_2CO_3 and shaking with ether (Ahrens, *A* 251, 312) Hygroscopic brittle mass Picric acid gives with a solution of mandragorine hydrochloride light yellow needles of the picrate. Iodine in $KIAq$ gives an oily periodide K_2FeCy_4 gives no pp Phosphotungstate gives a white pp The sulphate is crystalline and very deliquescent Dropped into the eye, its solution causes enlargement of the pupil— $B'HANCl_4$ [155°], yellow plates, sol hot water and $HClAq$ — $B'_2H_2P_2Cl_4$ [194°] Red nodules or yellow plates (from hot water)— $B'HCl_4HgCl_2$ [160°] Plates or needles (from water) or long slender needles (from alcohol) *V* sol alcohol (Ahrens, *B* 22, 2161)

Mandragora root also contains a second alkaloid of which the platinochloride [181°] and aurochloride [147°-153°] are crystalline (Ahrens)

MANGANATES Salts of the form M'_2MnO_4 derived from the hypothetical acid H_2MnO_4 , *v*. MANGANESE, OXYACIDS *op*, p 185

MANGANESE Mn At *w* 55 Mol *w* probably same as At *w* (*v infra*) [α 1800°-1900°] *S* G 6.85 to 8.01, according to Glatzel, *S* G of pure Mn is 7.3921 at 22° (*B* 22, 2857) *S* H 14° to 97° 1217 (Regnault, *A Ch* [3] 67, 427, specimen contained Si) Chief lines in emission spectrum are 6521, 6016, 6013, 4823, 4783, 4765, 4762, 4753, 4235, 4027 (Thalén) For absorption spectrum of Mn vapour *v* Lockyer *A Roberts* (*Pr* 23, 344)

Occurrence—The metal does not occur uncombined Mn compounds are widely distributed, the chief are *pyrolusite* MnO_2 , *braunite* Mn_2O_3 , *manganite* $Mn_2O_3 \cdot H_2O$, *hausmannite* Mn_3O_4 , *psilomelane* $(Mn, Ba, K)O_4 \cdot MnO_2$, *manganesc spar* $MnCO_3$, *manganese blende* MnS Small quantities of Mn compounds are found in sea water (Forchhammer, *Pr E* 2, 303), in many mineral waters (Buchanan, *Pr* 24, 593), in blood (Cotteneau, *J* 1849 530, Burn de Buisson, *J* 1852 377, Campani, *B* 5, 287), in the liver (Béchamp, *C R* 49, 895), in milk (Polacci, *Naturforscher*, 4, 122), in human urine (Horsford, *J* 1851 602), in wines, cereals, most vegetables used as human food, and in considerable quantities in tea (Vaumené, *C R* 98, 1056, 1416) Mn also occurs in the sun's atmosphere (Comu, *C R* 86, 315, 530)

Manganese dioxide was recognised as a compound of a distinctive metal by Scheele in 1774, it had previously been looked on as a compound of iron The metal Mn was first isolated by Gahn Native MnO was long known as *magnesia nigra* (probably because of its supposed magnetic properties), a new medicine was introduced in the early years of the eighteenth century, and was called *magnesia alba*, seemingly in contradistinction to *magnesia nigra*, when *magnesia nigra* was shown to contain a distinctive metal, this metal was called sometimes *magnesium* and sometimes *manganesium*, finally the name *magnesium* was retained for the metal of *magnesia alba*, and the name *manganesium* (hence *manganese*) was given to the metal of *magnesia nigra*

Formation—1 By reducing the oxides by C at a white heat.—2 By reducing MnF_3 or $MnCl_3$ by Na or by Mg—3 By heating Mn amalgam in a stream of H, the amalgam is made by the reaction of Na amalgam with $MnCl_2Aq$ (Giles,

P M [4] 24, 328, Roussin, *Bl* 6, 93) —4 By electrolysis MnCl_2 in a porous cell placed in a carbon crucible containing HClAq (Bunsen, *P.* 91, 619)

Preparation —1 Crystallised MnCl_2 is thoroughly dried by heating, it is then finely powdered and 100 grams are intimately mixed with 200 grams well dried and powdered KCl , the mixture is packed into a Hessian crucible which is loosely covered and heated in an air-furnace until the contents melt (the temperature must not be raised so high that white vapours begin to come off), the lid is removed and 15 grams Mg are thrown into the crucible in four or five portions, each weighing 3 to 4 grams, two to three minutes being allowed to elapse between the entrance of each piece, the lid is now replaced and the crucible is strongly heated for a few minutes, and then allowed to cool very slowly in the furnace. About 20 to 25 grams Mn are thus obtained as a compact regulus. If the temperature of the final heating is not sufficiently high, the regulus does not form a compact mass, if the temperature is too high and the heating is unduly prolonged, the KCl is vapourised and the surface of the Mn is oxidised (Glatzel, *B* 22, 2857) —2 Brunner (*P* 101, 264) recommends the following method —2 pts MnF_2 (obtained by dissolving moist MnCO_3 in HFAq , evaporating and drying at 100°) and 1 pt Na are arranged in alternate thin layers in a Hessian crucible, the mixture is pressed down and covered with NaCl over which is placed a layer of CaF_2 in small pieces (to prevent spitting), the crucible is covered and heated in an air furnace, at first gently, and then to near a white heat for about $\frac{1}{2}$ an hour, the crucible is then allowed to cool very slowly in the furnace —3 For an account of the older methods of preparing Mn from MnO_2 , *v* John (Gehlen's *Journ. Chem. Phys.* 3, 452), and Deville (*A Ch* [3] 46, 182). Tamm describes a method for obtaining approximately pure metal (99.91 p.c. Mn , 0.05 p.c. Fe , 1.15 p.c. Si , and 0.25 p.c. C) from MnO_2 (*C N* 26, 73, 111)

Properties —A white grey, lustrous, metal, very hard, brittle, may be highly polished, non magnetic (Glatzel). According to Glatzel (*B* 22, 2857), Mn prepared by reducing MnCl_2 by Mg (*v* Preparation No 1) is unchanged by keeping for months in a bottle closed with a glass stopper, but in moist air the surface undergoes slight oxidation. Mn is usually described as very easily oxidised in ordinary air, and as capable of decomposing water, with evolution of H , almost as rapidly as K . Mn obtained by Brunner (*v* Preparation No 2) by reducing MnF_2 by Na was scarcely oxidised in cold water. According to Bullock (*C N* 60, 20), Mn prepared by reducing the oxides by C is very easily oxidised, while specimens obtained by reducing MnCl_2 by Na are no more oxidisable than iron. It is probable that some specimens have contained small traces of Si and C which have affected the properties of the metal. Mn melts at a very high temperature ($c. 1800^\circ - 1900^\circ$), and is said to volatilise at a full white heat.

The atomic weight of Mn has been determined (1) by estimating Cl in MnCl_2 (Arfvedson, *S* 42, 202, Dumas, *A Ch* [3] 55, 151; Berzelius, *P* 18, 74), (2) by dissolving Mn in HNO_3 ,

and calcining the nitrate (Berzelius, *P* 8, 185), (3) by oxidising MnO to Mn_2O_3 by heating in air (*v* Hauer, *W A* 25, 133), (4) by reducing Mn_2O_3 in H , and weighing H_2O produced (Rawack, *P* 107, 605, 616), (5) by analysing Mn_2O_3 (Sohneider, *P* 107, 605), (6) by reducing AgMnO , and estimating Ag produced (Dewar, *A Scott*, *Pr* 35, 44), (7) by determining SH (Regnault, *A Ch* [3] 67, 427)

Molecular weight of manganese —Ramsay (*O J* 55, 521) has determined the lowering of the vapour pressure of Hg produced by dissolving Mn in Hg , the results render it probable that the molecular weight of Mn is the same as the atomic weight. This conclusion assumes the accuracy of Van't Hoff's law that equal volumes of dilute solutions contain equal numbers of molecules of the dissolved substances, and it also supposes the molecular weight of liquid Hg to be the same as the atomic weight of Hg .

Mn is metallic in its physical, and some of its chemical, properties, but in many of its reactions it behaves as a non metal. The oxides MnO and Mn_2O_3 are basic, forming salts, such as MnSO_4 and $\text{Mn}(\text{SO}_4)_2$, of which the managanous salts, corresponding with MnO , are much the more stable. The oxide MnO reacts as a basic peroxide, *e.g.* with H_2SO_4 it forms MnSO_4 , and O , it also reacts with some positive oxides, *e.g.* CaO , to produce salts of the form $x\text{MO} \cdot y\text{MnO}$, when this oxide is treated with molten KOH in presence of O the salt K.MnO_4 is formed, in which Mn forms part of the acidic radicle. The oxides MnO , and Mn_2O_3 have also been isolated, both are very unstable, both react with water to form managanic acid HMnO_3 , the former (MnO_2) at the same time also producing MnO_2 , these oxides are distinctly acidic. The acid HMnO_4 has been isolated, and many salts derived from the hypothetical acid H.MnO_4 are known.

Mn shows marked analogies with the iron metals Fe , Ni , and Co , it is also related, although less distinctly, to the halogens, and it is also related to the chromium metals, Cr , Mo , W , and U . In the classification of the elements based on the periodic law, Mn occupies a position different from that of any known element: it is placed in the same group as the halogens (Group VII), and in the same family (even-series members) as F , but no member of this family coming after Mn (*i.e.* with a greater atomic weight than Mn) has yet been isolated, Mn finds a place in series 4 (K , Ca , Sc , Ti , V , Cr , Mn), all the members of which are metallic, and it is immediately followed by Fe , Ni , Co , and Cu . The strongly marked negative character of the halogens is impressed on Mn , but this is counterbalanced by the positive characteristics of the metals which come before and after Mn in order of atomic weights (*v* HALOGEN ELEMENTS, vol. II p. 666, also IRON GROUP OF ELEMENTS, this vol. p. 65).

The atom of Mn is divalent in the gaseous molecule MnCl_2 .

Reactions —1 Mn is oxidised in air or oxygen, forming MnO —2 Mn reacts energetically with dilute mineral acids forming salts of MnO 3 Fused with potassium nitrate or chlorate KMnO_4 is produced —4 Heated in bromine MnBr_2 is formed —5 When Mn is heated in a stream

of hydrogen chloride MnCl_2 is produced—6 Mn decomposes water readily, with evolution of H_2 . The reactions of the metal have not been much examined.

Combinations—Mn forms alloys with many metals, and compounds with most of the non metals, but few have been formed by the direct union of their elements.

Detection and estimation—Mn is usually detected by the colour its compounds impart to a borax-bead, and by the ppn of buff-coloured MnS by addition of NH_4 sulphide to an alkaline or neutral solution of Mn salts. Mn is usually estimated gravimetrically by ppn with NH_4Aq , heating the pp in air, and weighing as Mn_2O_3 . As MnO_2 is much used in technical chemistry it is necessary to have accurate and rapid methods for determining the quantity of this oxide in specimens of Mn ores, for descriptions of these methods, and also for other methods for estimating Mn, reference must be made to *Manuals of Analysis*.

Manganese, alloys of Several alloys of Mn have been prepared, none has been made by directly alloying Mn with other metals.

1 With *aluminum* Wöhler & Michel (*A* 115, 102) obtained a crystalline powder, having the composition MnAl_3 , by fusing together MnCl_2 , Al, and a mixture of NaCl and KCl.

2 With *copper* Alloys of Mn and Cu are obtained by reducing mixtures of MnO , and Cu with charcoal, an alloy containing 25 p.c. Mn is white, fairly hard, very elastic, and tolerably easily melted (Allen, *C* 22, 184). Alloys containing from 3 to 20 p.c. Mn resemble bronze (Valenciennes, *C* R 70, 607, v also Schrotter, *D P J* 210, 355, Prieger, *ibid* 177, 303, Gintl, *ibid* 224, 653).

3 With *iron* Alloys of Mn and Fe, containing from 8 to 80 p.c. Mn, are used in the manufacture of steel, under the names of *spiegeleisen* and *ferromanganese*. These alloys are prepared by heating MnO with iron filings and charcoal in the blast furnace, or in graphite crucibles, or by reducing a mixture of FeO and MnCO_3 on the hearth of a Siemens' furnace, and then fusing under a reducing flame.

4 With *mercury* An amalgam of Mn is obtained by reducing MnCl_2Aq by Na amalgam (Giles, *P M* [4] 24, 328), or also by electrolyzing MnCl_2Aq in contact with Hg (Ramsay, *C J* 55, 532, Moissan, *Bl* [2] 31, 149). Alloys of Mn with Pb, Sn, and Zn are described by Allen (*l c*) and Valenciennes (*l c*).

Manganese, arsenates of, v vol 1 p 309

Manganese, arsenide of Mn and As combine when heated together to redness. An arsenide, approximately As_3Mn_4 , occurs native (Kane, *P* 19, 145).

Manganese, arsenite of, v vol 1 p 306

Manganese, boride of By heating Mn_2O (v *Manganese, carbides of*) with B_2O_3 in a graphite crucible, Troost and Hautefeuille obtained small greyish violet crystals of MnB_2 (*C R* 81, 1263). This compound decomposes water at 100° and dissolves in acids with evolution of H_2 , it reacts with moist HgCl_2 to produce MnCl_2 , boric acid, and HCl.

Manganese, bromides of Only one bromide, MnBr_2 , has been isolated, the tetrabromide also perhaps exists in solution.

MANGANOUS BROMIDE, MnBr_2 , Manganese dibromide The hydrate $\text{MnBr}_2 \cdot 4\text{H}_2\text{O}$ is obtained as clear, red, deliquescent, crystals, by dissolving MnCO_3 in HBrAq , or by digesting Mn with Br (Balard, *J pr* 4, 178, Marignac, *Ann M* [5] 12, 7), crystallises in monoclinic forms, $a b c = 845 : 1 : 166$ (Marignac). The anhydrous salt MnBr_2 is obtained by heating powdered Mn in Br vapour, it forms a rose red deliquescent mass, heated in air it gives Br and Mn_2O_3 , Thomsen gives the thermal data— $[\text{Mn}, \text{Br}, \text{Aq}] = 106,120$ (*Th* 3, 271).

MANGANOUS BROMIDE, MnBr_2 , (Manganese tetrabromide) This compound perhaps exists in the green solution obtained by treating Mn_2O_3 or MnO , with HBr gas and dry ether, the solution is easily decomposed with formation of MnBr_2 (Nicklès, *C R* 60, 79).

Manganese, carbides of According to Brown (*J pr* 17, 492), the compound MnC is obtained by heating $\text{Mn}(\text{SCy})_2$ and MnC , by heating MnCy_2 , Troost and Hautefeuille obtained graphite-like, lustrous, crystals of MnC , by melting Mn in a charcoal crucible, and cooling slowly (*C R* 80, 960).

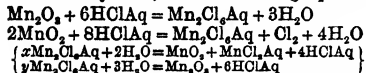
Manganese, chlorides of The only chloride of Mn which has been isolated is MnCl_2 . A solution of MnO_2 , or Mn_2O_3 , in conc. cold HClAq very probably contains Mn_2Cl_2 , perhaps also some MnCl_3 . The existence of MnCl_3 , described by Dumas, is very doubtful.

MANGANOUS CHLORIDE, MnCl_2 Mol w 125.74, Rose-coloured crystals of $\text{MnCl}_2 \cdot 4\text{H}_2\text{O}$ are obtained by dissolving MnCO_3 , or any oxide of Mn, in HClAq , and evaporating, by heating the dried crystals in a stream of dry HCl, the anhydrous salt, MnCl_2 , is obtained. MnCl_2 is also formed by heating Mn, MnCO_3 , or Mn_2O_3 , in a stream of dry HCl. By heating very finely powdered MnO with half its weight of NH_4Cl gradually to redness, MnCl_2 is formed. MnCl_2 is very deliquescent, Brandes (*P* 22, 263) gives S at $10^\circ = 62.16$, at $31.25^\circ = 85.72$, at $62.5^\circ = 122.22$, at $87.5^\circ = 122.22$, and at $106.25^\circ = 123.81$. S in alcohol at $11^\circ = 50$. SG of $\text{MnCl}_2 = 2.478$ (Schröder), of $\text{MnCl}_2 \cdot 4\text{H}_2\text{O} = 1.913$ (Schröder), 2.015 (Boedeker). The tetrahydrated salt is isomorphous with $\text{FeCl}_2 \cdot 4\text{H}_2\text{O}$, monoclinic, $a b c = 1.1409 : 1.16406$ (Marignac, *Ann M* [5] 15). All water is removed at 100° . Thomsen gives the thermal data $[\text{Mn}, \text{Cl}_2] = 111,990$, $[\text{MnCl}_2, \text{Aq}] = 16,010$, $[\text{MnCl}_2, 4\text{H}_2\text{O}] = 14,470$ (*Th* 3, 270). Heated in O_2 , a crystalline oxide containing 37 p.c. MnO_2 is produced (Schulze, *J pr* [2] 21, 407). MnCl_2 melts, in absence of air, at a red heat and sublimes at higher temperature. Scott found VD at c $1200^\circ - 1500^\circ$ to be 132.3 (*Pr E* 14, 410).

MnCl_2 forms double salts with the alkali chlorides, of the composition $\text{MnCl}_2 \cdot 2\text{XCl} \cdot 3\text{H}_2\text{O}$. The best examined are those in which $\text{X} = \text{NH}_4$, Rb, and Cs, the NH_4 salt contains one H_2O only according to Rammelsberg (*J pr* 65, 181, confirmed by Pickering, *C J* 85, 672). The double salts are obtained by mixing MnCl_2Aq or a solution of an oxide of Mn in HClAq with the alkali chloride, and evaporating slowly (v Godeffroy, *B* 8, 9, v Hauer, *J pr* 63, 436). Another double salt $\text{MnCl}_2 \cdot 3\text{CuO} \cdot 8\text{H}_2\text{O}$ is formed by boiling MnCl_2Aq with powdered CuO , filtering, and cooling (André, *C R* 106, 854). The double

salt $\text{MnCl}_2 \cdot \text{HgCl}_2 \cdot 4\text{H}_2\text{O}$, is described by Bonsdorff (*P* 17, 131)

MANGANIC CHLORIDE, Mn_2Cl_7 , and MANGANESE TETRACHLORIDE, MnCl_4 . Neither of these chlorides has been isolated. Mn_2O_3 and MnO_2 dissolve in cold conc HClAq to form deep-brown liquids, which slowly evolve Cl_2 and after a time contain MnCl_2 . Nicklès (*d Ch* [4] 5, 161), by passing HCl into ether in which MnO_2 was suspended, obtained a green liquid, of varying composition and very unstable, one analysis gave results approximately agreeing with the formula $\text{MnCl}_2 \cdot 12\text{C}_2\text{H}_5\text{O} \cdot 2\text{H}_2\text{O}$. These results are quite inconclusive of the formation of MnCl_2 (cf Pickering, *C J* 35, 672). Fisher's experiments (*C J* 38, 409) led him to conclude that a solution of MnO_2 in cold conc HClAq contains MnCl_2 , but the more complete experiments of Pickering (*C J* 35, 654) make it very probable that Mn_2Cl_7 , and not MnCl_4 , is produced when either Mn_2O_3 or MnO_2 is dissolved in cold conc HClAq . When the solutions are decomposed by adding water, the pp varies in composition but may always be expressed as $x\text{MnO}_2 \cdot y\text{MnO}$, x varying from 16 to 36 and y being usually 5. Pickering expresses the reactions of HClAq with Mn_2O_3 and MnO_2 , and the decomposition of the solutions by H_2O , in the following equations —



The average values of x and y are 4 and 1 respectively. Christensen (*J pr* [2] 35, 57) thinks that Mn_2Cl_7 is the product of the reaction of cold HClAq with MnO_2 , he supposes that some MnCl_2 is produced at 10° . According to C, ether holding HCl in solution produces a solution of MnCl_2 when shaken with Mn_2O_3 . According to Vernon (*C S Proc* 1890 58), a solution of MnO_2 in conc HClAq evolves less than half the Cl_2 at ordinary temperatures, required by the equations given by Pickering (*supra*), at -18°C is evolved very slowly, and at -26° only 35 p.c of the available Cl comes off when air is drawn through the solution for two hours. Vernon thinks that MnCl_2 is the only higher chloride formed by dissolving MnO_2 , Mn_2O_3 , or Mn_3O_4 in cold conc HClAq .

Franke (*J pr* [2] 36, 38) obtained *chloromanganic acid*, H_2MnCl_4 , by adding KMnO_4 to ether containing HCl , shaking with dry ether, and surrounding the deep-blue liquid thus produced with a freezing mixture.

MANGANESE HEPTACHLORIDE, MnCl_7 (?) Dumas (*B J* 7, 112, 8, 177) described a greenish gas, condensing at -15° to -20° to a green brown liquid, produced by adding excess of conc H_2SO_4 to KMnO_4 , and throwing in small pieces of fused KCl or NaCl , he gave the formula MnCl_7 , to this substance. Aschoff's analyses of the compound proved the presence of O in it, and led to the formula MnO_2Cl (*J pr* 81, 29). The exact composition of the substance is not yet settled.

Manganese, chromate of, v vol 11 p 155

Manganese, cyanides of, v vol 11 p 342

Manganese, ferri- and ferro cyanides of, v vol 11 pp 335, 339

Manganese, fluorides of. Two fluorides of Mn have been certainly isolated, MnF_3 and

Mn_2F_7 . The existence in solution of MnF_3 is doubtful. Wohler obtained a gas by the reaction of H_2SO_4 with a mixture of K_2MnO_4 and KF . To this gas he assigned the composition MnF_3 , but the composition of the substance cannot be regarded as settled. Nicklès asserts the existence of Mn_2F_7 .

MANGANESE FLUORIDE, MnF_3 (Manganese difluoride). A reddish crystalline powder, obtained by dissolving Mn_2CO_3 in excess of HFAq and evaporating, undecomposed by heating to redness (Berzelius).

MnF_3 forms double compounds with SiF_4 , &c (Berzelius, Stolba, *C C* 1883 292, Mari-gnac, *J pr* 83, 202). These compounds, better regarded as silicofluoride, titanofluoride, &c, of Mn , have the composition $\text{MnXF}_3 \cdot 6\text{H}_2\text{O}$, where $\text{X} = \text{Si}, \text{Ti}$, or Sn , they are isomorphous, crystallising in hexagonal forms, $a : c = 1 : 515$ (Mari-gnac, *Ann M* [5] 15). There is also a zirconofluoride of Mn , $\text{MnZnF}_3 \cdot 5\text{H}_2\text{O}$, which crystallises in monoclinic forms, $a : b : c = 2 : 09 : 1 : 2515$ (*M l c*).

MANGANESE SESQUIFLUORIDE, Mn_2F_7 . Crystals of $\text{Mn}_2\text{F}_7 \cdot 6\text{H}_2\text{O}$ were obtained by Christensen (*J pr* [2] 35, 57) by dissolving artificially prepared MnO_2 in HFAq , filtering through spongy Pt , evaporating, and placing over H_2SO_4 . Boiling or diluting the solution of MnO_2 in HFAq produces an oxyfluoride.

The double salts $\text{Mn}_2\text{F}_7 \cdot 4\text{KF} \cdot 2\text{H}_2\text{O}$, $\text{Mn}_2\text{F}_7 \cdot 4\text{NH}_4\text{F}$, $\text{MnF}_3 \cdot 4\text{NaF}$, and $\text{Mn}_2\text{F}_7 \cdot 2\text{AgF} \cdot 8\text{H}_2\text{O}$ are described by Christensen (*l c* and *ibid* p 161). They are obtained by adding solution of the alkali fluoride to solution of Mn_2O_3 or MnO_2 in HFAq , washing with water containing HF , and drying on Pt , the Ag -salt is obtained by dissolving freshly pptd AgCl in HFAq , adding Mn_2F_7 in HFAq , and evaporating. According to Christensen, the salt $\text{Mn}_2\text{F}_7 \cdot 4\text{KF} \cdot 2\text{H}_2\text{O}$ is identical with the compound to which Nicklès gave the formula $\text{MnF}_3 \cdot 2\text{KF}$ (*C R* 65, 107).

MANGANESE TETRA and HEPTA FLUORIDES, MnF_4 and Mn_2F_7 (?) The former compound was supposed by Nicklès to exist in the solution of MnO_2 in HFAq (*C R* 65, 107), by adding KF or NaF the double salts $\text{MnF}_3 \cdot 2\text{K}(\text{Na})\text{F}$ were said to be formed. The investigations of Christensen (*J pr* [2] 35, 57, 161) have made it very probable that Mn_2F_7 , and not MnF_4 , is formed when MnO_2 or Mn_2O_3 is dissolved in HFAq (*v supra*, cf *Manganese chloride, supra*).

The formula MnF_3 was given by Wohler (*P* 9, 619) to a purple yellow gas obtained by adding conc H_2SO_4 to a mixture of 2 parts commercial K_2MnO_4 and 1 part CaF_2 in a Pt retort. The gas dissolves in water to form HMnO_4Aq and HFAq , and on evaporation HF and O are evolved, and MnF_3 remains. The composition of the gas is still very doubtful, no analyses are given in Wohler's paper.

MANGANO MANGANIC FLUORIDE, Mn_2F_7 . According to Nicklès (*C R* 67, 448), brown crystals, having the composition $\text{Mn}_2\text{F}_7 \cdot 10\text{H}_2\text{O}$, are obtained by reacting on MnO_2 with warm HFAq and evaporating.

Manganese, haloid compounds of. The compounds MnX_2 ($\text{X} = \text{F}, \text{Cl}, \text{Br}, \text{I}$) have been isolated. Mn_2F_7 has also been obtained in definite form. There are very strong reasons in favour of the existence of Mn_2Cl_7 in the solutions obtained by dissolving Mn_2O_3 or MnO_2 in cold conc HClAq .

The existence even in solution, of tetra haloid compounds, MnX_4 , is doubtful. Nicklès claims to have obtained mangano manganic fluoride Mn_2F_7 , corresponding with Mn_2O_7 . The only haloid compound of Mn which has been gasified is $MnCl_4$. The general formula MnX_3 probably expresses the atomic composition of the molecules of the more stable haloid compounds of Mn, the formula MnX_2 may or may not be molecular. The existence of hepta-haloid compounds MnX_7 , might be expected from the position of Mn in the periodic scheme of classification, but the existence of these compounds is extremely doubtful.

Manganese, hydroxides of, v Manganese, oxides and hydrated oxides of, for $HMnO_3$, v Manganese, oxyacids of

Manganese, iodide of MnI_2 . Obtained, with $4H_2O$, in rose red deliquescent crystals, isomorphous with $MnCl_2 \cdot 4H_2O$, by dissolving $MnCO_3$ in $HIAq$, and evaporating. Turns brown when exposed to air and light, heated in absence of air, it is not decomposed, heated in O it burns like tinder, evolving vapours of I . Thomsen gives the thermal data $[Mn, I, Aq] = 75,700$ (*Th* 3, 271).

Manganese, oxides and hydrated oxides of. The oxides MnO , Mn_2O_3 , Mn_3O_4 , MnO_2 , probably also MnO_3 and Mn_2O_5 , have been isolated. A number of oxides intermediate between Mn_2O_3 and MnO_2 , having the general form $xMnO_2 \cdot yMnO$, also exist. Hydrates of most of these oxides exist, but their stability is generally small. $Mn_2O_3 \cdot H_2O = H_2Mn_2O_4$ is known, it is an acid. The acid corresponding to MnO_2 (H_2MnO_4) has not been isolated, but salts of this acid (manganates) are known. The oxides MnO , Mn_2O_3 , and Mn_3O_4 are basic. MnO reacts with acids to form manganous salts MnX_2 ($X = NO_3, \frac{1}{2}SO_4, \&c$), Mn_2O_3 forms manganic salts Mn_2X_3 , which are readily reduced to MnX_2 , Mn_3O_4 does not form corresponding salts, but with acids yields MnX_2 and MnX_3 , or in some cases MnX_2 and MnO_2 . Nicklès, however, asserts the production of Mn_2F_7 by the action of $HFAq$ on Mn_2O_3 (*v Mangano manganic fluoride*, p 180). MnO_2 reacts with acid to form manganic salts Mn_2X_3 , or manganous salts MnX_2 , according to temperature. It is possible that a few salts corresponding with MnO_2 may exist, but their isolation is doubtful. MnO_2 also combines with some oxides more basic than itself to form *manganites*, salts of the form $xMO \cdot yMnO$. MnO_2 is very unstable, it does not form salts. The existence of $(MnO_2)_2SO_4$ is probable. With water MnO_2 forms $H_2Mn_2O_4 \cdot Aq$ and MnO_2 . Mn_2O_3 is very unstable, with water it forms permanganic acid, $H_2Mn_2O_4$.

The molecular weight of none of the oxides of Mn is known with certainty, as none has been gasified.

MANGANOUS OXIDE, MnO (*Manganese monoxide or protoxide*). SG 5 09, crystalline (Ram-melsberg), 5 18, *manganosite* (Blomstrand, B 8, 180).

Occurrence—In small quantities, as *manganosite*, in bright green hexagonal forms (Blomstrand, l c).

Preparation—1 By heating to redness in a Pt vessel a mixture of equal parts $MnCl_2$ and Na_2CO_3 , with a little NH_4Cl , $MnCO_3$ is formed

and then decomposed, the formation of higher oxides is prevented by the NH_4Cl , the residue is washed and dried (Liebig, A. Wohler, P 21, 578). 2 By heating any of the higher oxides of Mn, very finely divided, to redness in a stream of H until the powder is green. According to Wright a Menke (C J 37, 28), pure MnO can be thus obtained, even from specimens of MnO_2 , containing c 10 p.c of potash—3 $MnCO_3$ or $Mn_2C_2O_4$ is strongly heated in absence of air, and the product is then heated in H (Liebig, A 95, 116).—4 Deville (C R 53, 199) obtained MnO in bright green regular octahedra by heating MnO_2 in H containing a very little HCl .

Properties and Reactions—A grass green powder, Deville's crystalline specimen (*v supra*) formed lustrous, diamond like, green regular octahedra. According to Moissan (A Ch [5] 21, 199, 251), MnO prepared by reduction of higher oxides by CO at 140° is pyrophoric. When pure, MnO does not oxidise by exposure to air (Wright a Menke, C J 37, 28 note), but if it contains minute quantities of potash oxidation occurs. Heated in air or O , Mn_2O_3 is produced, if the heating is done carefully till the weight is constant at dull redness, Mn_2O_3 is produced (*v Gorgeu, C R 106, 743*). MnO melts at white heat in absence of air. It is not reduced by heating in H or CO , or with C at 500° 600° (Wright a Luff, C J 33, 523). When heated in H_2S , MnS and H_2O are formed. MnO reacts with acids to form manganous salts, MnX_2 ($X = NO_3, ClO_3, \frac{1}{2}SO_4, \&c$).

HYDRATE OF MANGANOUS OXIDE, $MnO \cdot H_2O$. Occurs in small quantities in Sweden, in white crystalline tablets, as *pyrochroite*. Prepared, as small white hexagonal prisms, by adding 300 grm KOH in 500 c c air free water to an air free solution of 15-17 grm crystallised $MnCl$ in 15 c c air free water, in a vessel filled with H or coal gas, heating to 160° , and allowing to cool (A de Schulten, C R 105, 1265). Rapidly oxidises in air. When a manganous salt is present the compound $2MnO \cdot MnO_2 \cdot xH_2O$ is formed, when exposed to O for several years the product is $MnO_2 \cdot MnO$ (Gorgeu, C R 108, 948). When NH_4Aq is added to solution of a manganous salt, $MnO \cdot H_2O$ is not pptd, as it is soluble in NH_4Aq , but this solution rapidly absorbs O from the air, and after a time all the Mn is pptd as hydrates of Mn_2O_3 . The presence of NH_3 salts hinders the oxidation process, solutions of double NH_4 Mn salts are scarcely changed in air if free NH_3 is absent. Thomsen gives the thermal data $[Mn, O, H^+O] = 94,770$, $[MnO^+H^+O] = 21,560$, $[MnO^+H^+H^+SO^+Aq] = 26,480$ (*Th* 3, 271).

MANGANO MANGANIC OXIDE, Mn_2O_3 (*Red oxide of manganese*).

Occurrence—As *hausmannite*, in small brownish black tetragonal forms, a c = 1 1 1537, SG 4 8.

Preparation—1 Pure $MnCl_2 \cdot Aq$ is pptd by $Na_2CO_3 \cdot Aq$, the pp is thoroughly washed, dried, and then heated to whiteness for some time until the weight is constant (*cf* Wright a Luff, C J 33, 520, with Reissig, A 108 27).—2 By strongly heating $Mn_2C_2O_4$ in air (Laassaigne, A Ch [3] 40, 329).—3 Crystals of *hausmannite* were obtained by Debray by strongly heating a mixture of $MnSO_4$ and K_2SO_4 in a Pt crucible (C R 52, 985), also by passing a very slow stream of HCl

over amorphous Mn_2O_3 heated to redness (Dewille, *C R* 53, 199), also by keeping molten MnCl_2 in an atmosphere laden with moisture (Gorgeu, *C R* 96, 1044), also by melting amorphous Mn_2O_3 with borax (Nordenskjöld, *P* 114, 112, *v.* also Debray, *Ann M* [5] 1, 124, Sidot, *C R* 69, 201, *v.* Hauer, *J pr* 63, 425, Ebell, *D P J* 220, 64, 155)

It is generally stated that Mn_2O_3 is produced by heating any of the other oxides of Mn to redness in air, according to the experiments of Dittmar (*C J* 1864 294) the composition of the product of heating MnO_2 in a mixture of O and N varies according to the pressure of the O, if the pressure of the O is about 19 atmos the product is approximately Mn_2O_3 , while if the pressure of the O is greater than about 26 atmos the product approximates more or less closely to Mn_2O_3 . The experiments of Wright and Luff (*C J* 33, 520), on the effect of heating MnCO_3 in air, showed that unless the heating is continued for a long time and the temperature is kept very high the product contains more O than Mn_2O_3 . According to Gorgeu (*C R* 106, 743), finely powdered Mn_2O_3 can be oxidised to $x\text{MnO}_2 \cdot y\text{MnO}$, finally to Mn_2O_3 , by heating in air

Properties and Reactions—A reddish brown solid powder, crystalline Mn_2O_3 is brown SG crystalline 4856, amorphous 4918 Mn_2O_3 is not changed when heated to a very high temperature. It is reduced to MnO by H, CO, and O (Bell, *C N* 23, 258, Muller, *P* 186, 160), reduction by CO begins at c 100°, by H at c 240°, and by C at c 420° (Wright & Luff, *C J* 33, 520) Mn_2O_3 with conc H_2SO_4 forms a solution containing MnSO_4 and $\text{Mn}_2(\text{SO}_4)_3$, when the acid is hot, only MnSO_4 is formed and O is evolved. With boiling HNO_3 , $\text{Mn}(\text{NO}_3)_2$ and MnO are produced. Hot conc HClAq produces MnCl_2 and evolves Cl. Fused with alkalis, alkaline manganate is formed. Cl in presence of an alkaline solution reduces an alkaline permanganate. The reactions of Mn_2O_3 with acids, KIAq , &c., suggest the constitution $2\text{MnO} \cdot \text{MnO}_2$ (Pickering, *C J* 35, 657)

HYDRATES OF MANGANO-MANGANIO OXIDE, $\text{Mn}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ Such hydrates appear to exist, but their composition varies. Gorgeu (*C R* 84, 177) says a yellowish green hydrate is formed by shaking an aqueous solution of a manganese salt with alkali in presence of air. Hydrates of Mn_2O_3 are also said to be formed by placing finely-powdered MnO_2 in excess of an ammoniacal solution of MnCl_2 and heating (*cf J* Otto, *A* 93, 372) Veley (*C J* 41, 63) obtained a substance nearly agreeing in composition with the formula $8\text{Mn}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$, by heating $x\text{MnO}_2 \cdot y\text{MnO} \cdot z\text{H}_2\text{O}$ in H to 200°

MANGANIO OXIDE, Mn_2O_3 (*Sesquioxide of manganese*)

Occurrence—As *braunite* in quadrate octahedra, SG 4752 (Rammelsberg, *P* 124, 513) The hydrate $\text{Mn}_2\text{O}_3 \cdot \text{H}_2\text{O}$ occurs native as *manganite*

Preparation—1 The pp—approximately $\text{MnO}_2 \cdot x\text{H}_2\text{O}$ —obtained by passing Cl into $\text{Na}_2\text{CO}_3\text{Aq}$ holding finely powdered MnCO_3 in suspension, is made into a thin cream with conc H_2SO_4 and slowly heated on an oil bath to 100°, at which temperature O is suddenly evolved, and the mass becomes thicker and

greyish violet in colour, it is then heated to 138° until it is dark green. The impure $\text{Mn}_2(\text{SO}_4)_3$ thus produced is placed on a warm porous plate, by which H_2SO_4 is absorbed, it is then rubbed with conc HNO_3Aq , free from HNO_2 , again dried on a porous tile, and then warmed to 130°. The $\text{Mn}_2(\text{SO}_4)_3$ thus produced is exposed to air, when it rapidly deliquesces to form a violet solution, which afterwards becomes turbid from separation of $\text{Mn}_2\text{O}_3 \cdot \text{H}_2\text{O}$ ($\text{Mn}_2(\text{SO}_4)_3 + 4\text{H}_2\text{O} = \text{Mn}_2\text{O}_3 \cdot \text{H}_2\text{O} + 3\text{H}_2\text{SO}_4$) The brown solid which separates is washed, dried at 100° and then gently heated until the water is removed (Carius, *A* 98, 53)—2 According to Schreider (*P* 107, 605), Mn_2O_3 is obtained by heating MnO_2 , MnO , or Mn_2O_3 in O (but *v.* account of Dittmar's experiments under *Manganio manganio oxide*, *supra*) Moissan says that artificially prepared Mn_2O_3 goes to MnO when heated in O to 230° (*A Ch* [5] 21, 232). According to Berthelot (*A Ch* [5] 15, 185) and Knab, Mn_2O_3 is obtained by heating MnCl_2 , MnBr_2 , or MnI_2 in air or O. This oxide is also said to be formed, with evolution of O, by passing H_2O vapour over heated KMnO_4 , Gorgeu (*C R* 108, 1106) obtained MnO , by allowing ppd MnCO_3 to remain in contact with aerated water for 10 years, also by exposing $\text{MnO} \cdot \text{H}_2\text{O}$ to O in presence of excess of a manganous salt, also by exposing solutions of MnSO_4 , MnCl_2 , and $\text{Mn}(\text{C}_2\text{H}_3\text{O}_2)_2$ to sunlight

Properties and Reactions—A black powder, SG 4325, the mineral *braunite* forms brownish black, very hard, lustrous quadrate octahedra, SG 4752 Deoxidised at white heat to Mn_2O_3 . Soluble in conc H_2SO_4 , forming a reddish liquid, which evolves O on warming, and then contains MnSO_4 , boiled with dilute $\text{H}_2\text{SO}_4\text{Aq}$ or HNO_3Aq , MnO is separated, and manganous sulphate or nitrate goes into solution (Christensen, *J pr* [2] 23, 81) Soluble in cold conc HClAq , forming a brown liquid, which most probably contains Mn_2Cl_6 (*v. Manganio chloride*, p 180)

Mn_2O_3 is a basic oxide, the corresponding salts are not numerous, they are readily reduced to manganous salts. $\text{Mn}(\text{SO}_4)_2$ combines with alkali sulphates to form alums. Mn_2O_3 reacts with hot HClAq , with KIAq , and other reagents, as if it were $\text{MnO} \cdot \text{MnO}_2$ (Pickering, *C J* 35, 657) Laugier (*C R* 104, 1508) describes several compounds of Mn_2O_3 with SeO_2 , prepared by the reaction of MnO_2 with $\text{H}_2\text{SeO}_4\text{Aq}$

HYDRATE OF MANGANIO OXIDE, $\text{Mn}_2\text{O}_3 \cdot \text{H}_2\text{O}$ Occurs native as *manganite*, SG 4335, iso-morphous with *gibbsite* and *diaspore*, the corresponding Fe and Al compounds. The preparation of $\text{Mn}_2\text{O}_3 \cdot \text{H}_2\text{O}$ is described under *Manganio oxide* (*v. supra*), it forms a brownish black powder. Warmed with conc H_2SO_4 to c 100°, $\text{Mn}_2(\text{SO}_4)_3$ is formed without evolution of O (Carius, *A* 98, 53) According to Carius (*lc*), $\text{Mn}_2\text{O}_3 \cdot \text{H}_2\text{O}$ is not dissolved by dilute $\text{H}_2\text{SO}_4\text{Aq}$ even on gently warming, but if a little MnO is present solution occurs in the cold. $\text{Mn}_2\text{O}_3 \cdot \text{H}_2\text{O}$ is said by Hermann (*P* 74, 303) to dissolve in tartaric acid, forming a brownish red liquid, from which manganous tartrate separates on standing, the liquid becoming colourless and now containing formic acid and CO_2 . In a stream of H_2S , a little MnSO_4 , and also MnS

and Mn_2O_3 are formed (Wagner, *D P J* 195, 532) When moist $\text{Mn}_2\text{O}_3 \cdot \text{H}_2\text{O}$ is shaken with *magnesia alba*, K_2CO_3 , or Na_2CO_3 , or even with water, and much air, nitrates are produced according to Reichardt (*Henneberg's J für Landwirthsch* 26, 167)

MANGANESE PEROXIDE, MnO_2 (*Manganese dioxide*)

Occurrence—As *pyrolusite*, in iron black, opaque, rather brittle trimetric crystals, $a:b:c = 776:11066$, SG 4.82 to 4.97. The name is supposed to have been given from the use of the mineral to remove the colour from glass, coloured by compounds of iron (*pur* = fire, and *laver* = wash out)

Preparation—Pure MnCO_3 is dissolved in as small a quantity as possible of dilute HNO_3Aq , the solution is evaporated to a syrup, which is heated to $160^\circ\text{--}165^\circ$ for some hours, the product is thoroughly washed with boiling water, then dried over H_2SO_4 , and heated to $c\ 180^\circ\text{--}200^\circ$, until every trace of water is removed. Pure MnO_2 is thus obtained, exactly resembling *pyrolusite* (Gorgeu, *C R* 88, 796, Wright a Menke, *C J* 37, 45, cf. Schlösing, *C R* 55, 284, and Kuhlmann, *D P J* 211, 25). Even if the $\text{Mn}(\text{NO}_3)_2$ used contain a large quantity of KNO_3 , MnO_2 practically free from K_2O is obtained by this method.

Very many attempts have been made to obtain pure MnO_2 by ppn from Mn salts, e.g. by ppg with BrAq in presence of Na acetate, by passing Cl into an alkaline solution containing MnCO_3 in suspension, by reacting on a Mn salt solution with KMnO_4Aq , &c. The various methods have been examined by Gorgeu (*C R* 88, 796, *A Ch* [3] 66, 153), Guvard (*Bl* [2] 1, 89), Hannry (*C J* 33, 269, cf. Beilstein a Jawein, *B* 12 1530), Pickering (*C J* 35, 654), Vollhard (*A* 198, 318), Kesslet (*T* 18, part 1), Pattinson (*C J* 35, 365), Veley (*C J* 37, 581), Wright a Luff (*C J* 33, 504), and Wright a Menke (*C J* 37, 22). The outcome of the work is that pure MnO_2 cannot be obtained by any of the ppn methods, either the pp as $x\text{MnO} \cdot y\text{MnO}$, or, if all the Mn is present as MnO , the pp contains also K_2O or some other base besides H_2O . Vollhard's method—adding excess of KMnO_4Aq to MnSO_4Aq in presence of HNO_3 (exact quantities are given by V)—gave all the Mn as MnO accompanied by a small quantity of K_2O (c 3 p.c.), which could not be removed by washing (W a M, 1c).

Properties—A black, or brownish black, hard, crystalline powder. SG 5.02. Heated to moderate redness, Mn_2O_3 is formed, heated to whiteness, Mn_2O_3 remains (cf. *Mangano manganic oxide*, *Preparation*, p 182). MnO_2 is a conductor of electricity, it is strongly electro negative to the metals. Reacts with acids to form salts corresponding with MnO , with cold conc HClAq most probably forms MnCl_2 .

Reactions—1 Heated, gives Mn_2O_3 at moderate redness, and Mn_2O_4 at white heat—2 Heated in hydrogen, or carbon monoxide, is reduced to MnO , reduction in H begins at $c\ 190^\circ$, and in CO at $c\ 87^\circ$, when heated with carbon reduction begins at $c\ 390^\circ$ (Wright a Luff, *C J* 33, 518)—3 Heated with potassium chlorate, O is evolved and MnO_2 and KCl remain:

a little Cl is evolved, and at one stage of the change KMnO_4 is produced. The reaction between MnO_2 and KClO_3 probably produces KMnO_4 , Cl, and O, the KMnO_4 then decomposes to K_2MnO_4 , MnO_2 , and O, and the K_2MnO_4 reacts with Cl to form KCl , MnO_2 , and O. The following equations are given by McLeod as expressing approximately the various changes:

(1) $2\text{MnO}_2 + 2\text{KClO}_3 = 2\text{KMnO}_4 + \text{Cl}_2 + \text{O}_2$,
(2) $2\text{KMnO}_4 = \text{K}_2\text{MnO}_4 + \text{MnO}_2 + \text{O}_2$,
(3) $\text{K}_2\text{MnO}_4 + \text{Cl}_2 = 2\text{KCl} + \text{MnO}_2 + \text{O}_2$ (v McLeod, *C J* 55, 184, Hodgkinson a Lowndes, *C N* 59, 63, Fowler a Grant, *C J* 57, 272, Baudumont, *J Ph* [4] 14, 84, 161, Warren, *C N* 58, 247, Veley, *T* 1888 [1], 271, Spring a Prost, *Bl* 1889 340)—4 With hot conc sulphuric acid, MnSO_4 is formed and O evolved—5 With cold conc hydrochloric acid, MnCl_2 is very probably produced, on warming, Cl is evolved and MnCl_2 remains (cf. *Manganic chloride*, p 180)—6 Heated with sulphuric acid and oxidisable bodies, MnO acts as an oxidiser, e.g. $\text{H}_2\text{C}_2\text{O}_4$ is thus oxidised to CO_2 and H_2O —7 Not acted on by moderately dilute nitric acid alone, but in presence of an oxidisable body, $\text{Mn}(\text{NO}_3)_2$ and oxidised products are formed, e.g. with HNO_3 and HCl , Cl is evolved—8 Compounds of Mn O, with SeO_2 , were obtained by Laugier (*C R* 104, 1508), by heating ppd MnO_2 with selenious acid—9 With hydrogen peroxide and an acid, forms a salt of MnO and H_2O and evolves O, with hydrogen peroxide alone, O is evolved and the same quantity of MnO remains as was originally used (v Hydrogen peroxide, *Reactions* 11 and 19, vol II pp 723, 724)—10 Heated with magnesium chloride, MnCl_2 and MgO are formed and Cl is evolved—11 Fused with alkalis in presence of air or oxygen, manganates M^1MnO_4 , are formed (v Manganates under *Manganese*, oxyacids of, p 185).

12 MnO_2 combines with a number of basic oxides to form salts of the form $x\text{MO} \cdot y\text{MnO}_2$, known as *manganites*. According to Gorgeu (*A Ch* [3] 66, 153) MnO_2 decomposes neutral solutions of many salts of Ca, Ba, Ag, and Mn, making the solutions acid to litmus and combining with the bases. From dilute solutions of K_2CO_3 and Na_2CO_3 , MnO_2 withdraws and combines with 7 p.c. K_2O , and 4.1 p.c. Na_2O , respectively, according to Gorgeu, the *manganites* thus produced approximately correspond with the formula $\text{M}_2\text{O} \cdot 15\text{MnO}_2$. According to Jolles (*Chem. Zeitung*, 11, 1394) K_2MnO_4 is obtained, as a brown yellow solid, by mixing equivalent quantities of KMnO_4 and $\text{C}_2\text{H}_4\text{O}$. Wright and Menke (*C J* 37, 22) have examined the *potassium manganites* formed by ppg MnO_2 in presence of K salts, by passing Cl into $\text{K}_2\text{CO}_3\text{Aq}$ holding MnCO_3 in suspension, by adding MnSO_4Aq to KMnO_4Aq , &c., they conclude that the amount of K_2O ppd in combination with MnO_2 varies according to the relative masses of the reacting bodies, the temperature, the quantity of free acid present, &c., and that it tends to the maximum $5\text{M}_2\text{O} \cdot \text{K}_2\text{O}$. W a M failed to obtain $8\text{MnO}_2 \cdot \text{K}_2\text{O} \cdot 3\text{H}_2\text{O}$ described by Stigl a Morawski (*J pr* [2] 18, 91) as produced by acting on KMnO_4Aq with a reducing agent such as SO_2 , glycenn, or alcohol, they say that K_2O is ppd in combination with MnO_2 , but that the ratio of K_2O to MnO_2 varies much. Weldon (*C. N.* 20,

109) gave the formulae CaO MnO_2 and CaO 2MnO_2 to the *manganites* formed by oxidising $\text{MnO H}_2\text{O}$ by air in presence of lime. *Manganites* of the form MO 5MnO_2 , where $\text{M} = \text{Ca, Ba, Sr, Zn, and Pb}$, are described by Risler (*Bl* [2] 30, 110) as produced by heating KMnO_4 with slight excess of various metallic chlorides, and washing with water. By heating metallic chlorides with MnO_2 , and then adding MnCl_2 , Rousseau obtained *manganites*, which he classes as MO 5MnO_2 , MO MnO_2 , and 2MO MnO_2 (*C R* 101, 167, cf also Post, *B* 12, 1484, 1537, Rammelsberg, *B* 8, 233, Rousseau, *C R* 102, 425, 615, 103, 261, 104, 786, 1796). Various compounds of MnO with MnO_2 seem to exist, these may be regarded as *manganese manganites*, *v Oxydes intermediate between manganic oxide and manganese peroxide, infra*.

13 While MnO_2 reacts as an acidic oxide towards the oxides of the more positive metals, it also reacts as a basic oxide with some acids. The usual reactions of MnO_2 with acids are those of a basic peroxide (*v Reactions*, 3 to 6, *supra*). The salt MnO_2SO_4 is said by Fremy to be formed by carefully reacting on $\text{MnO}_2 \cdot x\text{H}_2\text{O}$ with conc H_2SO_4 , and allowing to stand in air (*C R* 82, 475). By adding MnSO_4 to a freshly prepared solution of PbO_2 in acetic acid, as long as PbSO_4 was pptd., Schönbein (*J pr* 74, 325) obtained a deep brownish red liquid, which had strongly oxidising properties, and from which MnO_2 separated on standing, or more quickly on boiling. Schönbein thought that this liquid contained an acetate of MnO_2 , but he did not succeed in isolating this salt.

HYDRATES OF MANGANESE PEROXIDE When approximately pure MnO_2 is pptd. by oxidising manganous salts in alkaline solutions, or by reactions between manganous salts and permanganates, or by decomposing KMnO_4aq by H SO_4 or HNO_3 , the pp always contains water, but it is very doubtful whether any definite, and fairly stable, hydrate of MnO_2 exists. When the process of formation results in ppn of all the Mn as MnO_2 , the composition of the pp approximates to $\text{MnO}_2 \cdot \text{H}_2\text{O}$, but this body loses water in dry air (*v* Wright & Menke, *C J* 37, 22). When the whole of the Mn is not pptd. as MnO_2 , the pp consists of compounds of the form $x\text{MnO}_2 \cdot y\text{MnO} \cdot z\text{H}_2\text{O}$ containing variable quantities of metallic oxides according to the conditions of formation. It appears to be possible to obtain hydrates which are stable for many hours within definite limits of temperature, but these hydrates are of the form $x\text{MnO}_2 \cdot y\text{MnO} \cdot z\text{H}_2\text{O}$ (*v* Velely, *C J* 37, 581, 41, 56).

References—The following papers contain the chief experiments bearing on the formation and composition of supposed hydrates of MnO_2 —Rammelsberg, *B* 8, 233, Fremy, *C R* 82, 1231, Van Bemmelen, *B* 13, 1466, Gorgeu, *A Ch* [3] 66, 154, *C R* 108, 948, Von Hauer, *W A B* 13, 453, Reising, *A* 103, 206, Böttcher, *J pr* 76, 235, Guyard, *Bl* 6, 81, Morawski a Stungl, *J pr* [2] 18, 90, 97, Volhard, *A* 193, 318, Suckow, *D P J* 177, 231, Wernicke, *P* 141, 116, Velely, *C J* 37, 581, 41, 56, Pickering, *C J* 35, 654, Wright & Luff, *C J* 33, 504, Wright & Menke, *C J* 37, 22, Franke, *J pr* [2] 86, 166, 451.

OXIDES INTERMEDIATE BETWEEN MANGANIC

OXIDE AND MANGANESE PEROXIDE These oxides belong to the general formula $x\text{MnO} \cdot y\text{MnO}_2$. The composition of the pp obtained by adding water to solutions of MnO , or Mn_2O_3 , in cold conc HClAq varies between $16\text{MnO} \cdot 5\text{MnO}_2$ and $36\text{MnO} \cdot 5\text{MnO}_2$ (Pickering, *C J* 35, 659). By passing Cl for a limited time into solution of Mn acetate, Velely obtained a pp approximating in composition to $5\text{MnO} \cdot \text{MnO} \cdot x\text{H}_2\text{O}$ (*C J* 37, 581, 41, 56, by heating this pp in a current of air, the compound $11\text{MnO} \cdot \text{MnO} \cdot \text{H}_2\text{O}$ was produced, and by heating the first compound in O , the body produced had the composition $23\text{MnO} \cdot \text{MnO} \cdot 2\text{H}_2\text{O}$. Wright & Menke (*C J* 37, 22) obtained a number of bodies $x\text{MnO}_2 \cdot y\text{MnO} \cdot z\text{H}_2\text{O}$ by various processes of oxidising manganous salts, and reducing permanganates, in every case, however, K O , or other alkali, was held in combination. Further accounts of these intermediate oxides, many of which were probably mixtures of the better defined oxides of Mn, will be found in the memoirs referred to under **HYDRATES OF MANGANESE PEROXIDE** (*v supra*), especially in the memoirs of Gorgeu.

MANGANFSE TRIOXIDE, Mn_2O_3 This compound is formed, in very small quantities, by slowly dropping a solution of KMnO_4 in cold conc H_2SO_4 (c f grams KMnO_4 in 100 cc H SO_4) on to dry Na_2CO_3 . The Na CO_3 is placed in a distilling flask surrounded by cold water. The flask is connected with a U tube filled with fragments of glass, and surrounded by a mixture of ice and salt, and this tube is connected with another U tube containing a little dilute $\text{H}_2\text{SO}_4\text{aq}$. As each drop of the green solution of KMnO_4 in H_2SO_4 falls on to the Na CO_3 , a pink cloud is formed, the cloud partly condenses in the first U tube, and part of it passes on and is dissolved in the H SO_4 in the second tube (Franke, *C J* [2] 36, 31, 166, *v* also Thorpe & Hambly, *C J* 53, 175). Only a very small quantity of MnO can be obtained. It appears as a reddish, amorphous, deliquescent mass. It slowly decomposes at ordinary temperatures, but is fairly stable if surrounded by ice and salt (*T a H*, l c). Mn_2O_3 is decomposed by water yielding HMnO_4aq and MnO_2 (*T a H*), according to Franke, H_2MnO_3 is produced, but quickly decomposes to MnO_2 , O , and HMnO_4 , and probably also $\text{H Mn}_2\text{O}_4$. Mn_2O_3 dissolves in conc H_2SO_4 , forming a green solution. Franke thinks this liquid contains $(\text{MnO}_2)_2\text{SO}_4$. Mn_2O_3 dissolves in KOH Aq , forming K_2MnO_4 . Mn_2O_3 liberates I from KI , and acts on Hg similarly to ozone (*T a H*).

MANGANESE HYPOXIDE, Mn_3O_4 , (*Permanganic anhydride*) This oxide was obtained by Thenard (*C R* 42, 382). Its composition was determined by Aschoff (*J pr* 81, 34). Pure KMnO_4 , free from Cl compounds, is added little by little to conc H SO_4 , *S G* 1845, kept cold by a freezing mixture. To the green solution thus obtained a few drops of water are added, when Mn_2O_3 slowly separates in dark redash-brown oily drops. If acid of the composition $\text{H}_2\text{SO}_4 \cdot \text{H}_2\text{O}$ (*S G* c 178) is used, oily drops of Mn_2O_3 are formed without addition of water. According to Franke (*J pr* [2] 86, 81), a solution of KMnO_4 in conc H_2SO_4 contains $(\text{MnO}_2)_2\text{SO}_4$, and this is decomposed by a little water, giving Mn_2O_3 and H_2SO_4 .

Terreil (*Bl* 1862 40) prepares Mn_2O_3 by dissolving $KMnO_4$ in well cooled H_2SO_4 , containing water in the ratio $H_2SO_4 : \frac{1}{2}H_2O$, placed in a stoppered retort, the neck of which passes into a glass balloon surrounded by a freezing mixture. Corks or organic material must not be used in any part of the apparatus. He heats the retort to $60^\circ-65^\circ$ (not over 70°), when purple red vapour is evolved and condensed to a thick greenish black liquid, which is Mn_2O_3 . Only a few drops of Mn_2O_3 can thus be prepared at a time. As soon as a certain quantity collects in the balloon, decomposition occurs, with slight detonation (v also Spiess, *J pr* [2] 1, 421).

P. Thenard (*J pr* 69, 58) describes Mn_2O_3 as a dark olive green liquid, with a smell recalling Cl compounds and ozone. Aschoff (*J pr* 81, 34) describes it as dark brownish red oily drops, which do not solidify at -20° . Mn_2O_3 slowly decomposes in air at ordinary temperatures, with evolution of O. It may be heated to $60^\circ-65^\circ$, under reduced pressure, without volatilising, at a somewhat higher temperature it is suddenly and violently decomposed to MnO_2 and O. Mn_2O_3 is very hygroscopic. It dissolves in water to form a purple liquid, probably containing $HMnO_3$. This liquid is slowly decomposed on warming into MnO_2 and O. Mn_2O_3 dissolves in cold conc H_2SO_4 , apparently with out decomposition, to form an olive green liquid (this liquid contains $(MnO_3)_2SO_4$ according to Franke, *J pr* [2] 36, 31). Mn_2O_3 is at once decomposed by contact with small quantities of MnO_2 , Ag O, or HgO (Aschoff, *l c*). Mn_2O_3 is a powerful oxidiser. It inflames paper or alcohol.

MANGANESE TETROXIDE (?) Franke (*J pr* [2] 36, 31, 166) states that a blue gas is obtained by leading air or CO_2 , saturated with water at $40^\circ-50^\circ$, over the green liquid formed by dissolving $KMnO_4$ in well cooled conc H_2SO_4 . To this blue gas he gives the formula MnO . Thorpe and Hamble repeated Franke's experiments (*C J* 53, 178), but failed to obtain any indications of the formation of a blue gas.

Manganese, oxyacids of, and their salts. Two series of salts derived from oxyacids of Mn are known—the *manganates* Mn_2MnO_4 and the *permanganates* $MnMnO_4$. The acid corresponding to the manganates, viz H_2MnO_4 , has not been isolated, but its anhydride MnO_3 is known. The anhydride of permanganic acid, viz Mn_2O_7 , is known, and the acid itself has also probably been isolated.

MANGANATES, M_2MnO_4 , derived from the hypothetical acid H_2MnO_4 . The anhydride of this acid, viz MnO_3 , is known. A solution of this oxide in water perhaps contains H_2MnO_4 , but it decomposes almost at once to $HMnO_3$ and MnO_2 (v *Manganese trioxide*, p 184). When an acid is added to solution of a manganate, the manganic acid produced at once decomposes to permanganic acid and MnO_2 . This change occurs even when CO_2 is passed into solution of a manganate. The manganates are isomorphous with the sulphates.

The fact that the product of fusing together pyrolusite, potash, and saltpetre dissolved in water to form a green liquid, which became blue, violet, and then red, on addition of much water, was known to Scheele. To the green substance Scheele gave the name '*Chameleon minerale*.'

Chevillot and Edwards, in 1817, showed that the green substance was a definite compound of potash with an acid of Mn (*A Ch* [2] 4, 287, 8, 337). Forchhammer (*Annals of Phil* 16, 310, 17, 150) and Fromherz (*P* 31, 677) investigated the manganates. Mitscherlich, in 1830, showed that two salts are obtained by the action of alkali on pyrolusite in presence of air or alkali nitrate (*P* 25, 287).

Manganates of the alkalis and alkaline earths are obtained by heating MnO_2 with KOH, $CaO.H_2O$, &c, to c 150° in absence of air, or to higher temperatures in presence of air, or by strongly heating any Mn salt with KOH, $CaO.H_2O$, &c, in presence of O or an oxidiser, e.g. $KClO_3$, K_2MnO_4 , and Na_2MnO_4 dissolve with out decomposition in water containing alkali. In pure water, $KMnO_4$ or $NaMnO_4$ is formed and MnO_2 ppt. Solutions of the alkali manganates are decolourised, with p.pn of MnO_2 , by easily oxidised bodies, e.g. SO_2 , As O_3 , H_2SAq , or ferrous salts. Ba and Sr manganates are in soluble in water.

Barium manganate $BaMnO_4$. A dark green powder. SG 4.85. Insol water. Unchanged in air, decomposed by acids. Prepared by calcining $Ba(NO_3)_2$ with MnO_2 , or by projecting finely powdered MnO_2 into a molten mixture of $KClO_3$ and $BaO.H_2O$, washing with hot water, and drying. By heating MnO_2 with $Ba(NO_3)_2$, $BaMnO_4$ is obtained as a green powder consisting of minute hexagonal crystals (Forchhammer, *Annals of Phil* 16, 130, 17, 150, Rosenstiehl, *J Ph* 46, 544, Schafarik, *J pr* 90, 16). The salt is also obtained by digesting $Ba(MnO_4)_2$ with $BaO.Aq$.

Didymium manganate $Dy_2(MnO_4)_3$. A black powder, insol water. Obtained by heating for 30 minutes 1 pt MnO_2 with 4 pts Dy_2NO_3 , and washing with water (Frerichs & Smith, *A* 191, 353).

Lanthanum manganate $La_2(MnO_4)_3$. Resembles the Dy salt, prepared similarly (*F a, S l c*).

Potassium manganate $KMnO_4$.

Formation—1 A mixture of equal parts of finely powdered MnO_2 and KOH is heated to bright redness in air, or in O, $3MnO_2 + 2KOH = K_2MnO_4 + Mn_2O_3 + H_2O$. $KMnO_4$ is formed by heating MnO_2 and KOH in absence of O to c 150° (Beketoff, *Bl* 1, 43), the reaction occurs in N at c 180° (Elliot & Storer, *P Am A* 5, 192). If the mixture is heated above 180° O must be present, because at c 190° K_2MnO_4 is decomposed with reformation of MnO_2 .—2 By boiling conc $KMnO_4$ with KOH.Aq; $2KMnO_4.Aq + 2KOH.Aq = 2K_2MnO_4.Aq + H_2O + O$ (Aschoff, *J pr* 81, 29). According to Thenard (*J pr* 69, 58), this reaction only occurs when the KOH contains some oxidisable substances, e.g. a little organic matter.—3 By long continued heating $KMnO_4$ to 240° , $2KMnO_4 = K_2MnO_4 + MnO_2 + O_2$ (Thenard, *l c*).—4 By fusing any oxide of Mn with KOH in presence of O, or an oxidiser such as $KClO_3$.

Preparation—2 pts KOH are dissolved in the smallest quantity of water, 1 pt $KClO_3$, and 2 pts very finely powdered MnO_2 are added, the mixture is dried, and then heated, nearly to redness, for a long time in a Pt dish, the fused mass when cold is treated with a little water,

the conc green solution is decanted, after standing until quite clear, and evaporated *in vacuo* over H_2SO_4 .

Properties and Reactions—Dark green crystals, isomorphous with K_2SO_4 and K_2CrO_4 . Soluble in water containing KOH without decomposition, dissolves in pure water with decomposition to $KMnO_4$ and MnO_2 . Dilute acids, even CO_2 , quickly decompose K_2MnO_4 , forming K permanganate and a manganous salt, e.g. with H_2SO_4 ,— $5K_2MnO_4 \cdot Aq + 4H_2SO_4 \cdot Aq = 2K_2MnO_4 \cdot Aq + MnSO_4 \cdot Aq + 3K_2SO_4 \cdot Aq + 4H_2O$. Heated in CS_2 , MnS , CO_2 , and polysulphides of K are formed (Muller, *P* 127, 404). Heated in a stream of water vapour, Mn_2O_3 , KOH, and O are produced.

$(2K_2MnO_4 + 2H_2O = Mn_2O_3 + 4KOH + 3O)$ By fusing MnO_2 with KOH in air, and then passing water vapour over the product, then again fusing the products together, and repeating these processes, O may be obtained from the atmosphere.

Sodium manganate $Na_2MnO_4 \cdot 10H_2O$. Obtained by long continued heating equal pts finely powdered MnO_2 and $NaNO_3$, boiling with water, filtering, and allowing to cool (Gentele, *J* pr 82, 58). Said to form colourless crystals, resembling Glauber's salt, and to dissolve in water with partial decomposition to form a green solution.

PERMANGANIC ACID AND PERMANGANATES The acid $HMnO_4$, or $H_2Mn_2O_8$, has probably been isolated, a series of salts derived from this acid is known. Permanganic anhydride, Mn_2O_7 , is a very unstable liquid, it dissolves in water to form $HMnO_4 \cdot Aq$, which slowly decomposes to MnO_2 and O. Permanganic acid and its salts readily part with O, and hence act as energetic oxidisers. The permanganates are isomorphous with the perchlorates $M'ClO_4$, they are purple red, all are soluble in water, the least soluble is the Ag salt. These salts are produced by the action of dilute acids, or of Cl, on the manganates, or by the reaction of several oxidisers, e.g. Pb_2O_3 and dilute $HNO_3 \cdot Aq$, on Mn salts—(Tamm, *J* 1872 910, Fresenius, *Fr* 11, 415, 425, Crum, *A* 60, 219, Gibbs, *Am S* [2] 14, 204, Chatard, *C N* 24, 196, Pichard, *C R* 75, 1821, Deshayes, *Bl* [2] 29, 541, Volhard, *A* 198, 354, Rose, *P* 105, 289). $KMnO_4$ is reduced to K_2MnO_4 by boiling conc $KMnO_4 \cdot Aq$ with potash (cf. POTASSIUM MANGANATE, *Formation* No 2, p 185). Solutions of permanganates are decomposed by $NH_3 \cdot Aq$ with separation of $xMnO_2 \cdot yMnO$, they are reduced to manganous salts by $HNO_3 \cdot Aq$, fairly conc $HCl \cdot Aq$, $SO_2 \cdot Aq$, $Na_2S_2O_4 \cdot Aq$, many organic compounds, &c. H_2SO_4 decomposes solid permanganates with separation of MnO_2 and evolution of O. Solutions of $HMnO_4$ and $KMnO_4$ absorb the green and yellow green rays of the spectrum, the absorption spectrum has been examined by Lecoq de Boisbaudran (*Spectres lumineux*, p 108). According to Vogel (*B* 8, 1534) $\frac{1}{10000}$ pt of Mn may be detected, in presence of Cu and Fe compounds, by boiling with P_2O_5 or Pb_2O_3 and dilute $HNO_3 \cdot Aq$ free from Cl compounds, and examining the absorption spectrum of the solution (of permanganate) produced.

Ammonium permanganate NH_4MnO_4 . Resembles $KMnO_4$, with which it is isomorphous, v sol water, decomposed by gently

heating. Prepared by rubbing together $AgMnO_4$ and $NH_4Cl \cdot Aq$ in the ratio $AgMnO_4$, NH_4Cl , filtering, and evaporating at the ordinary temperature over H_2SO_4 , also by decomposing $BaMnO_4$ by $(NH_4)_2SO_4$ (Böttger, *N R P* 25, 115).

Barium permanganate $BaMnO_4$. Small, hard, rhombic octahedra, almost black, with violet sheen. Prepared by adding excess of $H_2SiF_6 \cdot Aq$ to $KMnO_4 \cdot Aq$, filtering from $KSiF_6$ through asbestos, saturating with $BaO \cdot Aq$ containing $BaO_2 \cdot H_2$ in suspension ($BaCO_3$ should not be used as it causes ppm of MnO_2 , xH O), decanting from $BaSiF_6$, and evaporating (Rousseau a Brunesau, *C R* 93, 229). Böttger (*N R P* 25, 115) prepares $BaMnO_4$ by dissolving in water the product of the fusion of 2 pts KOH and 1 pt MnO_2 , filtering, and adding $BaCl_2 \cdot Aq$ until the green colour of the liquid disappears, he collects the violet blue pp, and washes it with cold water till the washings begin to appear reddish, he heats this pp, which is $Ba(MnO_4)_2$, with water, passes in CO_2 till the liquid is purple red, filters from MnO_2 , and evaporates.

Calcium permanganate $CaMnO_4 \cdot 5H_2O$. A crystalline, deliquescent mass, prepared by decomposing $AgMnO_4$ by $CaCl_2 \cdot Aq$, following directions given for obtaining NH_4MnO_4 (*v supra*).

Cobalt permanganate does not seem to have been isolated, but Klobb has prepared several luteo cobaltic permanganates, viz $Co_2(NH_4)_{12}(MnO_4)_6$, $Co(NH_4)_{12}Cl_4(MnO_4)_4$, $Co(NH_4)_{12}Br_4(MnO_4)_4$ (*C R* 103, 384, *Bl* [2] 48, 240).

Potassium permanganate $KMnO_4$.

Formation—1 By fusing together KOH and an oxide of Mn in the air or with $KClO_3$, dissolving the K_2MnO_4 in water, passing in CO_2 , filtering from MnO_2 , and evaporating.

$(3K_2MnO_4 \cdot Aq + 2CO_2 = 2KMnO_4 \cdot Aq + 2K_2CO_3 \cdot Aq + MnO_2)$ —2 By warming solution of a Mn salt with $KClO_3$ and $KClO_4 \cdot Aq$ —3 By melting MnO_2 with KHO , making a conc solution of the fused mass, adding saturated $MgSO_4 \cdot Aq$, and filtering.

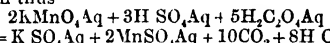
$(3K_2MnO_4 \cdot Aq + 2MgSO_4 \cdot Aq = 2KMnO_4 \cdot Aq + 2K_2SO_4 \cdot Aq + 2MgO + MnO_2)$ (Tessie du Mothay, *D P J* 186, 231)—4 By passing Cl into a solution of the fused mass obtained as in 1 and 3, and evaporating (Stadeler, *J* pr 103, 107). $(2KMnO_4 \cdot Aq + Cl_2 = 2KCl \cdot Aq + 2KMnO_4 \cdot Aq)$

Preparation—A mixture of 2 pts KOH and 1 pt $KClO_3$ is melted in a thin sheet iron crucible, the crucible is removed from the lamp, and 2 pts very finely powdered and sifted MnO_2 are added, little by little, heating is then continued, with stirring, until the mass gets quite hard, the temperature is then raised for a short time to full redness, when cold, the mass is broken up, and boiled with so much water that no crystals separate on cooling (c 40 pts water for 1 pt $KClO_3$ used), a rapid stream of well washed CO_2 is passed into the boiling liquid until a drop placed on paper makes a red stain (showing no green at the edges), the outer part of which soon becomes brown. After cooling the greater part of the liquid is poured off, the rest is filtered (from MnO) through glass wool, and the whole is evaporated until a drop placed on a cold surface quickly deposits crystals.

After standing 12 to 24 hours the greater part of the KMnO_4 separates out, a little is obtained by further evaporating the mother liquor 100 grms pyrolusite give about 30 grms KMnO_4 (Böttger, *J pr* 90, 156, for modification of this method, v Grager, *J pr* 96, 169, Wöhler, *P* 27, 626, Gregory, *J Ph* 21, 312, Mitscherlich, *P* 25, 287, Elliot a Storer, *P Am* 4, 5, 192) Béchamp (*A Ch* [3] 57, 293) heats MnO_2 with KOH , and oxidises by passing O over the fused mass, he then dissolves in water, passes in CO_2 , and evaporates

Properties—Dark purple red, almost black, rhombic crystals, $a, b, c = 79523 \text{ 1 6478}$ SG 2.71 (Kopp) Isomorphous with KClO_4 , S at $15^\circ = 6.25$ (Mitscherlich) Decomposed by heat, giving K_2MnO_4 and O Pure conc KMnO_4Aq can be boiled without change, in presence of oxidisable bodies KMnO_4Aq is quickly reduced

Reactions—1 Heated to 240° , MnO_2 , K_2MnO_4 , and O are produced (Chevillat a. Edwards, *A Ch* [2] 4, 290) At higher temperatures αMnO yk O is formed (Rousseau, *C R* 104, 786)—2 Sulphuric or nitric acid separates HMnO_4 , which decomposes with evolution of O and ppn of MnO_2 , Mn_2O_3 , or αMnO , γMnO , with considerable excess of warm $\text{H}_2\text{SO}_4\text{Aq}$ or HNO_3Aq , MnSO_4 or $\text{Mn(NO}_3)_2$ is produced Cold conc H_2SO_4 dissolves KMnO_4 , forming a green liquid, from which a few drops of water cause separation of MnO , (cf *Manganese heptoxide*, p 181, v also *Manganese trioxide*, p 184)—3 Hydrochloric acid in excess forms MnCl_2 , and evolves Cl Dry HCl evolves Cl , and forms H_2O , HCl , Mn_2O_3 , $\alpha\text{H}_2\text{O}$, MnCl_2 , and probably Mn_2Cl_2 (Thomas, *C J* 33, 372)—4 Sulphuric acid and oxidisable bodies either separate MnO , or αMnO , γMnO , the oxidisable body being simultaneously oxidised, or if there be enough acid to prevent ppn of αMnO_2 , γMnO , MnSO_4 is formed With $\text{H}_2\text{C}_2\text{O}_4$ and H_2SO_4 , CO_2 and H_2O (besides K_2SO_4 and MnSO_4) are produced (cf Berthelot, *A Ch* [5] 21, 176, Harcourt, *C J* [2] 5, 460, Jones, *C J* 33, 95) Harcourt (*lc*) represents the reaction thus



Jones (*lc*) says that H_2O , K_2CO_3 , and MnCO_3 are at first produced on adding KMnO_4Aq to $\text{H}_2\text{C}_2\text{O}_4\text{Aq}$, and that further addition of KMnO_4Aq forms K_2CO_3 , pppts Mn_2O_3 , and evolves CO_2 and O Alcohol gives Acetate and a pp containing MnO_2 , MnO , and K_2O , glycerin gives a similar pp and forms K_2CO_3 (Morawski a Stingl, *J pr* [2] 18, 78, cf Berthelot, *lc*)—5 According to Aschoff (*J pr* 81, 29) conc KMnO_4Aq is reduced to K_2MnO_4 by boiling with potash solution ($2\text{KMnO}_4\text{Aq} + 2\text{KOH}\text{Aq} = 2\text{K}_2\text{MnO}_4\text{Aq} + \text{H}_2\text{O} + \text{O}$), Thenard (*J pr* 69, 58) says that this action does not occur unless traces of oxidisable substances, e.g. organic matter, are present—6 KMnO_4Aq acidified by H_2SO_4 reacts with hydrogen peroxide to form H_2O , O , and MnSO_4 At low temperatures, c 12° , the KMnO_4 is decolourised, but O is not evolved, according to P Thenard (*C R* 75, 177), Berthelot (*A Ch* [5] 21, 176) thinks that H_2O_2 (?) is formed (v also Swietrowski, *A* 141, 205, Aschoff, *P* 111, 217)—7 KMnO_4Aq is decomposed with separation of MnO_2 , Mn_2O_3 , or αMnO , γMnO , by ammonia (Clodé a Guignet,

C R 47, 710, Wöhler, *A* 136, 256, Wanklyn a Gamgee, *C J* [2] 6, 25, Tamm, *C N* 25, 47), by hydrogen (Jones, *C J* 33, 96, Wanklyn a Cooper, *P M* [5] 6, 288), by phosphine, arsine, and stibine (Jones, *lc*, Schobig, *J pr* [2] 14, 289, Parsons, *C N* 1877 236), by chlorine dioxide (ClO_2) (Furst, *A* 206, 75)—8 With potassium sulphocyanide, K_2SO_4 and KCN are produced (Morawski a Stingl, *J pr* [2] 18, 78)—9 Sodium thiosulphate in boiling solution is wholly oxidised to Na_2SO_4 by fairly conc KMnO_4Aq , if the KMnO_4Aq is very dilute from 1 to 2 p.c. $\text{Na}_2\text{S}_2\text{O}_4$ remains (Glaser, *M* 7, 651)

Combination—From conc mixed solutions of KMnO_4 and K_2MnO_4 , the salt $\text{K}'\text{MnO}$, K_2MnO , separates in small, six sided, monoclinic tablets (Gorgeu, *J pr* 80, 123)

Sodium permanganate $\text{NaMnO}_4 \cdot 3\text{H}_2\text{O}$ Formed similarly to KMnO_4 , or by the reaction between AgMnO_4 and NaClAq V sol water, and hence obtained in crystals with difficulty

Silver permanganate AgMnO_4 Monoclinic crystals S 91 in cold water Obtained by mixing conc warm KMnO_4Aq with AgNO_3Aq , and allowing to cool (Dewar a Scott, *P* 35, 44) Klobb (*C R* 103, 384) obtained $\text{AgMnO}_4 \cdot 2\text{H}_2\text{O}$, by saturating KMnO_4Aq with NH_3 , and then adding an equivalent quantity of AgNO_3

Permanganates of copper, didymium (Freichs a Smith, *A* 191, 354), **lanthanum** (F a S, *lc*), **lead**, **lithium**, **magnesium**, **strontium**, and **zinc**, have been isolated

Permanganic acid, HMnO_4 The solution obtained by decomposing powdered $\text{Ba(MnO}_4)_2$ by an exactly equivalent quantity of $\text{H}_2\text{SO}_4\text{Aq}$, contains this acid Hunefeld (*Schweigger's Jahrb der Chem und Phys* 30, 133) says that the acid can be obtained, in indigo like crystals, by washing $\text{Ba(MnO}_4)_2$ with hot water, whereby it is decomposed to MnO and BaMnO_4 , filtering, adding exactly enough phosphoric acid to decompose the Ba salt, heating to 60° – 100° , filtering, and evaporating at a low temperature (No analyses are given)

Manganese, oxychlorides of Several oxy chlorides of Mn appear to exist, but they have not been satisfactorily examined P de Saint Gilles (*C R* 55, 329) says that $\text{MnCl}_2 \cdot 3\text{MnO}_2$ is formed as a black powder, resembling MnO_2 , by heating to 280° , in a partially closed vessel a mixture of MnCl_2 and NaNO_3 Gorgeu (*A Ch* [6] 4, 515) obtained an oxychloride, to which he provisionally assigned the formula MnCl MnO by heating MnCl_2 in water vapour According to Aschoff (*J pr* 81, 29) the gas which Dumas obtained by adding pieces of fused KCl or NaCl to a mixture of KMnO_4 and conc H_2SO_4 , and which he said was MnCl_2 , is probably an oxy chloride having the composition MnOCl

Manganese oxyfluoride of According to Nicklès (*C R* 65, 107) the compounds MnOF KF and $\text{Mn}_2\text{OF}_2 \cdot 2\text{HF}$ are ppd, when solution of MnO_2 in ethereal HCl is poured into boiling KFAq or NaFAq Christensen (*J pr* [2] 30, 57), however, asserts that the double compound obtained as described above is MnOF_2

Manganese, oxysulphide of The compound MnO MnS is said to be formed, by Arfvedson (*P* 1, 50), by heating MnSO_4 in H_2 , it is a green solid, which dissolves in acids with evolution of H_2S , heated in air it burns to Mn_2O_3

Manganese, phosphides of Several compounds of Mn with P have been described. Mn_3P an amorphous powder, S G 494, obtained by heating Mn and P together to low redness (Schrötter, *J pr* 51, 386). Also formed, according to H. Rose, by heating $MnCl_2$ in PH_3 . By heating Mn pyrophosphate mixed with sugar-charcoal, in a crucible filled up with charcoal, Struve (*J pr* 81, 321) obtained a brittle substance resembling pig iron, the composition of which varied between Mn_3P_2 and Mn_2P . Merkel and Wöhler (*A* 86, 371) obtained a crystalline, greyish, regulus, S G 595, by heating together 10 parts MnO_2 , 10 parts well burnt bones, 5 parts quartz sand, and 3 parts lamp black. The composition agreed with the formula Mn_3P_2 , but as part was soluble, and part insoluble, in $HClAq$, they regarded the substance as a mixture of two phosphides, Mn_3P_2 (soluble in $HClAq$) and Mn_2P_2 (insoluble in $HClAq$).

Manganese, salts of Compounds obtained by replacing the H of acids by Mn. Mn forms two series of salts: *manganous salts*, MnX_2 , and *manganic salts*, Mn_2X_3 , $X = NO_3$, ClO_3 , $\frac{1}{2}SO_3$, $\frac{1}{2}PO_3$, &c. One or two salts of the form MnX ,—e.g. $Mn(SO_4)_2$ —are also said to exist. The manganous salts are considerably more stable than the manganic salts. A number of double salts of manganic sulphate are known, those with the alkali sulphates are alums, e.g. $Mn_2(SO_4)_3 \cdot K_2SO_4 \cdot 24H_2O$. The manganous salts form a fair number of double salts and a few basic salts. The chief salts of Mn are the following: *antimonates*, *arsenates* and *arsenites*, *borate*, *carbonates*, *chlorate* and *perchlorate*, *chromates*, *ferric* and *ferro cyanides*, *iodate*, *molybdate*, *nitrate* and *nitrites*, *phosphates* and *phosphites*, *selenates* and *selenites*, *sulfates*, *sulphates* and *sulphites*, *thiosulphate*, *tungstate*, *vanadate* v **CARBONATES**, **NITRATES**, **SULPHATES**, &c.

Manganese, selenide of Said to be produced in combination with H_2O as a red powder, decomposed in air, by adding an alkali selenide solution to solution of a manganous salt.

Manganese, silicides of Mn and Si combine when heated together (Troost & Hautefeuille, *C R* 81, 264). Silicides of Mn, containing from 6.5 to 13 p.c. Si, have been obtained by Wohler (*A* 106, 54) by fusing together MnF_3 , Na, K silicate, and cryolite, or $MnCl_2$, $2NaCl$, CaF_2 , and K silicate, or $MnCl_2$, CaF_2 , K_2SiF_6 , and Na. These silicides form hard, brittle masses, they dissolve in $HClAq$ with evolution of H and SiH_4 .

Manganese, silicofluoride of. $MnSiF_6 \cdot 6H_2O$. Hexagonal crystals, $a = 1.515$, S G at $17.5^\circ = 1.9038$ (Stolba, *C C* 1893 292). Formed by dissolving $MnCO_3$ or MnF_2 in H_2SiF_6Aq , and evaporating. When heated, the crystals give off water, and then SiF_4 , and leave MnF_2 (Berzelius).

Manganese, sulphides of Two sulphides are known, MnS and Mn_2S_3 .

Manganous sulphide, MnS Occurs native, as *manganese blende*, in hexagonal crystals, S G 895. Produced by heating together MnO or $MnCO_3$ and S, not by heating S with Mr. Also formed by heating Mn in CS_2 (Gautier & Hallopeau, *C R* 108, 806), also by heating various Mn compounds in H_2S (Carnot, *Bl* [2] 32, 162). Sidot (*J* 1868 229) obtained MnS in hexagonal crystals by heating the amorphous substance in a stream of H_2S . MnS is pptd from solutions of

Mn salts by NH_4S in PH_3 , as a pale buff coloured amorphous solid, which quickly oxidises in air, if this ppt is rapidly dried and then warmed in H_2S , pure MnS is obtained. MnS is a greenish solid, decomposed by weak acids, e.g. acetic acid, with evolution of H_2S . Heated in air, SO_2 is evolved and Mn_2O_3 remains (P. W. Hofmann, *D P J* 181, 364). Heated in H_2O vapour, H_2S and H are given off and Mn_2O_3 is formed. MnS is slowly decomposed by Cl with formation of $MnCl_2$ and S_2Cl_2 .

There appear to be two forms of MnS obtainable by pptn from Mn salts by NH_4S , sulphide the ppt from conc. $MnCl_2Aq$ or $MnSO_4Aq$ is rose coloured, if NH_4Cl is present the ppt is greenish and consists of small 8 sided plates. The green sulphide is not produced when K_2S or Na_2S is used as pptant (Fresenius, *J pr* 82, 265; Muck, *Z* 1869 580; de Clermont & Guyot, *Bl* 27, 353; Geuther, *Z* 1865 347). The rose coloured sulphide is said to be changed to the green sulphide by heating with a little water, but it may be heated to 305° with much water without changing. The change from rose to green is accomplished by heating with NH_4Aq to 250° in a closed tube, and the reverse change by heating in NH_3 gas. The rose sulphide is not changed by heating alone to 250° but in presence of H S the green sulphide is formed at 220° (De C & G, *l c*). The green sulphide is thought by Muck to be an oxysulphide of Mn.

Double compound, $3MnS \cdot K_2S$ Obtained by heating a mixture of 1 pt dry $MnSO_4$, $\frac{1}{4}$ pt lamp-black, and 3 pts K_2CO_3 and S, and treating the fused mass with water, the compound remains insoluble in water, forming small lustrous dark red tablets (Voelcker).

MANGANIC SULPHIDE, Mn_2S_3 Occurs native as *hauertite*, in large, brown black, lustrous, regular octahedra, S G 3463. Said to be obtained as a red amorphous powder, by heating $MnSO_4Aq$ with solution of K polysulphides to 160° – 180° in a closed tube (Senarmont, *J pr* 51, 385). Not changed in air, decomposed by acids.

Manganese, sulphocyanide of, v vol 11 p 850 M M P M

MANGANITES Salts in which MnO_2 acts as the acidic radicle, v *Manganese peroxide*, *Reactions* 12, under **MANGANESE**, *Oxides and hydrated oxides of*, p 183.

MANGANOCYANIDES v vol 11 p 342

MANGOSTIN $C_{20}H_{22}O_5$ [α 190°] Contained in the husk of the fruit of *Garcinia mangostana*. The dry husks are boiled with water to extract tannin, then treated with hot alcohol, and the alcoholic extract left to evaporate. The mangostin which is deposited is dissolved in alcohol and pptd by lead subacetate. The ppt is decomposed by water and the mangostin finally crystallised from dilute alcohol (Schmid, *A* 93, 853). Thin golden laminae without taste or smell. Insoluble in water, v sol alcohol and ether. Warm dilute acids dissolve it without alteration. Hot conc. HNO_3 gives oxalic acid. Alkalis dissolve it with yellowish brown colour. Reduces chloride of gold solution. $FeCl_3$ gives a dark greenish black solution, decolourised by acids. Its solution is not pptd by any metallic salt except lead subacetate— $(C_{20}H_{22}O_5) \cdot 5PbO$ (dried at 100°), pptd by adding alcoholic lead acetate and ammonia to an alcoholic solution of mangostin.

MANNITE $C_6H_{12}O_6$

$CH_2(OH)CH(OH)CH(OH)CH(OH)CH(OH)CH_2(OH)$
 Mol w 182 (181 by Raoult's method, Brown a. Morris, *C J* 53, 620) [165°] SG 15 $[a]_D = -25$ S 15.6 at 18° (Berthelot, *A Ch* [3] 47, 801), 13 at 14° (Krusemann, *B* 9, 1467), 16 at 16.5° (Wanklyn a. Erlenmeyer, *J* 1862 480) S (alcohol) 07 at 14° HC v 728,200 HC p 728,500 (Berthelot a. Vieille, *Bl* [2] 47, 868, *A Ch* [6] 10, 456) HF 318,500 (B a V), 287,000 (Von Rechenberg)

Occurrence—Discovered by Proust (*A Ch* [1] 57, 143) Occurs to the extent of 30 to 60 p.c. in manna, the dried juice which exudes from the manna ash (*Fraxinus Ornus*) Mannite occurs in many other plants, e.g. the roots of *Aconitum napellus*, celery, *Apium graveolens*, *Meum athamanticum*, *Evanthe crocata*, *Polypodium vulgare*, *Scorzonera hispanica*, and *Triticum repens*, and in the root bark of *Tunica granatum* Mannite also occurs in the bark of *Cinnella alba* (8 p.c.), and of *Fraxinus excelsior*, in the leaves and young twigs of *Syringa vulgaris*, in the leaves of *Ligustrum vulgare* and of *Cocos nucifera*, and in the fruit of *Laurus Persca* and of *Cactus opuntia* Mannite also occurs in *Laminaria saccharina*, in olives, and in several fungi, e.g. *Lactarius vellereus*, *L. tupsis*, *L. pyrogalus*, and *L. pallidus* *Agaricus integer* contains 20 p.c. of its dry substance It also occurs in the cambium layer of *Conifera* (Payen, *A* 12, 60, Meyer a. Reiche, *A* 47, 234, Stenhouse, *A* 51, 349, Knop a. Schmedemann, *A* 49, 293, Dopping a. Schlossberger, *A* 52, 117, Muntz, *C R* 76, 619, 82, 210, Smith *J* 1850, 335, Roussin, *J* 1851, 550, Ludwig, *J* 1857, 503, De Luca, *J* 1861, 740, 1862, 505, Thorner, *B* 12, 1635, Reinsch, *J* 1863, 612, Bouquelot, *C R* 108, 568, Kiehler, *M* 7, 410)

Formation—1 In the lactic fermentation of sugar (Liebig, *J* 1847, 466, Pasteur, *J* 1857, 511, Diagenдорff, *Ar Ph* [3] 15, 47)—2 In the viscous fermentation of sugar, 100 pts of sugar yielding 51 pts mannite and 46 pts gum (Pasteur, *J* 1861, 728)—3 In the spontaneous fermentation of the juice of the sugar cane in tropical climates (Maicano, *C R* 108, 955)—4 By reducing glucose, levulose, or invert sugar with sodium amalgam (Linnemann, *A* 123 136, Dewar, *P M* [4] 39, 345, Bouchardat, *Bl* [2] 16, 38, Krusemann *B* 9, 1465, Scheibler, *B* 16, 3010)—5 By reducing mannose (L. Fischer, *B* 21, 1808), or 'gluco-one' (L. Fischer, *B* 22, 94), with sodium amalgam—6 From the dilactone of meta saccharic acid by treating with water and sodium amalgam (3 p.c.), acidifying with H_2SO_4 (Kilian, *B* 20, 2714, v. LÉVO-MANNITE, *infra*)

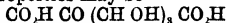
Preparation—1 Manna is extracted with boiling dilute alcohol, and the crystals which separate on cooling are recrystallised from water—2 Manna (2 pts) is boiled with water (1 pt) after addition of a little white of egg The crystals which separate from the filtrate are boiled with water (6 pts) to which some animal charcoal has been added, and the filtrate is allowed to crystallise (Ruspin, *A* 65, 203)

Properties—Needles or four sided prisms ∇ sol water, ∇ al sol alcohol, insol ether An aqueous solution, does not become syrupy on

spontaneous evaporation (difference from sugar) Only slightly sweet to the taste It has little, if any, action on light, but if borax be added to the solution it becomes dextrorotatory Thus in a 10 p.c. solution containing 12.8 g borax $\alpha = +22.5^\circ$, so that $[\alpha]_D = c + 225^\circ$ (Vignon, *A Ch* [5] 2, 493, *C R* 77, 1191, Muntz a. Aubin, *C R* 83, 1213, Pasteur, *C R* 77, 1192, Bouchardat, *C R* 80, 120, 84, 34) Arsenic acid slowly develops levorotation in a solution of mannite Salts of the alkalis and alkaline earths develop dextrorotation Free alkalis render the solution levorotatory A solution containing 8 g mannite and 8 g NaOH in 100 c.c. gives $\alpha = -3.4^\circ$, so that $[\alpha]_D = -42^\circ$ A solution of 12 g mannite and 4 g sodium tungstate made up to 100 c.c. shows $\alpha = +0.40'$ (Klein, *C R* 89, 484) These rotations vary with the amount of dissolved substances The mannite may be recovered from these solutions without having undergone any change in optical properties Mannite renders a borax solution acid, forming boric acid and sodium metaborate (Dunstan, *Ph* [3] 13, 257, 14, 41, Lambert, *C R* 108, 1016) Mannite does not reduce Fehling's solution It hinders the ppn of ferric and cupric salts by potash It is not turned brown by boiling KOH aq With a solution of ammonio sulphate of copper it gives a blue pp, sol in ammonia, forming a blue solution which is not affected by boiling (Guignet, *C R* 109, 528, 645) Mannite does not reduce boiling solutions of silver or mercurous nitrates, of $HgCl_2$, or of chloride of gold, but it reduces AgO and silver acetate (Hirzel, *A* 131, 50) When alcohol is added to a solution of mannite mixed with lime, strontia, or baryta water, pps $(C_6H_{12}O_6)_2CaO$, $(C_6H_{12}O_6)_2SrO$, and $(C_6H_{12}O_6)_2BaO$ are formed (Hirzel, *A* 131, 50, cf Ubaldini, *A Ch* [3] 57, 213) Ammoniacal lead acetate gives a pp On adding to an aqueous solution of mannite (1 mol) and of lead nitrate (2 mols) sufficient ammonia to neutralise two thirds of the nitric acid present in the salt, a voluminous white pp separates, which, if filtered off and dried over calcium chloride, forms a white crystalline powder $C_6H_{12}O_6.Pb(NO_3)_2.2H_2O$, S 110 at 14° , decomposing with explosion on heating, and separated into its constituents on long boiling with water and by carbonic anhydride with formation of lead carbonate (Smolka, *M* 6, 198)

Reactions—1 Mannite begins to sublime at c 200°, in a sealed tube it is unaltered at 250° At 280° it splits up into water and mannitan $C_6H_8O_6$ When distilled *in vacuo* it yields isomannosan $C_6H_{10}O_6$ —2 A mixture of HNO_3 and H_2SO_4 forms the nitrate, so called 'nitro mannite' (Flores Domonte a. Menard, *J* 1847, 1145)—3 Nitric acid oxidises it to saccharic, racemic, and oxalic acids, no mucic acid being formed (Backhaus, *J* 1860, 522, Carlet, *J* 1861, 367)—4 Boiling HIAq gives sec-hexyl iodide (Wanklyn a. Erlenmeyer, Domac, *M* 2, 309, cf Le Bel a. Wassermann, *C R* 100, 1589)—5 Boiling conc HCl aq slowly forms mannitan—6 Potash fusion yields formic, acetic, and propionic and oxalic acids and acetone (Gottlieb, *A* 52, 122)—7 On oxidation with $KMnO_4$ in alkaline solution it gives oxalic acid, formic acid, a little tartaric acid, a sugar which reduces Fehling's solution, and erythric acid $C_6H_8O_6$, which forms

the salts CaA' , 2aq, BaA' , 2aq, and $\text{C}_2\text{H}_5\text{BaO}$, 2aq (Hecht a Iwig, *B* 14, 1760, 19, 468, cf Pabst, *C* R 91, 728, who states that he obtained a dibasic acid $\text{C}_2\text{H}_2\text{O}_5$, which he names dioxyisoelectric acid) — 8 *As* and *platinum black* yield mannitic acid $\text{C}_2\text{H}_2\text{O}_5$, and mannitose $\text{C}_2\text{H}_2\text{O}_5$ — 9 MnO_2 and H_2SO_4 give formic acid and acrolein (Baokhaus) — 10 When mannite (2 pts) is heated with 80 p.c. formic acid (5 pts) for 8 hours a mixture of formyl derivatives of mannitan appears to be formed. This mixture yields on distillation CO , CO_2 , isomannide $\text{C}_2\text{H}_2\text{O}_5$, a liquid $\text{C}_2\text{H}_2\text{O}_5$ (?) (157° at 17 mm) and a liquid $\text{C}_2\text{H}_2\text{O}$ (108°), *S* G $\frac{1}{2}$ 9396, $[\alpha]_D = -168^\circ$, almost insol water, and smelling like a carbamine and mustard oil (Fauconnier, *C* R 100, 914, cf Henninger, *B* 7, 264) — 11 Distillation with *oxalic acid* yields formic acid, a formyl derivative of mannite or of mannitan being probably first formed (Lorin, *Bl* (2) 24, 437) — 12 *Electrolysis* of a solution of mannite acidulated by H_2SO_4 yields hydrogen, CO_2 , CO , oxygen, tri oxy methylene (? formic paraldehyde or mannitose), formic acid, oxalic acid, and a small quantity of an acid whose calcium salt is $\text{C}_2\text{H}_2\text{CaO}_2$, 2aq. This acid having reducing properties may be



(Rénard, *A Ch* [5] 17, 316) — 13 PCl_5 gives tetra chloro-hexamine (Bell, *B* 12, 1273) — 14 PI_2 forms methylene iodide (Butlerow, *A* 111, 247) — 15 *Organic acids* (e.g. acetic and butyric) heated with mannite at 200° form alkyl derivatives of mannitan — 16 Mannite heated with *water* suffers no change below 280°, but at that temperature a viscid product is got containing 'mannitic ether' $\text{C}_2\text{H}_2\text{O}_5$, $[\alpha]_D = -5.6$ and 'mannitone', a crystalline isomeride of mannitan. Both bodies are reconverted into mannite by heating with water at 295° (Vignon). — 17 Mannite does not react when heated with acetic, valeric, or benzoic aldehydes (Lochert, *A Ch* [6] 16, 65). But when *acetic aldehyde* is passed into a solution of mannite in H_2SO_4 or HClAq the aldehyde is absorbed and there is formed a compound crystallising in needles [174°], insol cold water *S* 1 at 95°. It is sl sol cold, v sol hot, alcohol. Boiling dilute (2 p.c.) H_2SO_4 resolves it into aldehyde and mannite. Paraldehyde gives the same compound. When a mixture of acetic and benzoic aldehydes is used the above product is formed first (Meunier, *C* R 103, 408) — 18 When a mixture of mannite (10 g), absolute alcohol (20 g), and fused ZnCl_2 (5 g) is saturated with HCl , left for two days, and then mixed with benzoic aldehyde, there is formed 'tribenzoic mannitoid' $\text{C}_2\text{H}_2\text{O}_5(\text{C}_6\text{H}_5\text{O})_3$, [207°]. This crystallises from benzene, and is insol water, sl sol alcohol. It is not affected by boiling alcoholic KOH , but dilute H_2SO_4 yields mannite and benzoic aldehyde (Meunier, *C* R 106, 1425, 1732). A solution of mannite in HClAq or H_2SO_4 also reacts with benzoic aldehyde (Meunier, *C* R 107, 910) — 19 A solution of mannite in alcohol containing ZnCl_2 and saturated with HCl reacts with BaCl_2 forming di-benzoyl mannide $\text{C}_2\text{H}_2\text{O}_5(\text{OBz})_2$, [152°] (Meunier, *C* R 107, 346) — 20 *Valeric aldehyde*, ZnCl_2 and HCl convert mannite in alcoholic solution into $\text{C}_2\text{H}_2\text{O}_5(\text{C}_4\text{H}_9\text{O})_2$, which is deposited as silky needles [91°] (Meunier) — 21 Not fermented by yeast. In contact with water, chalk, and cheese at 40° it gives

alcohol, carbonic, lactic, butyric, and acetic acids, and hydrogen, but no sugar or glycerin (Berthelot, *J* 1856, 664). When the fermentation is induced by a slit testicle a fermentible sugar is produced. Upon fermentation by Schizomycetes in presence of CaCO_3 and ammonium phosphate there is formed alcohol, *n* butyl alcohol, butyric, lactic, acetic, hexoic, and succinic acids (Fitz, *B* 10, 281, 11, 43). *Bacillus butyricus* gives similar products. Under other conditions formic acid is among the products of fermentation (Fitz, *B* 11, 1895, 15, 876, 16, 845) — 22 Heating with phenyl cyanate forms $\text{C}_2\text{H}_2(\text{OH})(\text{O CONHPh})$, [260°] (Tessmer, *B* 18, 968).

Pentanitrate $\text{C}_2\text{H}_2(\text{ONO})_2\text{O}$ *Pentanitroxyl derivative* [79°] *S* 2 at 60° *S* (alcohol, *S* G $\frac{1}{2}$ 150 at 12.8° *S* (ether) 130 at 9° *S*). Formed, together with nitro mannitan, by passing NH_3 into an ethereal solution of the hexanitrate (Tichanovitch, *J* 1864, 582). Long needles. Dextrorotatory. Explodes when struck, and also, more feebly, when heated. Ammonium sulphide reduces it to mannite.

Hexanitrate $\text{C}_2\text{H}_2(\text{ONO})_2\text{O}$ *Nitro mannite* Mol w 452 [113°] (Socloff) *S* G $\frac{1}{2}$ 1604 *S* (alcohol of *S* G $\frac{1}{2}$ 81) 3 at 12.8° *S* (ether) 5 at 9° (Tichanovitch) $\alpha = +12.4$ in a 5 p.c. solution, so that $[\alpha]_D = +400^\circ$ (Muntz a Aubin). Prepared by treating mannite (1 pt) with HNO_3 (5 pts of *S* G 1.5) at 0°, and adding H_2SO_4 (10 pts). The product is collected on a filter, washed with water and aqueous Na_2CO_3 , and recrystallised from alcohol (Domont a Menard, *J* 1847, 1145, Sobrero, *A* 64, 397, Strecker, *A* 73, 62, Socloff, *J* R 11, 136). Beautiful white needles, insol water, sol alcohol and ether. Explodes violently on being struck — it also explodes when suddenly heated, but if carefully heated it may be decomposed without explosion. Ammonium sulphide converts it into mannite (Dessaignes, *A* 81, 251). Iron and acetic acid and HIAq also convert it into mannite (Béchamp, *A Ch* [3] 46, 354, Mill, *J* 1864, 584). NH_3 passed into its ethereal solution forms crystalline $\text{C}_2\text{H}_2\text{O}(\text{NH}_3)_2$ (Tichanovitch).

Di sulphuric acid $\text{C}_2\text{H}_2(\text{SO}_3\text{H})(\text{OH})_2$. From mannite and conc H_2SO_4 (Favie, *B* 25, 560). The free acid is decomposed by boiling water. It gives no pp with BaCl_2 or CaCl_2 — $\text{Pb}_2\text{C}_2\text{H}_2\text{O}_5\text{O}_{12} \cdot 2\text{PbO}$ insoluble pp.

Tri sulphuric acid $\text{C}_2\text{H}_2(\text{SO}_3\text{H})_3(\text{OH})$. From mannite and conc H_2SO_4 (Knop a Schnedermann, *A* 51, 136). Both the acid and its salts are decomposed by water into mannite and H_2SO_4 — $\text{Na}_3\text{A}'''$ — $\text{K}_3\text{A}'''$ deliquescent gummy mass, insol alcohol — $\text{Ba}_3\text{A}'''$ crystal line powder, sol water, insol alcohol — $\text{Pb}_3\text{A}'''$ amorphous deliquescent mass.

Tetra sulphuric acid $\text{C}_2\text{H}_2(\text{SO}_3\text{H})_2(\text{OH})_2$ $[\alpha]_D = +9^\circ$. Obtained when, in the preparation of the hexa sulphuric acid, the neutralisation with BaCO_3 is delayed for two days — Ba_4A^{IV} .

Hexa sulphuric acid $\text{C}_2\text{H}_2(\text{SO}_3\text{H})_6$. $[\alpha]_D = +24^\circ$. Formed by adding mannite in small portions to chloro sulphuric acid (ClSO_3H). The product is dropped upon ice, and the ice cold solution neutralised with BaCO_3 — Ba_6A^{VI} 5aq. When alcohol is added to its solution this salt is thrown down as an oil, which

presently changes to a crystalline mass insol water. The other salts are very soluble, and do not behave thus (Claesson, *J pr* [2] 20, 10).

Boric acid When mannite (4 pts) is heated with boric acid (3 pts) for eight hours at 145°, and the product is dissolved in water and neutralised by BaCO₃, a filtrate is obtained which deposits barium borate on evaporating, after which alcohol gives a pp which, when dried at 100°, has the composition (C₆H₇B₂O₈) Ba (Klein, *Bl* [2] 29, 363).

Hexa acetyl derivative C₆H₇(OAc)₆ [119°] [α]_D = +18° (Bouchardat, *C R* 84, 34). From mannite (18 pts) and Ac₂O (80 pts) at 180°, with or without addition of a little ZnCl₂ (Bouchardat, *A Ch* [5] 6, 107, Schutzenberger, *A* 160, 94, Franchimont, *B* 12 2059). Limit of etherification from mannite (1 mol) and HOAc (6 mols) 26.4 (Menschutkin, *B* 13, 1814). Trimetric crystals (from HOAc). Insol water, cold alcohol, and ether, sl sol hot alcohol. Dextrorotatory. May be sublimed in a current of CO.

Penta benzoyl derivative C₆H₇(OBz)₅(OH) [c 80°]. From mannite (3 g), water (15 g), BzCl (20 g), and NaOHAq. The product is extracted with ether (Skraup, *M* 10, 391). Amorphous mass.

Hexa benzoyl derivative C₆H₇(OBz)₆ [149°]. From the preceding and BzCl. Crystalline grains, v sl sol alcohol.

Mannite dichlorhydrin C₆H₇(OH)₂Cl₂. *Di chloro tetra oxy heane* [171°] S 4.5 at 14° [α]_D = -3.7. Formed by heating mannite (1 pt) with conc HClAq (6 pts) at 100° for eight hours (Bouchardat, *C R* 75, 1187, 76 1550, *Bl* [2] 19, 199). Formed also by heating isomannide with fuming HClAq at 100° in sealed tubes (Fauconnier, *Bl* [2] 41, 119). Prepared by heating mannite (1 pt) with fuming HClAq (10 pts) at 106°, evaporating at low temperatures, dissolving the crystals in water, filtering the solution through animal charcoal, and recrystallising from hot alcohol (Siruloboff, *A* 233, 368).

Properties—Monoclinic crystals. Lavoatory. Its aqueous solution is neutral and tasteless, and is not pptd by AgNO₃. Insol alcohol and ether. Decomposed on fusion.

Reactions—1 Boiling water or conc NaOHAq rapidly convert it into C₆H₇O(OH)₂Cl₂.—2 A mixture of HNO₃ and H₂SO₄ gives C₆H₇(ONO₂)₂Cl₂, which crystallises from hot alcohol in needles [145°], insol water.—3 *Sodium amalgam* removes HCl in two stages, forming mannitan chlorhydrin, and finally (β) mannide [119°].

Mannite dibromhydrin C₆H₇(OH)₂Br [178°]. Obtained by heating mannite with conc HBrAq for 2 hours at 100° (Bouchardat, *A Ch* [5] 6, 120). Small colourless plates (from hot water). Decomposed on fusion. Insol cold water, alcohol, and ether. Converted by a mixture of HNO₃ and H₂SO₄ into C₆H₇(ONO₂)₂Br₂, which crystallises in long needles, sol hot alcohol, insol water.

Mannitan C₆H₇O₂, i.e. C₆H₇O(OH)₂. *First anhydride of mannite*. According to Alechin (*J R* 16, 888) the various mannitans described below are mixtures of isomannide and other bodies.

(a) *Berthelot's mannitan* C₆H₇O₂. Obtained in small quantity by heating mannite at 200°. A better method is by protracted boiling of mannite with conc HClAq (Berthelot, *A Ch* [3] 47, 306). Slightly sweetish syrup. Differs from mannite in being soluble in absolute alcohol. V sol water, insol ether. Slightly dextrorotatory (Bouchardat, *A Ch* [5] 6, 102). When exposed to the atmosphere it is partially reconverted into mannite. Boiling alkalis or dilute acids accelerate the change. Above 140° it partly volatilises.

(b) *Vignon's mannitan* C₆H₇O₂ [α]_D = +36.5°. Prepared by heating mannite for an hour or two with half its weight of water at 295°. Also by heating mannite (2 pts) with conc H₂SO₄ (1 pt) at 120°, saturating with BaCO₃, exhausting with alcohol, and evaporating (Vignon, *A Ch* [5] 2, 433). Deliquescent mass, v sol water and absolute alcohol, insol ether. Does not ferment with yeast. Boiling dilute H₂SO₄ or baryta water does not reconvert it into mannite. A mixture of mannitan (1 pt), H₂SO₄ (10 pts), and HNO₃ (4½ pts of SG 1.5) forms in the cold C₆H₇O(OH)(ONO₂), which is obtained on pouring into water and extracting with ether. It is dextrorotatory [α]_D = +53°.

(c) *Vignon's mannitone* C₆H₇O₂ [α]_D = -25°. Obtained by heating mannite with water for 3 hours at 280° and extracting the viscid product with alcohol (Vignon, *A Ch* [5] 2, 433). Crystals (from alcohol). Has a sweet taste. Levorotatory. Does not reduce Fehling's solution. In the preparation of this body the syrupy mother liquor yields a viscid mass, which appears to be the anhydride of mannite or 'mannitic ether' C₆H₇O₂, it does not reduce Fehling's solution, and is levorotatory, [α]_D = -5.6°. This mannitic ether is not converted into mannite or mannitan by boiling dilute acids or alkalis.

(d) *Crystalline mannitan* C₆H₇O₂ S 25 at 15°. When Berthelot's mannitan is allowed to stand for some months in a dry atmosphere it deposits a solid variety (Bouchardat). This crystallises in monoclinic tables. It is strongly levorotatory. Sl sol cold alcohol. Boiling water quickly converts it into mannite. This variety of mannitan is probably identical with Vignon's.

Mannitan tetra nitrate C₆H₇O(ONO₂)₄. Precipitated by adding water to the alcoholic mother liquor from which mannite pentanitrate has separated (Tichanovitch, *J* 1864, 583). Syrup, v sol alcohol and ether, insol water. Explodes when struck. Alcoholic KOH converts it slowly into syrupy mannitan.

Di acetyl derivative C₆H₇O(OH)₂(OAc)₂ [α]_D = +22.6. From mannite and HOAc at 210° (Berthelot). Mannite dissolves in boiling Ac₂O, and on cooling crystals of C₆H₇AcO₂ separate. Further action of Ac₂O yields hexa acetyl mannite and di acetyl mannitan (Schutzenberger, *A* 160, 74, Grange, *C R* 68, 1826). Extremely bitter substance, v sol water, HOAc, and alcohol. Dextrorotatory. Boiling baryta water converts it into acetic acid and mannitan.

Tetra acetyl derivative C₆H₇O(OAc)₄. Formed, together with hexa acetyl mannite, by heating mannite with Ac₂O at 180° (Bouchardat, *A Ch* [5] 6, 110). Amorphous viscid mass, which partially crystallises on long standing.

Insol water, v e. sol alcohol, ether, and HOAc. Dextrorotatory. Aqueous alkalis at 100° decompose it into acetic acid and mannitan. Ac_2O forms hexa-acetyl mannite. A mixture of HNO_3 and H_2SO_4 has no action in the cold.

Di-butyryl derivative

$\text{C}_6\text{H}_5\text{O}(\text{OC}_4\text{H}_7\text{O}_2)_2(\text{OH})_2$. From mannite and butyric acid at 200° (Berthelot, *C R* 38, 673, *A Ch* [3] 47, 319). Semi solid, partly crystalline, mass. Insol water, v sol alcohol and ether.

Tetra butyryl derivative

$\text{C}_6\text{H}_5\text{O}(\text{OC}_4\text{H}_7\text{O}_2)_4$. From mannite and excess of butyric acid at 200°–250°. Oil.

Di-benzoyl derivative

$\text{C}_6\text{H}_5\text{O}(\text{OBz})_2(\text{OH})_2$. From mannite and HOBz at 200° (Berthelot, *Chimie organique*, 2, 193). Soft resin, v e sol alcohol and ether.

Di-ethyl derivative $\text{C}_6\text{H}_5\text{O}(\text{OH})_2(\text{OEt})_2$. From mannite, KOH, and EtBr at 100° (Berthelot). Syrup, v sl sol water, sol alcohol, v e sol ether.

Mannitan chlorhydrin $\text{C}_6\text{H}_5\text{O}(\text{OH})_2\text{Cl}$. From mannite dichlorhydrin by boiling for two hours with 100 pts of water, neutralising with K_2CO_3 , evaporating, and extracting with ether (Bouchardat, *A Ch* [5] 6, 118). Solid mass, v e sol water, alcohol, and ether. Dextrorotatory. Conc HClAq at 100° reconverts it into mannite dichlorhydrin. Boiling water converts it into HCl and mannitan.

Mannitan dichlorhydrin

$\text{C}_6\text{H}_5\text{O}(\text{OH})_2\text{Cl}_2$. From mannite (1 pt) and fuming HClAq (15 pts) by heating for 3 days at 100° (Berthelot, *J* 1856, 661). Crystals, sol ether. KOHAq converts it into mannitan.

Mannitan dibromhydrin $\text{C}_6\text{H}_5\text{O}(\text{OH})_2\text{Br}$. From mannite dibromhydrin by boiling with water. V e sol water, alcohol, and ether. Dextrorotatory.

Mannide $\text{C}_6\text{H}_{11}\text{O}_4$. *Second anhydride of mannite*

(a) *Berthelot's mannide* $\text{C}_6\text{H}_{11}\text{O}_4$ (297°–317°). Obtained by heating mannite with butyric acid at 200°–250° (Berthelot, *A Ch* [3] 47, 312, Liebermann, *B* 17, 874). Thick deliquescent syrup, v e sol cold water and alcohol. Decomposes partially on distillation. In contact with the air it partially forms mannite.

(b) *(β)-Mannide* $\text{C}_6\text{H}_{11}\text{O}_4$ [119°] (212° at 16 mm). S 160 at 15°. S (alcohol) 27 at –16°, 73 at 12°. Obtained by treating mannite dichlorhydrin with sodium amalgam (Siwoloboff, *A* 233, 368). Prisms, v e sol water and alcohol, insol ether. Dextrorotatory. Sublimes at 14°, forming long needles. Does not yield mannite when heated with water. Air and platinum black oxidise it, forming a syrupy liquid.

(c) *Isomannide* $\text{C}_6\text{H}_{11}\text{O}_4$, *+* *e*

$$\begin{array}{c} \text{O} \quad \quad \text{O} \\ \diagdown \quad \diagup \quad \diagdown \quad \diagup \\ \text{CH}_2(\text{OH}) \text{CH} \text{CH} \text{CH} \text{CH}_2 \text{OH} \end{array}$$

[87°] (176° at 30 mm) (274° at 760 mm) $[\alpha]_D^{20} = +91.4$ in a 6 p c aqueous solution. Obtained by distilling mannite *in vacuo*, the product being fractionally distilled *in vacuo* (Fauconnier, *C R* 95, 991). Prepared by boiling mannite (200 g) with HClAq (2,000 g) for 24 hours, cooling, filtering, and distilling *in vacuo*. The distillate is allowed to stand for a fortnight, extracted with cold alcohol, the extract distilled *in vacuo*, and the frac-

tion boiling at 176° recrystallised from alcohol (Fauconnier, *Bl* [2] 41, 119). Large monoclinic, somewhat deliquescent, crystals. Boils at 274° with partial decomposition. V e sol water, m sol alcohol, sl sol chloroform, insol ether and benzene. Dextrorotatory. Not affected by heating with water at 150°. By heating with fuming HClAq in sealed tubes at 100° for 25 days it is converted into mannite dichlorhydrin $\text{C}_6\text{H}_5(\text{OH})_2\text{Cl}_2$ [174°]. It is not acted upon by POCl_3 , by bromine in the cold, or by sodium-amalgam.

Di-formyl derivative $\text{C}_6\text{H}_5\text{O}_2(\text{OCHO})_2$ [115°] (166° at 18 mm). Small plates, sl. sol cold water, v sol alcohol, sol ether.

Acetyl derivative $\text{C}_6\text{H}_5\text{O}_2(\text{OH})(\text{OAc})$ (186° at 25 mm.). From isomannide and Ac_2O . Colourless oil, sol ether.

Di-benzoyl derivative $\text{C}_6\text{H}_5\text{O}_2(\text{OAc})_2$ (198° at 28 mm). From isomannide (1 pt) by boiling with Ac_2O (3 pts) for 8 hours. Viscid liquid, with bitter taste. Not altered by further treatment with Ac_2O .

Di-benzoyl derivative $\text{C}_6\text{H}_5\text{O}_2(\text{OBz})_2$ [132°]. From mannite, alcohol, ZnCl_2 , HCl, and BzCl (Meunier, *C R* 107, 346). Monoclinic or triclinic crystals, v sl sol water and cold alcohol, sol chloroform and benzene. Not decomposed by aqueous acids or alkalis.

Methyl derivative $\text{C}_6\text{H}_5\text{O}_2(\text{OH})(\text{OMe})$ [48°] (174° at 24 mm). From isomannide, KOH, MeI, and a little water at 150°. Crystals.

Ethyl derivative $\text{C}_6\text{H}_5\text{O}_2(\text{OH})(\text{OEt})$ (165° at 17 mm). Formed by heating mannite, EtI, and conc KOHAq in sealed tubes for 4 hours at 120°. Colourless mobile liquid, sol water, alcohol, and ether.

Isomannide dichlorhydrin $\text{C}_6\text{H}_5\text{O}_2\text{Cl}_2$ [49°] (143° at 43 mm). From isomannide (1 pt) by warming with PCl_5 (2 pts). White needles or plates, v sol ether, m sol alcohol. Insol cold, sol hot, water. Has a strong aromatic odour and a peppery taste. May be distilled with steam. It is very stable, not being attacked by alcoholic KOH at 150°, by PCl_5 at 125°, or by sodium amalgam.

Anhydride of mannide $\text{C}_2\text{H}_5\text{O}_4$, *+* *e* ($\text{C}_2\text{H}_5\text{O}_2(\text{OH})_2\text{O}$) (?) Deposited on standing from a sample of butyric acid (Geuther, *A* 221, 59). Gummy.

Lævo mannite $\text{C}_6\text{H}_{11}\text{O}_4$ [164°]. Formed by reducing lævo mannose with sodium amalgam. Globular groups of fine needles, v sol water, sl sol absolute alcohol, much more sol methyl alcohol. It tastes sweet, and does not reduce Fehling's solution. A solution of this mannite containing borax turns the ray of polarised light to the left. This substance is probably the one obtained by Kiliani (*B* 20, 2714) by reduction of the double lactone of metasaccharic acid (Emil Fischer, *B* 23, 375).

Inactive mannite *a Acrite* [168°]. Formed by reducing inactive mannose with sodium-amalgam. Small prisms (from water), v sol water, m sol hot glacial acetic acid, sl sol methyl and ethyl alcohols. It is easily distinguished from ordinary mannite by its crystalline form and by its optical inactivity. A solution containing borax remains wholly inactive. This mannite is identical with (*a*) acrite, a substance obtained by the action of sodium amalgam on

(e)-acrose, a sugar obtained by the action of alkalis on acrolein dibromide (Fischer, *B* 22, 100) By oxidation with dilute HNO_3 inactive mannose is formed (Emil Fischer, *B* 23, 383)

The mannites may be distinguished optically by means of their phenyl hydrazides 1 g of the phenyl hydrazide dissolved in 1 c c of cold conc HClAq and 5 c c of water gives in a tube 100 mm long a rotation of $+1.2^\circ$, -1.2° , and 0° , with the levo-, dextro, and inactive mannite respectively In the course of 3 or 4 hours the rotation vanishes, the phenyl hydrazide being decomposed by the HCl When the di phenyl dihydrazide is used 1 g is dissolved in warm HOAc , cooled, and examined in a 100 mm tube It gives a rotation of $+0.85^\circ$, -0.85° , or 0° in the case of the levo-, dextro-, and inactive compounds respectively The dextro and levo mannites themselves scarcely exhibit a rotation, but 15 g of the mannite dissolved in 5 c c water containing 37 g borax exhibits a rotation of $+0.85^\circ$, -0.85° , or 0° , according as the mannite is dextro-, levo-, or inactive (Fischer)

MANNITIC ACID $\text{C}_6\text{H}_{12}\text{O}_7$ Prepared by mixing mannite (2 g) with platinum black (4 g), moistened with water, and exposing the mixture at 30° to 40° to the air for 3 weeks The mass is then exhausted with water, the solution ppd. by lead subacetate, the pp decomposed by H_2S , and the solution evaporated in the cold over H_2SO_4 (Gorup Besanez, *A* 118, 257) Gummy mass, sol water and alcohol, nearly insol ether It is a strong acid, and dissolves zinc with evolution of hydrogen It begins to decompose at 80° It reduces silver nitrate solution and hot Fehling's solution It is ppd by baryta and by lime water— CaA'' amorphous powder, ppd by adding alcohol to its aqueous solution CuA'' green amorphous mass left by evaporating its solution— PbA'' granular— AgA'' curdy pp

MANNITINE $\text{C}_6\text{H}_{12}\text{N}_2$ (170°) Prepared by distilling a mixture of mannite (1 mol) and ammonium chloride (2 mols) The distillate is mixed with KOHAc and shaken with ether (Seichlone a Denaro, *G* 12, 416, Etard, *C R* 92, 795) Brown oil, with strong odour, sol alcohol and ether, m sol water Gives an orange yellow pp with sodium phosphomolybdate, a reddish yellow pp with iodine in $\text{KI} \text{Aq}$, a pink pp with HgCl_2 , and a black pp with auric chloride Hypodermically injected it produces diminution of the cardiac systole, and irregular respiration, followed by insensibility and death

MANNITOSE $\text{C}_6\text{H}_{12}\text{O}_6$ Produced, together with mannitic acid, by the atmospheric oxidation of mannite in the presence of platinum black (Gorup Besanez, *A* 118, 273) Optically inactive Fermentable Reacts like glucose with alkalis, Fehling's solution, basic bismuth nitrate, and Na_2CO_3 , but it does not unite with NaCl Its alcoholic solution gives with alcoholic potash a pp of $(\text{C}_6\text{H}_{11}\text{O}_6)_2\text{K}_2\text{O}$ Mannitose is perhaps identical with levulose (Dafert, *B* 17, 228, 19, 911)

MANNONIC ACIDS $\text{C}_6\text{H}_{10}\text{O}_7$

Inactive mannonic acid *Lactone* $\text{C}_6\text{H}_{10}\text{O}_6$ [155°] Formed from inactive mannose by oxidation with bromine (E Fischer, *B* 23, 876) Long glittering prisms, grouped in stars of

needles (from alcohol), v sol hot water, sl sol hot alcohol Has a sweet taste and does not reduce Fehling's solution By means of the strychnine salt it can be split up into dextro-mannonic acid and arabinose carboxylic acid, the strychnine salt of the latter being extremely sl sol alcohol The morphine salts may also be employed, morphine dextro mannonate crystallising out *Penicillium glaucum* partly decomposes inactive mannonic acid, liberating some arabinose carboxylic acid— $\text{Ca}(\text{C}_6\text{H}_9\text{O}_6)_2$ Groups of slender needles Less soluble than calcium levo mannonate

Phenyl hydrazide $\text{C}_{12}\text{H}_{18}\text{N}_2\text{O}_6$ [230°] Colourless cubes, sl sol hot water, v sl sol alcohol Split up by hot baryta water into phenyl hydrazine and inactive mannonic acid

Dextro-mannonic acid Formed from gluconic acid by heating with twice its weight of quinoline for 40 minutes to 140° (Emil Fischer, *B* 23, 801) Formed also as above and by oxidation of dextro mannose by bromine By heating dextro mannonic acid with quinoline, gluconic acid is formed For this purpose 20 g of dextro mannonic acid, 5 g water, and 40 g quinoline are heated to 140° for 40 minutes The unchanged mannonic acid is separated by means of its brucine salt Dextro mannonic acid, like the other mannonic acids, when liberated from its salts condenses at once to the lactone $\text{C}_6\text{H}_{10}\text{O}_6$ [149°–153°] Its rotation, $[\alpha]_D = 53.8$, is equal and opposite to that of the lactone of arabinose-carboxylic acid, which is therefore levo mannonic acid

Salts— $\text{Ca}(\text{C}_6\text{H}_9\text{O}_6)_2$ 2aq— SrA' , 3aq— BaA' , (dried at 100°)

Phenyl hydrazide $\text{C}_{12}\text{H}_{18}\text{N}_2\text{O}_6$ [216°] Small colourless prisms, v sol, hot water Decomposed by hot baryta water into dextro mannonic acid and phenyl hydrazine

Levo-mannonic acid Lactone $\text{C}_6\text{H}_{10}\text{O}_6$ *Arabinose carboxylic acid* [145°–150°] $[\alpha]_D = -54.8$ Formed from arabinose (v ARABIC ACID and SUGARS) by treatment with HCy and saponification of the product (Kiliani, *B* 19, 3033) Formed also from inactive mannonic acid, as above

MANNOSE $\text{C}_6\text{H}_{12}\text{O}_6$ 16
 $\text{CHO CH(OH) CH(OH) CH(OH) CH(OH) CH}_2\text{(OH)}$
Dextro mannose $[\alpha]_D = +13^\circ$ Formed by heating 3 kilos of mannite with 20 litres of water and 10 litres of HNO_3 (S G 141) in the water bath with shaking to 40° – 45° until a test portion rendered neutral with soda gives a thick pale-yellow pp of the hydrazide with phenyl-hydrazine acetate The whole is then cooled with ice to about 25° , made feebly alkaline with crystallised Na_2CO_3 , acidified with acetic acid, and treated with 1 kilo of phenylhydrazine dissolved in dilute acetic acid The phenylhydrazide crystallised from hot water is converted into the sugar by solution in HCl (S G 119), allowing to stand, cooling, filtering, and neutralising the diluted filtrate with pure carbonate of lead The whole is again filtered, made alkaline with Ba(OH)_2 , and shaken with ether The aqueous solution separated from ether contains the sugar, which remains as a syrup on evaporating, and is ppd from absolute alcohol by ether (Emil Fischer a Josef Hirschberger, *B* 21, 1806, 22, 365), Mannose may

more easily be got from vegetable ivory nuts, *v* *Seminosae infra*

Properties—Light yellow syrup, with sweet taste. *V* sol water and alcohol. Reduces Fehling's solution, 1 cc corresponding to 0.04307 g of mannose. Turned brown by heating with 20 p.c. HClAq, the solution then containing levulic acid. Ferments easily with yeast, giving CO_2 and alcohol. The aqueous solution is dextrorotatory, but less so than glucose. Sodium-amalgam (2 p.c.) reduces mannose to mannite. Furfuraldehyde is formed by heating a dilute solution of mannose at 110° for four hours. AsCl forms a substance like acetochlorhydrone.

Phenyl hydrazide $\text{C}_{12}\text{H}_{15}\text{N}_3\text{O}_5$ [195° – 200°] *S* 1 at 100° . Prepared as above. Slender prisms (from water). *Sl* sol alcohol and acetone. Its solution in dilute HClAq is levorotatory. Concentrated mineral acids convert it into mannite and phenyl hydrazine even in the cold. When it is heated with phenyl hydrazine hydrochloride (3 pts), NaOAc (4 pts) and water (80 pts) at 105° for 4 hours there is formed the 'osazone' of glucose $\text{C}_{18}\text{H}_{22}\text{N}_4\text{O}_8$, which crystallises from alcohol in slender yellow needles [206°].

Oxime $\text{C}_6\text{H}_{11}\text{O}_2\text{N}$. Crystalline, *v* sol hot water, *insol* alcohol. When slowly heated it melts at 176° – 180° , but when quickly heated, at 184° (Fischer a Hirschberger, *B* 22, 1155).

Inactive mannose. Formed by reducing the lactone of inactive mannonic acid with sodium-amalgam (E Fischer, *B* 23, 381). Colourless syrup, *v* sol water, *sl* sol absolute alcohol, *m* sol hot methyl alcohol. Resembles (dextro) mannose except in being inactive. When subjected to fermentation levomannose is left. Bromine oxidises it to inactive mannonic acid.

Phenyl hydrazide [195°]. Decomposed on fusion. *Sl* sol water. Its solution in HCl is optically inactive. Cold conc HCl splits it up into sugar and phenyl hydrazine.

Di-phenyl hydrazide $\text{C}_{18}\text{H}_{22}\text{N}_4\text{O}_8$ [218°]. Formed by heating the inactive mannose with phenyl hydrazine (2 mols), the corresponding quantity of acetic acid and water. Fine yellow glittering needles, *sl* sol water, cold alcohol, and ether. Decomposes on fusion. At 45° conc HCl forms the corresponding osone. The solution in 60 pts glacial acetic acid is optically inactive. The properties of this di-phenyl-hydrazide are the same as those of (a) acrosazone (the di-phenyl hydrazide of (a) acrose), and Emil Fischer (*B* 23, 383) considers that they are identical.

Levo-mannose. To prepare this body 1 pt of the lactone of arabinose carboxylic acid is dissolved in 10 pts water, acidified with H_2SO_4 , and cooled to 0° . $2\frac{1}{2}$ p.c. sodium amalgam is added in small portions. The liquid must remain acid. Colourless syrup, *v* sol water, *sl* sol absolute alcohol, *m* sol methyl alcohol. The aqueous solution is slightly levorotatory. It ferments little, if at all, with yeast.

Phenyl hydrazide $\text{C}_{12}\text{H}_{15}\text{N}_3\text{O}_5$ [195°]. Formed by adding phenyl-hydrazine acetate to a solution of the sugar. Slender, almost colourless crystals. It is more sol water than the dextro mannose phenyl hydrazide. The hydrochloric acid solution is dextrorotatory. Cold HCl splits up the compound into phenyl-hydrazine and mannose.

Di-phenyl hydrazide [205°]. Formed by heating the levo mannose phenyl hydrazide with phenyl hydrazine acetate and 30 pts water to 100° . Fine yellow needles (from water). Decomposed on fusion. It is extremely like the di-phenyl hydrazide of dextro mannose, levulose, and dextrose. It, however, turns the ray of polarised light strongly to the right (Emil Fischer, *B* 23, 373).

Isomannitose, a sugar obtained, together with glucose, by boiling salep with acids, is probably identical with mannose (Tollens, *B* 21, 2150). Its phenyl hydrazide $\text{C}_{12}\text{H}_{15}\text{N}_3\text{O}_5$ [188°] is *sl* sol alcohol.

Seminosae, obtained by digesting with dilute H_2SO_4 , the cellulose composing the thick walled cells of the endosperm of vegetable ivory nuts, is identical with mannose (Reiss, *B* 22, 609, Schultz, *B* 22, 1192, Fischer a Hirschberger, *B* 22, 3218).

'Mannose carboxylic acid' $\text{C}_6\text{H}_{11}\text{O}_5$ *etc.* $\text{C}_6\text{H}_{11}(\text{CO}_2\text{H})\text{O}_2$. From mannose (50 g) by dissolving in water (250 g) and adding anhydrous HCl (18 cc) and a few drops of ammonia. After three days the mixture is heated to 50° for 4 hours. The product appears to contain the ammonium salt of the acid and its amorphous a nide [183°], which may be pptd by alcohol (Fischer a Hirschberger, *B* 22, 365). The acid changes to anhydride when set free from its salts. In preparing the lactone (*v infra*) an acid [c 169°], probably the free mannose carboxylic acid, is sometimes formed.

Salts—*BaA'*. Obtained by boiling the ammonium salt with baryta. Colourless, semi-crystalline mass, *m* sol hot, *v* *sl* sol cold, water, *insol* alcohol. Boiling conc HIAq reduces it slowly to *n* heptonic acid.

Lactone $\text{C}_6\text{H}_{10}\text{O}_5$ [150°]. Obtained by decomposing a hot solution of the *Ba* salt with the theoretical quantity of H_2SO_4 . Needles, *v* sol water, *m* sol alcohol, *insol* ether.

Phenyl hydrazide $\text{C}_{18}\text{H}_{22}\text{N}_4\text{O}_8$, *Ph* [220° – 223°]. Formed by boiling the *Ba* salt with aqueous NaOH , and adding HOAc and phenyl hydrazine acetate (Fischer a Passmore, *B* 23, 2732). Very small prisms, *sol* hot water. Decomposed on fusion.

MANTLE MUCIN *v* PROTEIDS, *Appendix C*.

MARGARIC ACID $\text{C}_{17}\text{H}_{33}\text{O}_2$ (*Heptadecanoic acid*) *Mol* *v* 270 [60°]. The name margaric acid was originally applied to an acid $\text{C}_{17}\text{H}_{33}\text{O}_2$ supposed to occur in the product of the saponification of solid natural fats, but this acid was shown by Heintz to be a mixture of palmitic acid $\text{C}_{16}\text{H}_{32}\text{O}_2$ and stearic acid $\text{C}_{18}\text{H}_{36}\text{O}_2$. An acid $\text{C}_{17}\text{H}_{33}\text{O}_2$ occurs, however, according to Ebert (*B* 8, 775), in adipocere, a substance formed in dead tissues of animals. Formed by boiling its nitrile (cetyl cyanide) with alcoholic potash (Heintz, *P* 102, 272). White crystals—*BaA'*, white amorphous powder—*AgA'* white amorphous powder.

Nitrile $\text{C}_{16}\text{H}_{31}\text{CN}$ [53°] (?) (Köhler, *J* 1856, 579, cf Becker, *A* 102, 213). According to Heintz (*J* 1857, 445), margaritrile prepared from potassium cetyl sulphate and KCN is an oil.

Margaric acid $\text{C}_{17}\text{H}_{33}\text{O}_2$ [60° uncor.]. (277° at 100 mm). Prepared by the oxidation

of methyl-heptadecyl-ketone (Krafft, *B* 12, 1672) Probably identical with the preceding acid

MARJORAM OIL $\text{SG} \approx 911$ (Bruylants)

The essential oil obtained by steam distillation from the flowers of sweet marjoram (*Majorana hortensis* or *Origanum majorana*) Yellow oil which becomes brown on standing It has a pungent smell, a hot peppery taste, and an acid reaction According to Bruylants (*J Ph* [4] 30, 33, cf Mulder, *A* 31, 69) it begins to boil at 185° and the thermometer remains stationary at 215° – 220° Bruylants found it to consist of 5 p.c. of dextrorotatory terpene, 85 p.c. of a mixture of dextrorotatory camphor and borneol, and 10 p.c. of resin According to Kane (*A* 32, 285) the essential oil of wild marjoram (*Origanum vulgare*) contains a terpene which boils at 161° and has SG 867 According to Beilstein a Wiegand (*B* 15, 2855) oil of sweet marjoram contains a terpene $\text{C}_{10}\text{H}_{18}$ (178° 1 V) $\text{SG}^{18.5}$ 846, which absorbs HCl , forming liquid $\text{C}_{10}\text{H}_{17}\text{HCl}$ By distilling the oil of marjoram over sodium Beilstein a Wiegand obtained liquid $\text{C}_{10}\text{H}_{18}\text{O}$ (200° – 220°)

Cretan oil of marjoram from *Origanum hirtum* has $\text{SG} \approx 951$, and 100 mm produce a rotation of -0.4° It contains 50 p.c. of carvacrol [2 $^\circ$] and dissolves in half its volume of 15 p.c. soda solution (Jahns, *Ar Ph* [3] 15, 1, 16, 277) The oil also contains a mixture of terpenes, and a phenol that is coloured violet by FeCl_3 , Jahns found a little carvacrol in the oil from *Origanum vulgare*

MARRUBIUM [160°] The bitter principle of white horehound (*Marrubium vulgare*) (Kro Mayer, *Ar Ph* [2] 108, 257, Harms, *J* 1863, 593) Prepared by exhausting the dried herb with hot water, evaporating, and extracting with alcohol Large tables (from ether) or needles (from alcohol) Almost insol cold, sl sol hot, water Not affected by alkalis Metallic salts do not ppt it Not pptd by tannin Hot conc HClAq does not affect it Conc H_2SO_4 gives a brownish yellow solution Cold HNO_3 has no action, hot HNO_3 forms a yellow solution

MARSH GAS v METHANE

MARTYLAMINE v p AMIDO DIPHENYL

MASOPIN $\text{C}_{22}\text{H}_{40}\text{O}$ [155°] Occurs in a resin used in Mexico for chewing, said to be the dried juice of a tree called Dschlite The resin occurs in porous lumps, smelling like rotten cheese, but with little taste The masopin is obtained by boiling the resin with water, and crystallising the residue from alcohol (Genth, *A* 46, 124) White silky needles (from ether) After fusion, it melts a second time at 70° On distillation it yields a terpene and a crystalline acid, of which the Ag salt contains 45.5 p.c. of silver

MASTIC A resin obtained by incisions in the bark of *Psittacia lentiscus*, a tree growing in Chios Small round transparent grains with faint agreeable odour Used for making varnishes and, from the earliest times, for chewing Softens when masticated SG 1.074 Aqueous alcohol dissolves the greater part $\text{C}_{15}\text{H}_{26}\text{O}_2$, leaving masticin $\text{C}_{16}\text{H}_{28}\text{O}_2$ undissolved (Johnston, *P T* 1839, 182) Flückiger (*Ar Ph* 219, 170) found 2 p.c. of a dextrorotatory terpene in mastic.

MATEZITE $\text{C}_{15}\text{H}_{26}\text{O}$, [187°] $[\alpha]_D = -64.7$

Occurs in Madagascar caoutchouc (mateza rorina) (Girard, *Bl* [3] 21, 220, *C R* 110, 84) Identical with (8) Pinite (Combes, *C R* 110, 46). Crystalline nodules, v sol water, m sol alcohol Sublimes above 200° Resembles dambonite but is dextrorotatory Fuming HIAq splits it upon heating into Mel and matezo dambose $\text{C}_{15}\text{H}_{26}\text{O}_2$ [246°] $[\alpha]_D = 67.6$, which is dextrorotatory and more soluble in water than dambose Matezo dambose is identical with the (8) inosite of Maquenne (*C R* 109, 812) It forms small tetrahedra when pptd by adding alcohol to its aqueous solution

MATICIN A bitter substance which remains dissolved in the water in the retort in the preparation of the essential oil of matico by distilling the leaves of *Piper aspersifolium* with steam It is sol water and alcohol, and insol ether (Hodges, *P M* [3] 25, 204, *C S Mem* 1, 123) Its solution is not pptd by lead acetate

MATICO-CAMPHOR $\text{C}_{15}\text{H}_{26}\text{O}$ [94°] Occurs in the essential oil of matico leaves (*Piper angustifolium*), from which it is obtained by distilling off the greater part and allowing the residue to crystallise (Kugler, *B* 16, 2841) Hexagonal crystals V sol alcohol, ether, benzene, petroleum ether, &c With HCl it gives a violet colour, passing into blue, and finally becoming green. H_2SO_4 colours it yellow, then red, and lastly violet A mixture of H_2SO_4 and HNO_3 first produces a yellow, which then becomes violet, and finally a splendid blue

MAUVANILINE $\text{C}_{15}\text{H}_{11}\text{N}_3$ A by product in the preparation of rosaniline by the oxidation of crude aniline (De Laire, Girard a Chapoteaut, *C R* 64, 410, *Bl* [2] 7, 366), cf Girard a Pabst, *Bl* [2] 34, 37) Light brown crystals (containing $\frac{1}{2}$ aq, which is given off at 130° with decomposition) Insol cold, v sl sol hot, water, sol alcohol, ether, and benzene Its salts form lustrous bronze green crystals, v sl sol cold water, and dye silk and wool mallow red Tri ethyl-mauvaniline $\text{C}_{15}\text{H}_{11}\text{Et}_3\text{N}_3$ dyes bluish violet, while tri phenyl mauvaniline $\text{C}_{15}\text{H}_{11}\text{Ph}_3\text{N}_3$ is a blue dye

MAUVEINE $\text{C}_{15}\text{H}_{11}\text{N}_3$ The first aniline dye introduced (Perkin, 1856) Obtained by adding a cold dilute solution of a salt of crude aniline to a cold dilute solution of $\text{K}_2\text{Cr}_2\text{O}_7$, and leaving the mixture to stand for 12 hours The resulting black pp is dried, extracted with benzene, dissolved in alcohol, and the filtrate evaporated When aniline sulphate is used the product is $(\text{C}_{15}\text{H}_{11}\text{N}_3)_2\text{SO}_4$, aniline hydrochloride yields $\text{C}_{15}\text{H}_{11}\text{N}_3\text{Cl}$ (Perkin, *A* 131, 200, *C J* 35, 717) On adding aqueous KOH to a solution of a salt of mauveine the base separates as a violet black crystalline substance It dissolves in alcohol, forming a violet solution, which on addition of acids turns purple Mauveine is insol benzene and ether With dilute acids it forms purple solutions, with stronger acids, blue Conc H_2SO_4 forms a dirty green solution Dyes silk mauve Used for postage stamps Oxidised by PbO_2 and boiling HOAc it gives para safranine $\text{C}_{15}\text{H}_{13}\text{N}_3$

Salts — B/HCl tufts of small prisms with green metallic lustre Insol cold, sl sol hot, water, m sol alcohol, nearly insol ether — B/HCl blue, with copper lustre Unstable. Becomes B/HCl when dissolved in alcohol —

B⁺ H₂PtCl₆, green crystalline powder, v sl sol alcohol —B⁺ H₂PtCl₆, blue —B⁺ H₂SO₄, —B⁺ HBr —B⁺ Hl lustrous prisms —B⁺ H₂SO₄, —B⁺ H₂CO₃, prisms, with green metallic lustre On boiling its solution CO₂ is given off

Ethyl mauveine C₂₂H₂₀EtN₂, Formed by heating mauveine with EtI at 100° Its colour is redder than that of mauveine During the manufacture crystals of C₂₂H₂₀EtN₂HI, sometimes appear —C₂₂H₂₀EtN₂HI Forms a reddish purple solution in alcohol —(C₂₂H₂₀EtN₂HI).PtCl₆, Golden green lustre

Pseudo-mauveine C₂₂H₂₀N₂ (Perkin, G. J. 85, 725) Present in commercial mauveine It is more soluble in alcohol than mauveine It is a strong base, and dyes silk almost exactly like mauveine Pure aniline gives pseudo mauveine on oxidation, hence it does not contain methyl Mauveine is probably derived from aniline and *p* toluidine The violet colouration given by bleaching powder to a dilute solution of aniline is probably due to pseudo-mauveine —C₂₂H₂₀N₂HI Greenish-golden lustre —(B⁺ HCl).PtCl₆

MAYER'S REAGENT v ALKALOIDS, Reaction 3

MAYNAS RESIN *Calaba* or *Galba* of the Antilles (Lewy, A. Ch. [3] 10, 380) A resin extracted by incision from *Calophyllum calaba* Alcohol extracts from it C₂₁H₃₄O₄, which separates on cooling in yellow monoclinic prisms Maynas resin is insol water, but v sol KOHAq and ammonia It melts about 105°

MECCA BALSAM *Balm of Gilead* The produce of *Balsamodendron gileadense*, a shrub growing in Arabia Felix There appear to be several varieties of it It contains a fragrant volatile oil, an acid resinol alcohol, and a resin, insoluble in alcohol (Bonastre, A. 3, 147, cf Trommsdorff (Trommsd. Neues Journal, 16, 62)

MECONIC ACID C₁₅H₈O₆, 3aq, *tc* C₁₅H₈O₆(OH)(CO₂H)₂, Mol w 200 S 25 at 100° Heat of neutralisation (4 mols NaOH) 87,382 (Gal a Werner, B1 [2] 47, 162) Electrical conductivity Ostwald (J. pr. [2] 32, 368)

Occurrence —In opium (Serturmer, Am. S. 55, 72, 57, 183, 64, 65, Robiquet, A. Ch. 5, 232, 51, 236, 53, 425, Liebig, A. 7, 87, 26, 118, 147)

Preparation —Opium is exhausted with water at 38°, the extract is neutralised with CaCO₃, evaporated to a syrup, and mixed with a concentrated solution of CaCl₂, which ppts calcium meconate The pp (1 pt) is suspended in conc HClAq (3 pts) mixed with boiling water (20 pts), and kept near 100° till dissolved On cooling, acid calcium meconate separates in crystals These (1 pt) are again dissolved in a mixture of HClAq (3 pts) and boiling water (20 pts), and, on cooling, meconic acid separates (Gregory, A. 24, 43) It may be further purified by crystallising its ammonium salt from water (How, A. 83, 350)

Properties —Micaceous scales or small trimetric prisms (containing 3aq) (Burghardt, C. J. 27, 937) It gives off its water of crystallisation at 100°, then becoming a white opaque mass Has a sour taste and strongly reddens litmus Sl sol cold water, v sl alcohol, sl sol ether FeCl₃ colours its solutions blood red This colouration, like that with sulphocyanides, is

much weakened by oxalic acid and by metaphosphoric acid (Dupré, C. N. 32, 15), but it is not destroyed by boiling or by dilute HClAq Meconic acid may be regarded as a derivative of the hypothetical $\text{CH} \begin{smallmatrix} \text{CH} & \text{CH} \\ \diagdown & \diagup \\ \text{CH} & \text{CH} \end{smallmatrix} > \text{O}$

Reactions —1. Heated at 180° it gives off CO₂, and comenic acid C₈H₆O₄(OH)(CO₂H) remains At a higher temperature the comenic acid is further resolved, partly into CO₂ and pyromeconic acid C₈H₆O₄(OH), and partly into water, acetic acid, and oily and carbonaceous products 2 When boiled with water or with HClAq it gives CO₂ and comenic acid —3 Nitric acid oxidises it, forming oxalic acid —4 Boiling conc KOHAq also yields oxalic acid —5 Boiling aqueous ammonia forms comenamic acid C₈H₆NO(OH)CO₂H Comenamic acid yields pyridine on dry distillation with zinc-dust (Lieben a Hattinger, B. 16, 1263) The comenamic acid is di oxy picolinic acid, and comenic acid is probably an intermediate product in its formation —6 Bromine, acting on its aqueous solution, forms a brominated comenic acid —7 Aqueous HI gives comenic acid —8 Sodium amalgam reduces it to hydro meconic acid —9 Gives with alcohol and HCl a di ethyl ether, C₈H₆O₄(OH)(CO₂Et)₂, whence a silver compound, C₈H₆O₄(OAg)(CO₂Et)₂, which gives with EtI a tri ethyl derivative C₈H₆O₄(OEt)₃(CO₂Et)₂ [61°] This gives no colour with FeCl₃ (Öst a Mennel, J. pr. [2] 23, 439) —10 PCl₅ converts it into a liquid chloride, probably C₈H₆Cl₂(COCl)₂, whence ice water forms 'chloro meconic' acid, C₈H₆ClO(OH)(CO₂H)₂, aq [165°], whereas warm water forms only oxalic acid (Hilsebein, J. pr. [2] 32, 136) Chloro meconic acid gives a green colour, with solution of ferric chloride It forms an ether, C₈H₆ClO(OH)(CO₂H)(CO₂Et)₂, [148°], which gives an acetyl derivative C₈H₆ClO(OAc)(CO₂H)(CO₂Et)₂, [70°] It also forms barium salts Chloro meconic acid is decomposed by sublimation into carbonic acid and yellowish needles of *pyro chloro meconic acid*, C₈H₆ClO(OH)₂, aq [174°] The solution of this acid is turned dark green by FeCl₃ Its alcoholic solution turns the skin red It forms a crystalline calcium salt, (C₈H₆ClO₂)₂Ca Chloro meconic acid dissolves in conc NH₃, the solution turning successively red, violet, and blue, owing to the formation of two nitrogenous colouring matters, 'meconic blue' and 'meconic red' Chloro meconic acid is reduced by treatment with sodium amalgam, forming *di-hydro chloro meconic acid* C₈H₆ClO(OH)(CO₂H)₂, [145°] Fuming HI at 100° reduces chloro meconic acid to an *oxy pentane di carboxylic acid*, C₈H₁₀O(CO₂H)₂ This body forms groups of needles [149°] Its silver salt, Ag₂A'', is a crystalline powder, insol water Its barium salt, BaA'', forms white prisms Its ether, Et₂A'', is an oil, boiling about (250°) Oxy-pentane di carboxylic acid, when distilled with lime, yields amyl alcohol, (c 130°) (Hilsebein J. pr. [2] 32, 129–153)

Salts —(NH₄)₂HA''' aq granular crystals, sl sol water —(NH₄)₂HA''' aq slender needles —A barium salt is ptd by adding BaCl₂ to a solution of an alkaline meconate, in white flakes, soluble in acetic acid —CaHA'''₂ aq ptd by adding CaCl₂ to aqueous meconic acid —CaHA''' aq yellow gelatinous pp, formed by

adding CaCl_2 to a solution of ammonium meconate.—The cupric salts are green pps.— $\text{Pb}_2\text{A}''$, 2aq white flocks, ppd by adding lead acetate to a solution of meconic acid (Stenhouse, *A* 51, 281).—A sparingly soluble ferric salt may be obtained by the action of ferric sulphate on ammonium meconate (Stenhouse)— $\text{Fe}_2\text{Ca}_2\text{A}'''$, 6aq (Rennie, *C N* 42, 75)— $\text{Ag}_2\text{HA}''$ white pp got by adding AgNO_3 to a solution of meconic acid— $\text{Ag}_2\text{A}''$ yellow pp from ammonium meconate and AgNO_3 .—Aniline salt $(\text{C}_6\text{H}_5\text{NH}_2)_2\text{C}_2\text{H}_3\text{O}_2$, crystalline (Korff, *A* 138, 195)—Urea salt $(\text{CON}_2\text{H}_4)_2\text{C}_2\text{H}_3\text{O}_2$, (Hlasiwetz, *J* 1856, 699)—Narceine salt [126°] from narceine (1 mol) and meconic acid (1 mol) Lemon yellow crystals, sol hot water (Merck, *C G* 1889, 384).—The acid morphine salt could not be obtained by Dott (*Ph* [3] 17, 690) It will be observed that meconic acid has but little tendency to form tribasic salts, so that it might perhaps be more correctly regarded as a dibasic acid (cf Dittmar & Dewar, *Pr E* 1867, Dott, *Ph* [3] 11, 576)

Ethyl derivative $\text{C}_2\text{H}_3\text{O}_2(\text{CO}_2\text{H})_2(\text{OEt})$ [200°] Formed by boiling $\text{C}_2\text{H}_3\text{O}_2(\text{CO}_2\text{Et})_2(\text{OEt})$ for forty eight hours with water, and evaporating the solution Small prisms grouped in lumps Decomposed by fusion Gives no colour with FeCl_3 Gives, with most metallic salts, no pp Lead acetate is an exception Heated alone till it melts, CO_2 is given off, and the ethyl derivative of meconic acid formed Its aqueous solution is converted by bromine into a brominated derivative of meconic ether— PbA'' 1½aq Crystalline pp

Mono ethyl ether

$\text{C}_2\text{H}_3\text{O}_2(\text{CO}_2\text{Et})(\text{CO}_2\text{H})(\text{OH})$ [179°] Prepared by dissolving meconic acid in alcohol, heating to 100°, and subjecting to a current of HCl till crystals begin to appear The liquid on cooling deposits crystals of the mono ethylic ether (E Mennel, *J pr* [2] 26, 450) It gives a red colour with FeCl_3 .—Silver salt $\text{C}_2\text{H}_3\text{O}_2(\text{CO}_2\text{Et})(\text{CO}_2\text{Ag})(\text{OH})$, aq—Barium salts $\text{BaC}_2\text{H}_3\text{O}_2$, and $\text{Ba}(\text{C}_2\text{H}_3\text{O}_2)_2$.

Di ethyl ether $\text{C}_2\text{H}_3\text{O}_2(\text{CO}_2\text{Et})_2(\text{OH})$ [112°] Prepared by passing HCl into an alcoholic solution at 100° until the crystals (of mono ethylic ether) which are first formed are dissolved up again On pouring into water, plates of the di ethylic ether separate (How, *A* 83, 350, Mennel, *J pr* [2] 26, 453) Plates From dilute solutions it crystallises with ½aq as needles Gives a red colour with FeCl_3 An amorphous pp is formed by adding AgNO_3 and then exactly neutralising with NH_3 .— $\text{NH}_4\text{C}_2\text{H}_3\text{O}_2$, needles (from alcohol), v sol water Its solution is ppd by various metallic salts

Ethyl derivative of the diethyl ether $\text{C}_2\text{H}_3\text{O}_2(\text{CO}_2\text{Et})_2(\text{OEt})$ [61°] From $\text{C}_2\text{H}_3\text{O}_2(\text{CO}_2\text{Et})_2(\text{OAg})$ and EtI Gives no red colour with FeCl_3 Boiled with water it changes to the ethyl derivative of meconic acid (q v) Not acted upon by bromine*

Mono-amide

$\text{C}_2\text{H}_3\text{O}_2(\text{CONH}_2)(\text{CO}_2\text{H})(\text{OH})$ aq **Meconamic acid** From the above by the action of aqueous NH_3 , followed by HCl Crystallises in hard lumps from water Not decomposed by cold NaOH , converted by boiling NaOH into meconate **Copper salt** $\text{C}_2\text{H}_3\text{O}_2\text{Cu}$ 2aq

Basic ammonium salt of amide $\text{C}_2\text{H}_3\text{O}_2(\text{CONH}_2)(\text{CO}_2\text{NH}_2)(\text{ONH}_2)$ This yellow pp is the first product of the action of NH_3 upon mono ethylic meconate

Di amide $\text{C}_2\text{H}_3\text{O}_2(\text{OH})(\text{CONH}_2)_2$ From the diethyl ether and boiling aqueous NH_3 (How) Powder, sl sol cold water

Oxim $\text{C}_2\text{H}_3\text{O}_2\text{NH}$ Formed by treating meconic acid with hydroxylamine hydrochloride (Oderheimer, *B* 17, 2081) Small colourless needles (containing H_2O) V sol water, sl sol alcohol, ether, and chloroform, insol ligroine Decomposes suddenly at 190° It gives a red colour with FeCl_3 It reduces Fehling's solution By boiling with HCl hydroxylamine is split off— $\text{A}''\text{Na}_2$ colourless crystals— $\text{A}''\text{Ag}$, aq white sparingly soluble pp— $\text{A}''\text{Ba}$ 10aq sparingly soluble concentric needles— $\text{A}''\text{Ca}$ 2aq, sparingly soluble needles

Hexahydrate $\text{C}_2\text{H}_3\text{O}_2$, **Hydromeconic acid** Obtained by reducing meconic acid suspended in water by adding sodium amalgam (Korff, *A* 138, 191) Separated by decomposing the lead salt by H_2S and evaporating the filtrate Strongly acid syrup, v sol water, m sol alcohol, insol ether Ppd as deliquescent flocks on adding ether to its alcoholic solution Not affected by HI Gives no colour with FeCl_3 — BaA'' 2aq amorphous pp, sol water, insol alcohol— PbA'' 1½aq amorphous pp— $\text{Ag}_2\text{A}''$ ½aq white granular pp

Reference—PYROMECONIC ACID AND PYROMECONAZONIC ACID

MECONIDINE $\text{C}_{21}\text{H}_{29}\text{NO}_4$, [58°] An alkaloid contained in the aqueous extract of opium (Hesse, *A* 153, 47) The aqueous extract is ppd by Na_2CO_3 , the pp dissolved in ether, the ethereal solution shaken with dilute acetic acid, and the acid solution poured into aqueous NaOH After 24 hours the pp of thebain and papaverine is removed by filtration, the filtrate neutralised by HCl , rendered alkaline by NH_3 , and shaken with chloroform The chloroform is then shaken with acetic acid, and the acid solution exactly neutralised by ammonia, which ppts lanthopine The filtrate is mixed with KOH and shaken with ether Codamine and laudanum separate, and the mother liquor is then shaken with dilute acetic acid, the solution saturated with NaCl , the ppd meconidine hydrochloride decomposed by NaHCO_3 , and the base extracted with ether and dried at 90°

Properties—Brownish yellow transparent amorphous mass Cannot be sublimed Tasteless Insol water, v sol alcohol, ether, benzene, chloroform, and acetone Its alcoholic solution turns red litmus blue Sl sol NH_4Aq , v sol NaOHAq Can be extracted by ether from its solution in ammonia but not from that in caustic soda Its solution in dilute H_2SO_4 becomes rose coloured in a few minutes, and purple red on standing for some time, or immediately on boiling Conc H_2SO_4 gives an olive-green solution Conc HNO_3 colours it orange red The dilute solutions of the salts have an intensely bitter taste— $\text{B}'\text{H}_2\text{PtCl}_6$ yellow amorphous pp which soon turns reddish The aurochloride is a dingy yellow amorphous pp

MECONIN $\text{C}_{11}\text{H}_{13}\text{O}_6$, &c

$\text{C}_2\text{H}_3(\text{OMe})_2 < \text{CO} > \text{O}$. [1.6.3] **Anhydride of**

Meconic acid. *Opranyl* Mol w 194 [99°] (M & F), [102.5°] (Wegscheider, M 3, 351, Prinz, J pr [2] 24, 371) S 4 in the cold, 5 at 100°. A neutral substance existing in opium (Dublanc, A Ch [2] 49, 17, Couerbe, A Ch 49, 11, 50, 337, 59, 148) It may also be extracted by ether from the root of *Hydrastis canadensis* (Freund, B 22, 459)

Formation—1 Together with cotarnine, opianic acid, and hemipic acid, by the action of warm dilute nitric acid on narcotine (Anderson, Tr E 20, 347, 21, 204)—2 From opianic acid by the action of sodium-alumalgar, or of zinc and H₂SO₄ (Matthiessen & Foster, C J 16, 849)—3 Together with hemipic acid by the action of caustic alkalis on opianic acid (M & F)

Preparation—1 A concentrated aqueous extract of opium is pptd by dilute ammonia, and the filtrate evaporated to crystallisation. The crystals are extracted with alcohol, which after evaporation deposits crystals of meconin, which are recrystallised from water and ether successively (Couerbe)—2 The aqueous extract of opium is precipitated by chloride of calcium, the precipitated meconate of calcium is filtered, the filtrate evaporated to the crystallising point, and separated from the deposited hydrochloride of morphine, and the dark mother liquor is diluted with water, filtered from the flocks which separate, and treated with ammonia, which precipitates narcotine, thebaine, and a large quantity of resin. The filtrate is mixed with acetate of lead the excess of lead is removed from the filtered liquid by dilute sulphuric acid, and the filtrate is neutralised with ammonia, and evaporated to the crystallising point at a moderate heat, whereupon narcotine separates out, and then sal ammoniac by further concentration. The mother liquor is repeatedly digested with $\frac{1}{2}$ vol ether at 26°, and the ether is distilled off from the extracts, a brown syrup then remaining. On treating this syrup with dilute hydrochloric acid, papaverine dissolves, and meconin remains in the form of a dark-grey crystalline powder, which, to free it from resin and purify it completely, must be several times crystallised from boiling water, with addition of animal charcoal (Anderson)

Properties—Colourless hexagonal prisms. Appears tasteless at first, but afterwards acid. May be sublimed. Sol cold water, m sol alcohol and ether. Sol KOHAq, nearly insol NH₄Aq. Inactive. Its aqueous solution ppts lead subacetate but not lead acetate. Dilute H₂SO₄ forms a colourless solution which becomes dark green when evaporated (Couerbe). Conc H₂SO₄ forms a colourless solution which becomes purple on heating.

Reactions—1 Conc HClAq at 100° gives MeCl and methyl-normeconin C₁₀H₇O₄ (M & F). HI acts in like manner.—2 Baryta water dissolves meconin forming 'barium meconinate' (C₁₀H₇(OMe)₂(CH₂OH)CO₂)₂ Ba. This salt crystallises in needles, but the free acid splits up at once into water and its anhydride meconin (Hessert, B 11, 240, Prinz, J pr [2] 24, 373).—3 Potash-fusion forms methyl-normeconin C₉H₇O₄ [125°] and, finally, protocatechuic acid.—4 Oxidised by MnO₂ and dilute H₂SO₄ to opianic acid.—5. KMnO₄ oxidises it to hemipic acid [180°].

Chloro-meconin C₁₀H₇ClO₄ [175°] Obtained by passing chlorine into a cold saturated aqueous solution of meconin (Anderson, A 98, 47). Colourless needles. May be sublimed. Sol cold water, v sol alcohol and ether. Hot H₂SO₄ gives a greenish blue colour.

Bromo-meconin C₁₀H₇BrO₄ [167°] (Anderson), [177°] (Salomon, B 20, 888). From meconin and bromine-water. Colourless needles, sl sol water.

Iodo-meconin C₁₀H₇IO₄ [112°] From meconin and chloride of iodine (A). Needles, nearly insol water, m sol alcohol and ether.

Nitro-meconin C₁₀H₇(NO₂)O₄ [158°] From meconin and HNO₃. White needles (from alcohol).

Amido-meconin C₁₀H₇(NH₂)O₄ [171°] Obtained by reducing nitro-meconin with iron and acetic acid (S). M sol hot benzene.

Methyl-normeconin C₉H₇O₄ + e

C₉H₇(OH)(OMe) < $\begin{matrix} \text{CO} \\ \text{CH} \end{matrix}$ > O [125°] Obtained

by heating meconin with conc HClAq (Matthiessen & Foster), by heating meconin with KCy at 180° (Bowman, B 20, 890), or by potash fusion from meconin or narcotine (Beckett & Wright, J 1876, 810). Monoclinic prisms, v sol hot water and alcohol, sl sol ether. FeCl₃ colours it blue. It reduces silver salts in the cold. Potash fusion converts it into protocatechuic acid—CaA'—BaA'.

Meconin acetic acid C₁₁H₁₁O₆ + e

C₁₀H₇(OMe) < $\begin{matrix} \text{CO} \\ \text{CH} \end{matrix}$ > O [167°] Obtained

by heating opianic acid with a mixture of malonic acid, acetic acid, and sodium acetate for 10 hours at 100° (Liebermann & Kleemann, B 19, 2290). Glistening needles. By boiling with baryta water it is converted into opianyl acetic acid C₁₀H₇(OMe)(CO₂H)CH(OH)CH CO₂H. By heating with HI it is dimethylated, giving

C₁₀H₇(OH) < $\begin{matrix} \text{CO} \\ \text{CH} \end{matrix}$ > O [228°], of which the ethyl ether melts at 181°.

Salts—A'A₂Ca white slightly soluble crystals. —A'A₂Cu needles, sl. sol water. —A'A₂Cu* blue crystalline pp.

Methyl ether A'Me [124°], glistening plates.

Ethyl ether A'Et [83°], plates, sol alcohol, ether, and hot water.

Nitro meconin acetic acid

C₁₀H(NO₂)OMe < $\begin{matrix} \text{CO} \\ \text{CH} \end{matrix}$ > O [176°] Nearly

colourless crystals. Obtained by nitration of meconin acetic acid. It dissolves in cold H₂SO₄ with a yellow colour, which becomes red on warming from formation of an indigo derivative. By tin and HCl it is reduced to the lactone of (Py 1.8)-di-oxy-(B'2.8) di methoxy-di hydroquinoline-(B 1)-carboxylic acid.

C₉H(OMe) < $\begin{matrix} \text{CO} \\ \text{CH} \end{matrix}$ > O [256°].

Salts—A'A₂Ca. yellow needles. —A'A₂G* curdy pp. —A'A₂Cu* : green pp.

Ethyl ether AEt [129°], glistening needles, insol water, sol alcohol and benzene (Liebemann & Kleemann, *B* 19, 2295)

ψ **Meconin** $C_8H_4(OMe)_2 \langle \frac{CH}{CO} \rangle O$ [163°] ?

Di methyl derivative of di oxy phthalide [124°] When hemipimide, the imide of hemipic acid $C_8H_4(OMe)_2 \langle \frac{CO}{CO} \rangle NH$, is heated with tin and hydrochloric acid it is reduced to hemipimidine $C_8H_4(OMe)_2 \langle \frac{CH}{CO} \rangle NH$ [181°], which yields a

nitroso derivative • $C_8H_4(OMe)_2 \langle \frac{CH}{CO} \rangle N NO$ [156°], which on treatment with dilute NaOH in the cold yields pseudo meconin (Salomon, *B* 20, 883) Obtained also by boiling hemipimide with zinc dust and acetic acid Long colourless needles (from water) Sl sol cold water, sol benzene, alcohol, and ether Unlike meconin it is not affected by boiling with dilute H₂SO₄ and MnO₂. Dilute HNO₃ at 150° forms nitro hemipic acid and nitro ψ meconin, whereas meconin yields only nitro meconin Fusion with KOH or KCy does not affect ψ meconin

Bromo ψ meconin $C_{10}H_6BrO$, [142°] From ψ meconin and bromine White flocculent pp, sol benzene, insol petroleum

Nitro ψ meconin $C_{11}H_6(NO_2)_2O$, [166°] From ψ meconin and fuming HNO₃ Yellow needles, which yield oxalic acid on further treatment with nitric acid

Amido ψ meconin $C_{10}H_6(NH_2)_2O$, [165°] Obtained by reducing the preceding More basic than amido meconin

MECONOISIN $C_{15}H_{10}O$, [88°] S 37 Occurs in opium, and found in the mother liquor after separation of meconin When this is allowed to stand, crystals are deposited, which are washed with alcohol and recrystallised from water (T a H Smith, *Ph* [3] 8, 981) Large, leaf shaped crystals, v e sol hot water It gives a green colour when heated with dilute H₂SO₄

MEDULLIC ACID $C_{15}H_{22}O_2$, [72.5°] A fatty acid said to be produced, together with stearic and palmitic acids, by the saponification of beef-marrow (Eylerts, *Ar Ph* [2] 104, 129)

MELAM v CYANIC ACIDS

MELAMINE v *Cyanuramide* in the article CYANIC ACIDS

MELAMPYRITE v DULCITE

MELANILINE v DI PHENYL GUANIDINE

MELANIN C 60, H 4.8, N 10.8, ash 2.2 A black pigment covering the choroid membrane of the eye (Scherer, *A* 40, 63) Insol water, alcohol, and ether

Phymatorhusin C 55.7, H 6.0, N 12.3, S 8 to 9, Fe 0.7 to 2 A pigment occurring in melanotic urine and tumours It is obtained by ppg with baryta, dissolving in Na₂CO₃, and ppg with dilute H₂SO₄ (Berdez a Nencki, *J Th* 1886, 477, Möerner, *H* 11, 81) It is a brownish-black amorphous powder, insol water, alcohol, ether, chloroform, and dilute mineral acids, v e sol ammonia, aqueous NaOH, Na₂HPO₄, and Na₂CO₃ Ppd from its solution in NaOH by baryta, MgSO₄, and BaCl₂ With potash fusion it gives skatole, volatile fatty acids, nitriles, KCy, and K₂S Hot H₂SO₄ forms pyridine Phymatorhusin is accompanied by another black substance, which differs from it in being soluble

in 50-75 p c acetic acid. It contains 59 p c of sulphur and 2 p c Fe

Hippomelanin C 53.5-55.6, H 2.7-3.9, N 10.5-10.9, S 2.8-3.0 Occurs in melanotic tumours of horses May be obtained by diluting the emulsion of a melanotic spleen with water and adding CaCl₂ and Na₂HPO₄ The pp is washed at 40°, and then treated with a digestive fluid until the solution ceases to give a reaction for peptones The residue is washed with aqueous soda, alcohol and ether (Miura, *C C* 1887, 250). Brownish black powder, insol water, alcohol, and ether Dissolves on warming in dilute acids and alkalis Potash fusion gives KCy, succinic acid, formic acid, and hippomelanin acid Hippomelanin acid is a black amorphous body, sol ammonia, and reppd by HCl It contains less S and more C than the melanin (Nencki a Sieber, *C C* 1888, 587) The black pigment in dark hair and in bird's feathers after purifying by alcoholic NH₃ and dilute H₂SO₄ may be represented by the formula $C_{10}H_7N_2O_4$ (Hodgkinson a Sorby, *J* 1876, 936) Black powder Not affected by dilute acids and alkalis

Sepiatic acid C 56.3, H 3.6, N 12.3, S 5, O 27.2 Obtained by digesting the pigment from the ink bag of *sepiæ* with 15 pts of 10 p c potash Sol alkalis, ppd from ammoniacal solution by ammonia zinc chloride or copper sulphate (Nencki a Sieber)

MELANTHIN $C_{20}H_{14}O$, [205°] Occurs in the seeds of *Nigella sativa* (Greenish, *Ph* [3] 10, 909, 1013) Prepared by extracting the seeds with alcohol, evaporating the extract, dissolving the pp in alcohol, and fractionally ppg with water Minute grey crystals (from alcohol) Insol water, benzene, CS₂, and light petroleum, v sol alcohol, sol alkalis, sl sol chloroform Conc H₂SO₄ gives a red colouration H₂SO₄ and sugar give a violet blue colour Boiling dilute HCl splits it up into a sugar and melanthigenin $C_{11}H_{12}O_2$, which forms minute crystals, sl sol water

MELANURENIC ACID v AMMELIDE

MELEM v CYANIC ACIDS

MELENE $C_{12}H_{10}$, [62°] (370°-380°) S G 89 VD 10-11.8 S (alcohol) 13 in the cold, 3.6 at 78° An olefine (?) produced by the dry distillation of bees' wax (Ettling, *A* 2, 252, Lewy, *A Ch* [3] 5, 395, Brodie, *A* 71, 156) — White nacreous plates (from ether) Insol water, sl sol cold alcohol, v sol ether Not attacked by cold H₂SO₄, slightly attacked by boiling HNO₃ Attacked by chlorine

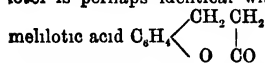
MELEZITOSE $C_{12}H_{20}O_{11}$, [148°] $[\alpha]_D = 94^\circ$ (B), 88.5° (V) A sugar discovered by Bonastre in the manna of Briançon which exudes from the young branches of 'mélèze' (*Pinus Larix*) (Berthelot, *A Ch* [3] 65, 282) It occurs also in 'taranjoline' or Turkestan manna (Villiers, *Bl* [2] 27, 98, Alekhine, *Bl* [2] 46, 824) Obtained by extracting the Briançon manna with boiling alcohol, and evaporating After a few weeks melezitose separates, it is then recrystallised from alcohol Monoclinic crystals containing aq (V), or anhydrous crystals (A) About as sweet as glucose Dextrorotatory V sol water, nearly insol cold, sl sol boiling, alcohol, insol ether Its aqueous solution is ppd by ammoniacal lead acetate Boiling dilute H₂SO₄ forms glucose It is turned brown by alkalis.

It does not reduce Fehling's solution Does not ferment with yeast It forms a compound with phenyl hydrazine It gives an octo acetyl derivative.

MELIDO-ACETIC ACID $C_8H_5N_2O_2$ *18* $(CN)_2N_2H_2CH_2CO_2H$ Formed by treating cyanamide with chloroacetic ether and NaOEt (Drechsel, *J pr* [2] 11, 332) Amorphous mass, but obtained in a crystalline state by ppg its ammonium salt with HCl Decomposed by heat without previous fusion Insol alcohol and ether, v sl sol cold water Sol aqueous HClAq, baryta, and KOHAq, but insol NH₄Aq — HA'HCl needles, sl sol cold water — HA'HNO₃Aq — AgA'HNO₃Aq — (HA')₂H₂SO₄ 4aq

MELILOTIC ACID *v* OXY PHENYL PROPIONIC ACID

MELILOTOL $C_{10}H_{10}O_2$ (?) An oil found, together with coumarin, in the yellow melilot (*Melilotus officinalis*) Extracted by distilling the plant, when in flower, with steam, and extracting the distillate with ether (Phipson, *C N* 32, 25, *C R* 86, 830) Liquid, v sl sol water, alcohol, and ether Has an agreeable odour When boiled with conc KOHAq it yields oxy phenyl propionic (mehlotic) acid $C_8H_{10}O_2$ Melilot is perhaps identical with the lactone of



MELINOIN - TRI - SULPHONIC ACID $C_8H_5O_3(SO_3H)_3$

Formation — 1 By warming a mixture of *p* oxy benzoic aldehyde, (8) naphthol, and H₂SO₄ 2 From benzoic aldehyde, (8) naphthol, and H₂SO₄ — 3 By heating Baeyer's condensation product $C_8H_5O_2$ (from benzoic aldehyde and (8) naphthol) with H₂SO₄ (Trzciński, *B* 16, 2835, 17, 500) — Yellow crystalline powder Insol absolute alcohol, tolerably sol water, the dilute solution being of a rose red colour with a green fluorescence It dissolves in strong H₂SO₄ or HNO₃ with a splendid green fluorescence, and is not attacked even on boiling It forms unstable compounds with HCl and H₂SO₄, although it is itself a strong acid

Salts — A'''K, easily soluble fine colourless needles — A'''Ca, aq colourless soluble crystals — A'''Ba, sparingly soluble white amorphous pp or microscopic needles

MELISSIC ACID $C_{10}H_{16}O_2$ *18* $C_{10}H_{16}CO_2H$, or $C_7H_{12}O_2$ *18* $C_3H_7CO_2H$ [90°] Occurs in bees' wax (Natzger, *A* 224, 225) Formed by heating the myricyl alcohol of bees' wax with soda lime at 270°-300° in absence of air (Brodie, *A* 71, 156, Von Peverling, *A* 183, 344, Sturcke, *A* 223, 295, Schwalb, *A* 235, 106) Silky scales, composed of minute needles Sol alcohol, light petroleum, chloroform, and CS₂, sl sol ether According to Schalleff (*B* 12, 697) melissic acid is a mixture of acids

Salts — PbA₂ [119°] Insol alcohol and ether, sl sol boiling toluene, chloroform, and glacial acetic acid — MgA₂ — CaA₂ — AgA₂ [95°], amorphous pp

Methyl ether MeA' [71°] Needles (from lignin)

Ethyl ether EtA' [70°] (Schwalb), [73°] (P) Needles (from alcohol) Split up on heating into ethylene and the free acid

Isoamyl ether C₅H₁₁A' [69°].

MELISSYL ALCOHOL *v* MYRICYL ALCOHOL.

MELITOSE *v* RAFFINOSE and SUGARS

MELLITIC ACID $C_{12}H_6O_{12}$ *18* $C_6(CO_2H)_6$

Mellitic acid Benzene hexa-carboxylic acid. Mol w 342 H G v 790,800 H C p 788,200 H F 546,800 (Stohmann, Kleber, a Langbein, *J pr* [2] 40, 141) Heat of neutralisation 84,034 (Gal a Werner, *Bl* [2] 47, 162)

Occurrence — As aluminium mellitate in honey stone or mellite (Klaproth, *A D* 1799, Baeyer, *A Suppl* 6, 1)

Formation — 1 By the oxidation of hexamethyl benzene by cold alkaline KMnO₄ (Friedel a Kratts, *A Ch* [6] 1, 470) — 2 By the oxidation of charcoal by KMnO₄ (Schulze, *B* 4, 802, 806) — 3 By the electrolysis of acidified water or of KOH using a positive electrode of gas carbon (Bartoli a Papasogli, *G* 11, 468, *C C* 1831, 327) — 4 In very small quantity by oxidising coal, animal charcoal, or lampblack by alkaline NaOCl (Bartoli a Papasogli, *G* 15, 446)

Preparation — Powdered honey stone is boiled with ammonium carbonate, ammonia is added, and the filtrate evaporated to crystallisation The ammonium mellitate is recrystallised with addition of a little ammonia, dissolved in water, ppd by lead acetate, and the pp decomposed by H₂S (Wohler, *A* 37, 263, Schwarz, *A* 66, 47, Claus, *B* 10, 559)

Properties — Stellate groups of delicate silky needles (from alcohol), v sol water and alcohol Decomposed by heat into CO and pyromellitic acid $C_{10}H_4O_8$ (Erdmann, *J pr* 52, 432), which solidifies in the neck of the retort as a radio-crystalline mass When strongly heated with glycerin it yields CO₂ and trimelic acid $C_6H_4O_6$ When heated to redness with soda lime it yields benzene (Baeyer, *A Suppl* 7, 5) Not decomposed by boiling HNO₃, sulphuric acid, HIAq, chlorine, or bromine When its aqueous solution is electrolysed, CO₂, hydrogen, CO, and oxygen are given off (Bunge, *J R* 12, 421)

Salts — (NH₄)₂C₁₂O₁₂ 3aq trimetric crystals (Schwarz, *A* 66, 47) Decomposed at 150° with formation of paramide and ammonium euchoate — (NH₄)₂H₂C₁₂O₁₂ 4aq trimetric prisms From ammonio cupric mellitate and H₂S — K₂A^{VI} 9aq trimetric crystals — K₂H₂A^{VI} 8aq — K₂H₂A^{VI} 6aq large prisms, more soluble in water than the normal potassium salt — (K₂H₂A^{VI})(KNO₃)₂ 9aq (?) — Na₂A^{VI} 12aq needles (from a hot conc solution) — Na₂A^{VI} 18aq large striated triclinic crystals — Ba₂A^{VI} 3aq white gelatinous pp, changing to scales — Mg₂A^{VI} 18aq crystalline mass — Mg(NH₄)₂A^{VI} 15aq large glassy prisms — Zn₂A^{VI} 15aq — Zn₂A^{VI} 9aq — Mn₂A^{VI} 18aq minute needles, more sol cold than hot water S (hot water) 125 — Co₂A^{VI} 18aq minute prisms (from boiling water) — Ni₂A^{VI} 24aq v sl sol water — Al₂A^{VI} 18aq *Mellite* S G 16 SH 336 (Bartoli, *G* 14, 105) Occurs in lignite at Asten in Thuringia, Bilin in Bohemia, and near Walschau in Moravia Massive nodules Dimetric crystals *a c* = 1 745 — Fe₂A^{VI}Fe₂O₉ 9aq minute lemon yellow crystals, v sl sol water — Cu₂A^{VI} 12aq From boiling mellitic acid and cupric acetate Amorphous pp, becoming crystalline — Cu₂H₂A^{VI} 12aq — Cu₂(NH₄)₂A^{VI} 12aq from cupric sulphate and ammonium mellitate. — Hg₂A^{VI} 6aq (dried at 100°), granular mass —

$\text{Hg}_2\text{A}^{\text{VI}}$ 6aq (dried at 100°). granular pp — $\text{Pb}_2\text{A}^{\text{VI}}$ (dried at 180°) bulky white pp — $\text{Pd}_2\text{A}^{\text{VI}}(\text{NH}_3)_2$ 6aq colourless trimetric crystals (Karmrod, *A* 81, 164) — $\text{Ag}_2\text{A}^{\text{VI}}$ white crystal line powder (Wöhler, *A* 30, 1)

Methyl ether $\text{Me}_2\text{A}^{\text{VI}}$ [187°] From silver mellitate and MeI (Kraut, *J* 1862, 281, *A* 177, 273) Plates H C 1,825,600 HF 487,400 (Stohmann, *J pr* [2] 40, 350)

Ethyl ether $\text{Et}_2\text{A}^{\text{VI}}$ [73°]

Isoamyl ether $(\text{C}_4\text{H}_9)_2\text{A}^{\text{VI}}$ Oil

Chloride $\text{C}_6(\text{COCl})_2$ [190°] (Claus, *B* 10, 561) Formed, together with an oxychloride $\text{C}_6\text{O}_2\text{Cl}_2$, from PCl_5 and mellitic acid. Hard vitreous prisms (from ether) Sublimes in laminae at 240° . Slowly decomposed by water into HCl and mellitic acid.

Tri imide $\text{C}_{12}\text{H}_2\text{N}_2\text{O}_8$ *se*

$\text{C}_6\left\{\begin{array}{c} \text{CO} \\ \text{CO} \end{array}\right\} \text{NH}$ *se* *Paramide* Formed, together with euchroic acid, by heating ammonium mellitate at 160° as long as NH_3 escapes. Water extracts ammonium euchroate from the residue leaving paramide undissolved (Wöhler, *A* 37, 268, Schwarz, *A* 66, 52). White amorphous powder, insol water and alcohol. Sol H_2SO_4 , but reprecipitated by water. Boiling with water slowly converts it into $(\text{NH}_4)_2\text{H}_2\text{C}_6\text{O}_{12}$. Alkalis convert it first into euchroic and then into mellitic acid. AgNO_3 forms in its ammoniacal solution a gelatinous pp which gives off NH_3 when heated, leaving $\text{Ag}_2\text{N}_2\text{C}_{12}\text{O}_{12}$. An ammoniacal solution of paramide dropped into HClAq gives a white crystalline pp called 'paramic acid' $\text{C}_{12}\text{H}_2\text{N}_2\text{O}_8$ (?) or

$\text{CO}(\text{NH}_2)_2 > \text{C}_6\left\{\begin{array}{c} \text{CO} \\ \text{CO} \end{array}\right\} \text{NH}$ *se*

Tri phenyl tri imide

$\text{C}_6\left\{\begin{array}{c} \text{CO} \\ \text{CO} \end{array}\right\} \text{NPh}$ *se* Formed by heating mellitic acid or ammonium mellitate (1 mol) with aniline (6 mols) at 160° for 7 hours (Hotte, *J pr* [2] 32, 238). White amorphous mass

Di imide $\text{C}_{12}\text{H}_2\text{N}_2\text{O}_8$ *se*

$(\text{CO}_2\text{H})_2\text{C}_6\left\{\begin{array}{c} \text{CO} \\ \text{CO} \end{array}\right\} \text{NH}$ *se* *Euchroic acid*. Prepared by heating ammonium mellitate at 160° for several hours, dissolving in water, and adding HCl to the hot solution. The crystals which separate on cooling are recrystallised from water (Wöhler, *P* 52, 610, Schwarz, *A* 66, 49). Small prisms (containing 2aq), al sol cold water. Strongly acid. Melts about 280° . Euchroic acid is not decomposed by boiling water or HClAq . When a solution of euchroic acid is treated with zinc the metal becomes covered with a dark blue deposit of 'euchrone'. This deposit detaches itself when the zinc is immersed in dilute HClAq . After drying, euchrone is a black mass which becomes colourless when heated in air, being re-oxidised to euchroic acid. Euchrone forms a deep purple solution in NH_4Aq or KOH Aq , but the solution is quickly decolourised by the air. When a solution of euchroic acid is electrolysed, the negative platinum becomes coated with the blue deposit.

Salts — $(\text{NH}_4)_2\text{C}_{12}\text{H}_2\text{N}_2\text{O}_8$ white crusts — An acid ammonium salt separates from hot solutions in yellowish crystals — BA^{VI} 6aq pale-yellow powder — PbA^{VI} 6aq — PbA^{VI} 6aq — $\text{Ag}_2\text{A}^{\text{VI}}$ sulphur-yellow powder.

Hexahydrate $\text{C}_6\text{H}_4(\text{CO}_2\text{H})_6$ *Hydromellitic acid* Formed by treating an ammoniacal solution of mellitic acid with sodium amalgam, neutralising with acetic acid, ppg by lead acetate, and decomposing the pp with H_2S (Baeyer, *A Suppl* 7, 16). Formed also in the electrolysis of aqueous KOH with a positive electrode of carbon (Bartoli a Papasogli, *C* 1881, 327). Hygroscopic syrup, v e sol water. By heating with H_2SO_4 (5 pts) there is formed, together with CO_2 and SO_2 , trimelic acid $\text{C}_6\text{H}_4(\text{CO}_2\text{H})_3$, prehnitic acid $\text{C}_6\text{H}_2(\text{CO}_2\text{H})_4$, mellophanic acid $\text{C}_6\text{H}_2(\text{CO}_2\text{H})_4$, and prehnomalic acid $\text{C}_{10}\text{H}_8\text{O}_8$.

Salts — The alkaline salts are gummy, v e sol water — The calcium salt is more soluble in hot than in cold water — $\text{Pb}_2\text{A}^{\text{VI}}$ (dried at 150°) Amorphous pp — $\text{Ag}_2\text{A}^{\text{VI}}$ amorphous pp

Isohexahydrate $\text{C}_6\text{H}_4(\text{CO}_2\text{H})_6$ *Isohydromellitic acid* Formed by heating the preceding hexahydrate with conc HClAq for 3 hours at 180° (Baeyer) Prisms (from water) v sol water, but ppd on addition of HCl . Not affected by heating with fuming HClAq at 300° . Decomposed in the same manner as its isomeride when heated with H_2SO_4 . Chromic acid mixture oxidises it to acetic acid and CO_2 . — $\text{Pb}_2\text{A}^{\text{VI}}$

Methyl ether of the Isohexahydrate $\text{Me}_2\text{C}_{12}\text{H}_2\text{O}_{12}$ [125°] Needles, v e sol also hol

References — HEMI MELLITIC ACID, PYROMELLITIC ACID

TRIMELLITIC ACID $\text{C}_6\text{H}_2\text{O}_8$ *se*

$\text{C}_6\text{H}_2(\text{CO}_2\text{H})_3$ [1 2 4] *Benzene tricarboxylic acid* Mol w 210 [218°]

Formation — 1 Together with isophthalic acid and pyromellitic anhydride by heating the tetrahydride of pyromellitic acid $\text{C}_6\text{H}_4(\text{CO}_2\text{H})_4$ with 5 pts of H_2SO_4 (Baeyer, *A Suppl* 7, 40) — 2 By oxidising the acid $\text{C}_6\text{H}_4\text{Me}(\text{CO}_2\text{H})_2$, prepared from ψ cumene, by KMnO_4 (Krinno, *B* 10, 1491) — 3 By oxidising alizarin carboxylic acid with dilute HNO_3 (Hammerschlag, *B* 11, 88) — 4 By oxidising the acid $\text{C}_6\text{H}_2\text{Me}(\text{C}_6\text{H}_4)\text{CO}_2\text{H}$ with dilute nitric acid (S G 1 12) at 240° (Effront, *B* 17, 2338) — 5 By the action of cuprous cyanide upon the diazo compound from amido terephthalic acid, and saponification of the product with aqueous KOH (Ahrens, *B* 19, 1634) — 6 By heating mono potassic (8) sulphophthalate with sodium formate, 1-ophthalic acid being also formed (Graebe a Rce, *C J* 49, 532)

Preparation — 100 g pulverised colophony and 2 litres of nitric acid (1 vol commercial acid to 2 vols water) are introduced into a retort with its beak directed upwards, and the liquid is heated to boiling, whereupon the colophony melts, and is quickly attacked, with copious evolution of red vapours, and converted into an orange yellow viscid mass. After the boiling has been continued for 6 or 8 hours, fresh resin and strong nitric acid are added alternately, till about 1 kg resin has been introduced, the contents of the retort being kept in constant ebullition, this process takes about a fortnight. The resulting wine yellow liquid, which on addition of water should merely show turbidity, and not yield any lumpy precipitate, is then distilled off till the remaining liquid begins to froth, and

this liquid, while still hot, is poured into a ten fold volume of cold water, decanted after twenty-four hours from the varnish like resin which separates, and evaporated to a syrupy consistence. The whole then solidifies after a while to a thick crystalline pulp, which may be freed from the syrupy mother-liquor by means of an aspirator and porous earthenware plates, and afterwards boiled with water. The extract on cooling deposits isophthalic acid in slender needles, and on further concentration, first a little more of the same acid and then trimellitic acid, which may be purified by solution in alcohol and crystallisation from water. 30 g of trimellitic acid may be thus obtained from 50 g of resin (Schreder, *A* 172, 93).

Properties—Rosettes of small needles, m sol water and ether. When heated above its melting point the anhydride distils over, condensing in concentrically arranged groups of needles. On fusion with NaOH it gives benzene and diphenyl (Barth & Schreder, *B* 12, 1257). It also yields benzene when distilled with lime.

Salts— $\text{Ba}_2\text{A}'''$, 4aq (when air dried). Gives off 3aq at 160° . Sl sol water— $\text{Ba}_2\text{A}'''$, 3aq (when dried over H_2SO_4)— $\text{Ag}_2\text{A}'''$ white pp, not much affected by light. Sl sol water.

Anhydride $\text{C}_6\text{H}_2(\text{CO}_2\text{H})_2 \xrightarrow{\text{CO}} \text{CO}_2\text{O}$ [158°]

Formed by heating the acid (Baeyer, *A* 166, 340). Crystalline mass, sl sol cold, v sol hot, water.

Reference—SULPHO TRIMELLITIC ACID

MELLITYL ALCOHOL v **PENTA METHYL BENZYL ALCOHOL**

MELLOGEN $\text{C}_{11}\text{H}_{10}\text{O}_4$. On the electrolysis of water by a battery of 1,200 Daniell's cells, the anode being gas carbon, the liquid becomes of an inky colour, and there is deposited in the voltameter a black residue. This is washed with water until the filtrate is quite colourless, from the washings the mellogen is ppd with a small quantity of dilute hydrochloric acid (Bartoli & Papasogli, *G* 11, 468, 12, 117, 13, 37, 15, 464, *C* R 94, 1839, *A* Ch [6] 7, 364). A black solid, of conchoidal fracture, sol water and alkalis, insol alcohols and hydrocarbon, and in most mineral acids and salts. It is not readily combustible, and is converted by oxidising agents, e.g. NaOCl, into mellitic and pyromellitic acid and their hydrides. At ordinary temperatures it has the composition $2\text{C}_{11}\text{H}_{10}\text{O}_4 + 3\text{H}_2\text{O}$, at 100° it may be represented by $2\text{C}_{11}\text{H}_{10}\text{O}_4 \cdot \text{H}_2\text{O}$. The aqueous solution of mellogen is dark coloured, and is ppd by acids and by many salts. With baryta mellogen gives an insoluble pp. Nitric acid (8 G 136) oxidises mellogen forming an amorphous compound $\text{C}_{11}\text{H}_8\text{O}_5$, sol water, alcohol, and ether, a soluble acid $\text{C}_{11}\text{H}_8\text{O}_6$, 2aq, forming the salts $\text{Ba}_2(\text{C}_{11}\text{H}_8\text{O}_6)_2$ and $\text{Ag}_2\text{C}_{11}\text{H}_8\text{O}_6$, and a black compound resembling mellogen.

MELLONE v **CYANIC ACIDS**

MELLOPHANIC ACID $\text{C}_{12}\text{H}_6\text{O}_8$, $\text{C}_{12}\text{H}_6\text{O}_8$, $\text{C}_{12}\text{H}_6\text{O}_8$ [1 2 3 5]. *Benzene t-tetra carboxylic acid* [238°]. Obtained, together with prehnitic acid $\text{C}_6\text{H}_2(\text{CO}_2\text{H})_4$ [1 2 3 4], by heating the hydride of pyromellitic acid $\text{C}_6\text{H}_2(\text{CO}_2\text{H})_4$ [1 2 4 5] with H_2SO_4 (Baeyer, *A* 166, 327). Formed also by the oxidation of iso-durene $\text{C}_{12}\text{H}_{16}\text{Me}_2$ (Jacobson, *B* 17, 2516). Small four-sided prisms, v sol water, but ppd by HCl from a conc. solu-

tion. Converted into an anhydride on fusion. Calcium acetate gives in hot solutions of the acid a pp which re-dissolves on cooling. Baryta-water gives a pp which becomes crystalline on heating. Lead acetate gives a flocculent pp, insol acetic acid. Sodium amalgam forms a hydride of mellophanic acid.

Anhydride [238°]. Formed by fusing the acid. Insol ether.

MELOLONTHIN $\text{C}_{12}\text{H}_{12}\text{N}_2\text{SO}_4$. Found, together with leucine and hypoxanthine, in cockchafer (*Melolontha vulgaris*). The insects are extracted with water, the extract boiled, ppd with lead subacetate, and the filtrate freed from lead by H_2S and evaporated. The crystals are freed from leucine by treatment with alcohol (Schreiner, *B* 4, 763). 30lbs of cockchafer yield 1.5 g melolonthin. Silky needles (from water), sl sol cold, m sol hot, water, v sl sol dilute alcohol and ether, v sol acids and alkalis. Does not act on light.

MELMESIDINE v **DI MESITYL GUANIDINE**

MENAPHTHYLAMINE $\text{C}_{11}\text{H}_9\text{N}_2$

$\text{C}_{10}\text{H}_7\text{CH}_2\text{NH}_2$. (292°). Formed by treating the amide of (a)-thionaphthoic acid $\text{C}_{10}\text{H}_7\text{CSNH}_2$ with zinc and HCl (Hofmann, *B* 1, 100). Alkaline liquid, readily absorbing CO_2 from the air— B^+HCl long, sparingly soluble crystals— $\text{B}^+\text{H}_2\text{PtCl}_6$ yellow crystalline pp.

MENISPERMINE $\text{C}_{18}\text{H}_{24}\text{N}_2\text{O}_2$ (?) [120°]

Occurs in the seeds of *Menispermum cocculus* (Pelletier & Caventon, *A* Ch [2] 54, 178). To extract the menispermine, the alcoholic extract of the seeds is first treated with cold water, then exhausted with hot acidulated water, the brown solution is ppd by an alkali, and the pp is exhausted with very weak acetic acid, which leaves a brown black mass undissolved. Or the seeds may be made up into a heap, and exhausted with alcohol of specific gravity 0.833, the alcohol distilled off, the residue boiled with water, and the liquid filtered at the boiling heat, it then, on cooling, deposits crystals of picrotoxine, especially if a small quantity of acid has been added. The part insol boiling water is then treated with acidulated water, and ppd by an alkali, a granular pp is thereby formed, from which alcohol extracts a peculiar yellow substance, and the residue is finally dissolved in ether, which deposits menispermine in the crystalline form. The ether leaves undissolved a viscous substance, which may be dissolved in absolute alcohol and the solution, evaporated at 45° , ultimately yields crystals of paramenispermine.

Properties—Prisms, insol water, sol alcohol and ether. Is not poisonous. Its sulphate forms prismatic needles, sol water.

Paramenispermine [250°]. Isomeric with menispermine. Insol water, sl sol ether, v sol boiling alcohol. Dissolves in dilute acids.

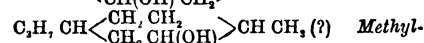
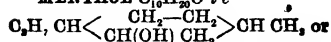
MENTHENE $\text{C}_{10}\text{H}_{16}$. Mol w 138 (167.4°) (Atkinson & Yoshida, *C* J 41, 53), (163°) (Oppenheim, *C* J 15, 29). VD 4.94 (calc 4.78). SG 1.814, $d_{20}^{20} 0.8226$, $d_{20}^{20} 0.8073$ (A & Y). CE (0° – 10°) 0.00994, (0° – 60°) 0.0100 [α_D] = +13.25 (A & Y). $R_D 74.0$ Specific refractive energy, 548. Specific dispersive energy, 0313 (Gladstone, *C* J 49, 622). Obtained by distilling menthol (1 pt) with ZnCl_2 (2 pts), or with P_2O_5 (Walter, *A* 32, 288). Formed also by the action

of conc H_2SO_4 on menthol (Beckmann, *A* 250, 358)

Properties—Colourless oil, smelling like cymene, m sol ether and alcohol, v sol benzene, turpentine, and petroleum Dextrorotatory

Reactions—1 Combines with fuming HCl , forming a yellow oil $\text{C}_{10}\text{H}_{19}\text{Cl}$ Combines also with HI , forming $\text{C}_{10}\text{H}_{21}\text{I}$, identical with the product obtained by heating terpinene dihydride $\text{C}_{10}\text{H}_{16}2\text{HI}$ with HI (Bouchardat & Lafont, *C R* 107, 916)—2 Combines with bromine, forming $\text{C}_{10}\text{H}_{18}\text{Br}_2$, which is split up by heat into HBr and cymene (Beckett & Wright, *C J* 29, 1) 3 Fuming nitric acid oxidises it to glutaric acid

MENTHOL $\text{C}_{10}\text{H}_{18}\text{O}$



propyl phenol hexahydride Mol w 156 $[\alpha]_D^{20}$ (212°) VD 5 62 (calc 5 41) SG $\frac{1}{2}$ 890 $[\alpha]_D^{20}$ = -59.3 (Moriya), $[\alpha]_D^{20}$ = -59.6 (Oppenheim), = -49.4 in a 5 p.c. alcoholic solution at 22° (Arth, *A Ch* [6] 7, 438) R_D^{20} = 77.4 in a 21 p.c. benzene solution (Kanonnikoff, *J pr* [2] 31, 348) HC 1,509,100 (Lougounine, *A Ch* [5] 23, 387) Deposited in crystals when the essential oil of peppermint is kept for a long time or cooled to a low temperature (Dumas, *A* 6, 252, Blanchet & Sell, *A* 6, 293; Walter, *A* 32, 288; Kane, *P M* 16, 418; Laurent, *Rev Sci* 14, 341; Oppenheim, *C J* 15, 24) Menthol is imported from Japan in the solid state as 'peppermint camphor' (Moriya, *C J* 39, 77) White crystals, smelling of peppermint, sl sol water, v sol alcohol, ether, CS_2 , HOAc , and fixed and volatile oils Insol aqueous alkalis From alcoholic NaOH it crystallises in long needles Sodium dissolves in it with evolution of hydrogen Conc acids dissolve menthol, but it is reppd on dilution with water Menthol has the refractive and dispersive energy of a saturated compound (Gladstone, *C J* 49, 621) The rate of etherification of menthol by acetic acid is that of a secondary alcohol (Menschutkin, *J R* 13, 569)

Reactions—1 With $\text{K}_2\text{Cr}_2\text{O}_7$ and H_2SO_4 in sealed tubes at 120° gives an inactive oil 'menthone' (205°), $\text{C}_{10}\text{H}_{18}\text{O}$, SG $\frac{1}{2}$ 9032, sol alcohol, ether, and chloroform—2 Fuming HNO_3 forms an explosive oil, SG $\frac{1}{2}$ 1 061, which may be reduced to a yellow oil $\text{C}_{10}\text{H}_{19}\text{NH}_2$ (185°–190°) 3 Excess of fuming HNO_3 forms an acid, probably glutaric (*q v*)—4 Bromine in acetic acid forms $\text{C}_{10}\text{H}_{17}\text{Br}$ Unstable oil—5 Conc HCl aq at 100° slowly forms menthyl chloride PCl_5 forms the same body—6 ZnCl_2 or P_2O_5 dehydrate menthol, producing menthene—7 KMnO_4 in acid solution forms (8) pimelic acid $\text{C}_8\text{H}_{12}\text{O}_4$, [87°], of which the amide [191°] crystallises from water in prisms (Arth, *A Ch* [6] 7, 440, *C R* 107, 107) Formic, propionic, butyric, and oxymenthyl acids are formed at the same time The oxymenthyl acid $\text{C}_{10}\text{H}_{18}\text{O}_4$ boils at 174° under 15 mm pressure—8 Boiled with HI (SG 1.7) for 3 days it forms a mixture of hydrocarbons chiefly consisting of $\text{C}_{10}\text{H}_{16}$ (168.6°) VD 67.25, $[\alpha]_D^{20}$ = +5.2 SG $\frac{1}{2}$ 8254, $\frac{1}{2}$ 8111, CE (0°–10°) 000929, (0°–60°) 000694 R_D^{20} 78.28 Colourless oil, v sol petroleum or

benzene, less so in alcohol or ether Odour resembles cymene (Atkinson & Yoshida, *C J* 41, 54)—9 Menthol does not react with hydroxylamine (Nageli, *B* 16, 499)—10 On heating with sodium and ether on the water bath, and, after cooling, adding CS_2 , there is formed menthyl-xanthogenic acid $\text{C}_{10}\text{H}_9\text{O CS SH}$, a liquid which undergoes spontaneous decomposition, but forms a yellow crystalline copper salt CuA_2 (Bamberger & Lodter, *B* 23, 213)

Acetyl derivative $\text{C}_{10}\text{H}_{17}\text{OAc}$ *Menthyl acetate* (223°) $[\alpha]_D^{20}$ = -114° From menthol and HOAc at 120° (Oppenheim, *C J* 15, 26) Liquid Not decomposed by cold alkalis, but saponified by alcoholic NaOH at 120°

Butyryl derivative $\text{C}_{10}\text{H}_{17}\text{O CO C}_3\text{H}_7$, *Menthyl butyrate* Obtained by heating menthol with butyric acid at 200° for 30 hours (Oppenheim) (230°–240°) $[\alpha]_D^{20}$ = -89°

Benzoyl derivative $\text{C}_{10}\text{H}_{17}\text{OBz}$ [54°] (230°) $[\alpha]_D^{20}$ = -91° In a one p.c. benzene solution, trimetric crystals partially decomposed on distillation into menthene and benzoic acid (Arth, *A Ch* [6] 7, 479)

Carbamyl derivative $\text{C}_{10}\text{H}_{17}\text{NO}$, *Menthyl carbamate* [165°] $[\alpha]_D^{20}$ = -85° in a 58 p.c. chloroform solution at 21° Formed by passing dry cyanogen into a solution of sodium menthol in dry toluene The product is washed with water and the toluene distilled off The crystalline mass is then recrystallised from alcohol, from which the carbamyl derivative separates in slender prismatic needles (Arth, *C R* 94, 872, 98, 521, *A Ch* [6] 7, 433) Needles, m sol alcohol and benzene Sublimes at 100° Decomposes at about 200°, yielding cyanuric acid Alcoholic KOH forms menthol and potassium cyanate Ac_2O at 130° forms $\text{C}_{10}\text{H}_{17}\text{OAc}$ Benzoic aldehyde and HCl form $(\text{C}_{10}\text{H}_{17}\text{OCO NH})_2\text{CHPh}$ [143°]

Carbonyl derivative $(\text{C}_{10}\text{H}_{17}\text{O})_2\text{CO}$ *Menthyl carbonate* [105°] Obtained in the preparation of the carbamyl derivative by evaporating the alcoholic mother liquor and boiling the residue with water as long as menthol and ammonia come off On cooling, the carbonyl derivative is deposited as a crystalline mass, sol boiling alcohol (Arth) Plates (from alcohol) or prisms (from toluene), v sl sol alcohol, v sol ether and benzene Alcoholic KOH at 100° saponifies it

Phenyl-carbamyl derivative $\text{C}_{10}\text{H}_{17}\text{O CO NHPh}$ *Menthyl phenyl urethane* *Menthyl phenyl carbamate* [111°] From menthol and phenyl cyanate (Leuckart, *B* 20, 116) Silky needles (from hot alcohol)

Succinoyl derivative $\text{C}_{10}\text{H}_{17}\text{O CO CH}_2\text{CH}_2\text{CO}_2\text{H}$ [62°] $[\alpha]_D^{20}$ = -59.6° in a 1.4 p.c. benzene solution Formed by heating menthene (1 mol) with succinic anhydride (1 mol) at 110° (Arth) Acicular crystals, v sol alcohol, v sl sol hot water Its Na and K salts are very deliquescent, the Ca and Ba salts are white pps— $\text{AgC}_{10}\text{H}_{17}\text{O}$, small needles (from hot water)

Succinyl derivative $(\text{C}_{10}\text{H}_{17})_2\text{C}_2\text{H}_2\text{O}_4$ [62°] (220°) $[\alpha]_D^{20}$ = -81.5° in a 1.9 p.c. benzene solution From menthol (1 mol) and succinic acid (2 mols) at 150° Trimetric crystals (from alcohol) Decomposes on distillation into succinic acid and menthene.

Phthaloxyl derivative $C_{18}H_{24}O_4$, *tc* $C_{18}H_{18}O_4$ $COO C_6H_4 CO_2H$ [110°] $[\alpha]_D = -105.6^\circ$ in a 16 p c benzene solution. Formed by heating menthol (1 mol) with phthalic anhydride (1 mol) at 110° (Arth). Minute needles, insol cold water, v sol alcohol and ether. Gives white pps with Ba and Ca salts $-Mg(C_{18}H_{24}O_4)$ laminae, almost insol cold water.

Phthalyl derivative $C_{18}H_{20}O_4$, *tc* $(C_{18}H_{18})_2C_6H_4O_2$ [133°] $[\alpha]_D = -94.7^\circ$ in a 2 p c benzene solution at 20°. From phthalic anhydride (1 mol) and menthene (2 mols) at 140°. Trimetric crystals (from ether), sl sol boiling alcohol.

Menthyl chloride $C_{10}H_{17}Cl$ (c 204°). Obtained by treating menthol with PCl_5 (Walter, *A* 82, 292) or with conc $HClAq$ for a week at 100° (Oppenheim). The same body appears to be formed by heating menthene with fuming $HClAq$ at 120° (Arth, *C R* 97, 323). Light oil, sl sol water, m sol alcohol. Not attacked by alcoholic KOH . With sodium menthol it gives menthene and menthol.

Menthyl bromide $C_{10}H_{17}Br$. From menthol and bromide of phosphorus. Non volatile liquid. Gives with bromine $C_{10}H_{17}Br_2$.

Menthyl iodide $C_{10}H_{17}I$. From menthol and conc $HI Aq$. Liquid, decomposed by alcoholic K_2S into menthene and HI .

Tetra menthyl silicate $Si(OC_{10}H_{19})_4$. [82°] (350°) at 155 mm. Formed by the action of $SiCl_4$ upon menthol, the yield is 75 p c of the theoretical (Hertkorn, *B* 18, 1695). Colourless prisms, v sol ether, benzene, ligroin, chloroform, CS_2 , and warm alcohol, sl sol cold alcohol.

Oxymenthyllic acid $C_{16}H_{24}O_4$ (175°) at 15 mm (280°). A product of the oxidation of menthol by $KMnO_4$ (Arth, *A Ch* [6] 7, 448). Colourless liquid, partially decomposed on boiling at 280°, v sl sol water, v sol alcohol and ether. The salts of alkalis and alkaline earths are very soluble. A solution of the sodium salt is ppd by salts of Pb , Fe''' , Cu , and Pt . It forms no acetyl derivative $-AgA'$ $-NaA'$ white deliquescent crystals.

Methyl ether MeA' (137°) at 17 mm. Liquid.

Ethyl ether EtA' (145°) at 18 mm. Thick liquid.

MENTHONE $C_{10}H_{18}O$ (206.3° cor) $[\alpha]_D = +21^\circ$ S.G. $\frac{4}{20}$ 9126, $\frac{4}{20}$ 8972 C.E. (0°-10°) 000862, (0°-100°) 000923 R_∞ 75.3. From menthol (30 g), $K_2Cr_2O_7$ (10 g), and H_2SO_4 (10 g) at 135°. The light oil is subjected several times to the action of the same oxidising mixture (Atkinson & Yoshida, *C J* 41, 49). Colourless oil, miscible with alcohol, chloroform, benzene, and CS_2 . Smells like diluted peppermint. Does not combine with $NaHSO_4$. If a solution in petroleum is treated with Na and CO_2 successively and then shaken with water crystals of menthol are got [42°], but having $[\alpha]_D = -39^\circ$ not -59° .

Lævo-menthone $C_{10}H_{18}O$ *tc*

$CHP < \begin{smallmatrix} CH_2 & CH_2 \\ CH_2 & CO \end{smallmatrix} > CHMe (?)$ (207°) S.G. $\frac{4}{20}$ 8960 $[\alpha]_D = -28.2^\circ$. Menthone is a mixture of two isomerides—one dextro, the other lævo-rotatory. By using very little acid the lævo-menthone may be obtained free from dextro-menthone and menthol. The best proportions are 60 g. $K_2Cr_2O_7$, 50 g. H_2SO_4 , 300 g. water, and

45 g. menthol, the mixture being kept at 80° to 55° (Beckmann, *A* 250, 325). The menthone is extracted with ether, washed with dilute alkalis, and distilled with steam.

Properties—Mobile liquid, smelling faintly of peppermint, sl sol water, miscible with alc. hohol, ether, and petroleum ether. The molecular weight determined by Raoult's method agrees with the formula. The molecular re-

fraction $\frac{(n^2-1)P}{(n^2+2)d}$ is 46.3. Lævo menthone is

converted into a mixture of dextro and lævo menthone by acids, bases, water, heat, or even on keeping.

Oxim $C_{10}H_{17}NOH$. Mol w 169 [58°]. $[\alpha]_D = -42.5^\circ$. Crystalline, sol dilute alcohol, ether, and petroleum ether. Raoult's method gives, when acetic acid is used, the mol w 169, but when benzene is used it gives 251. With HCl the oxim forms a compound $(C_{10}H_{17}NOH)HCl$ [119°], a white crystalline powder $\rightarrow C_{10}H_{15}NONa$.

Dextro menthone $C_{10}H_{18}O$. Mol w 154 (208°) S.G. $\frac{20}{20}$ 900 $[\alpha]_D = +28.14^\circ$. Obtained by freezing a mixture of H_2SO_4 (10 pts) and water (1 pt), adding menthene (2 pts), and shaking. The temperature is slowly raised to 30°, the product poured into ice, and the menthone extracted by ether (Beckmann, *A* 250, 334). Colourless mobile oil, sl sol water, miscible with alcohol, ether, and petroleum ether. The mol w determined by Raoult's method is 150. The action of acids, bases, water, heat, and time convert dextro menthone into a mixture of dextro and lævo menthone.

Oxim $C_{10}H_{17}NOH$. Thick oil, slightly lævo rotatory. The mol w determined by Raoult's method is 165 when acetic acid is used, and 228 when benzene is employed as solvent $-(C_{10}H_{17}NOH)HCl$ [c 97°] Deliquescent $\rightarrow C_{10}H_{15}NONa$.

MENYANTHIN $C_{20}H_{32}O_{11}$ [60°-115°]. A bitter glucoside found in buckbean (*menyanthes trifoliata*) (Brandes, Kromayer, *Ar Ph* [2] 108, 257, 124, 37, *Fr* 1, 15). Prepared by treating the aqueous extract with animal charcoal at 65°, and subsequently extracting the charcoal with alcohol. Amorphous yellowish mass, sl sol cold water, v sol hot water and alcohol, insol ether. Dissolves in aqueous alkalis. Its solutions are not ppd by metallic salts. It is ppd by tannin. When distilled with dilute H_2SO_4 it gives off volatile menyanthol, while glucose $C_6H_{12}O_6$ aq remains behind.

Menyanthol $C_8H_{16}O$ is an oil smelling like benzoic aldehyde. It reduces ammoniacal $AgNO_3$. By exposure to the air, or by potash-fusion, it is converted into a crystalline sublimable acid.

Menyanthin. This name has also been applied to inulin when prepared from buckbean.

MERCAPTALS. Compounds of the form $RCH(SR_2)$. They may be viewed as thio acetals or as the sulphur compounds corresponding to the alkyl derivatives of ortho aldehydes. **V. THIO ALDEHYDES**

MERCAPTAN C_2H_5S *tc* C_2H_5SH *Ethyl mercaptan Thio alcohol*. Mol w 62 (36.5°). S.G. $\frac{4}{20}$ 8891 V.D. 2.11 (calc 2.15). S.V. 76 (Loosen, *A*, 264, 71) H.F.p. 14,480. H.F.v.

18,271 (Thomsen) R_{∞} 18 62 (Nasini, *G* 18, 301)

Formation—1 By distilling calcium ethyl sulphate $\text{CaEt}_2(\text{SO}_4)_2$ with a solution of barium sulphhydrate (Zeise, *P* 31, 369)—2 A mixture of alcohol and H_2SO_4 is neutralised with KOH, decanted from $\text{ppd K}_2\text{SO}_4$, mixed with excess of KOH, saturated with H_2S , and distilled (Wohler)—3 A solution of KOH (*S* G 13) is saturated with H_2S , mixed with an equal volume of a conc solution of K_2SO_4 , and distilled from a water bath (Liebig, *A* 11, 14, 23, 34) The product is freed from Et_2S by fractional distillation (Claesson, *J pr* [2] 15, 193)—4 From KSH and EtCl (Regnault, *A Ch* [2] 71, 390)—5 From KSH and EtI (Baudrimont, *C R* 51, 616)—6 By reducing with HI at 160° the solid thio aldehyde obtained by passing H_2S into a mixture of aldehyde solution and iodine (Bottinger, *B* 11, 2203)—7 Together with HETSO_4 and ether, by heating alcohol saturated with SO_2 in sealed tubes (Endemann, *A* 140, 336, Pagliani, *B* 11, 155)

Preparation—A cold mixture of alcohol (1 litre) and H_2SO_4 (500 c c) is diluted with ice and then added to a solution of crystallised sodium carbonate (4 kilos) The product is concentrated until Na_2SO_4 separates The mother liquor is further concentrated, and mixed with KOH (800 g) dissolved in water (1,500 c c) previously saturated with H_2S The whole is heated on a water bath, and the gases are passed first through a strong solution of caustic potash (50 g) and then into a solution of KOH (350 g) in water (700 g) The small amount of H_2S contained in the latter solution is ppd by lead acetate, and the mercaptan liberated by HCl It is dried with K_2CO_3 and distilled (Klason, *B* 20, 3407)

Properties—Colourless, very mobile, liquid, having a peculiarly persistent alliaceous odour V sl sol water, miscible with alcohol and ether Very inflammable, and burns with a blue flame A drop of it when briskly agitated on the end of a glass rod becomes solid, possibly through formation of a hydrate Neutral to litmus An alcoholic solution forms a yellow pp with lead salts, and white pps with cupric acetate, mercuric salts, and trichloride of gold The alcoholic solution is turned blue by FeCl_3 (Rathke, *A* 161, 148) Mercaptan is a reducing agent thus it converts nitro benzene into azo benzene Mercaptan decomposes salts of phenyl with formation of sodium phenylate and a thio ether (R Seiffert, *J pr* [2] 31, 462) But with phenyl salicylate it acts thus $\text{C}_6\text{H}_5(\text{OH})\text{CO}_2\text{Ph} + \text{NaSEt} = \text{C}_6\text{H}_5(\text{ONa})\text{CO}_2\text{Ph} + \text{HSEt}$ A trace of isatin dissolved in sulphuric acid gives, with a trace of mercaptan, a fine green colouration (Berthelot, *C R* 108, 350)

Reactions—1 Boiling nitric acid (*S* G 14) becomes red, deposits an oil, and ultimately forms ethane sulphonic acid Weaker nitric acid (*S* G 128) forms EtSO_3SEt —2 Potassium forms hydrogen and KSEt —3 P_2S_5 forms Et_2PS and Et_2HPS (Carius, *A* 112, 190)—4 Bromine forms EtBr and S_2Br_2 (Friedel & Ladenburg, *A* 145, 189)—5 Heated in sealed tubes with diazo-compounds (diazo salicylic acid diazo phenol chloride) it forms di ethyl disulphide, the diazo-compound being reduced exactly as when

boiled with alcohol (R Schmitt & O Mittenzwey, *J pr* 126, 192) $\text{C}_6\text{H}_5(\text{OH})\text{NCl} + 2\text{EtSH} = \text{Et}_2\text{S}_2 + \text{N}_2 + \text{C}_6\text{H}_5\text{OH} + \text{HCl}$ —6 PCl_5 at 100° forms thiophosphorous ether $\text{P}(\text{SEt})_3$, a fetid oil, resolved by distillation into phosphorus and Et_2S_2 —7 CCl_4 , acting on NaSEt , forms $\text{C}(\text{SEt})_4$, a light oil, *S* G 101, decomposed on distillation CCl_4 with NaSEt gives $\text{C}_2(\text{SEt})_4$, a heavy fetid oil, solidifying at very low temperatures C_2Cl_4 , heated with NaSEt at 100° gives $\text{C}_2(\text{SEt})_4$, crystallising in rhombohedra [54°]—8 S_2Cl_2 forms HCl and Et_2S_2 , which is resolved by distillation into Et_2S_2 and S_2 —9 TiCl_4 gives EtSHTiCl_3 and $(\text{EtSH})_2\text{TiCl}_2$ (Demarçay, *Bl* [2] 20, 132)—10 Reacts with ketones forming 'mercaptates' or thioketates Thus HCl passed into a mixture of acetone and mercaptan forms $(\text{CH}_3)_2\text{C}(\text{SEt})_2$ (Baumann, *B* 18, 887)—11 Mercaptan suspended in ice water and mixed with diazobenzene sulphonic acid and soda gives explosive yellow needles of $\text{SO}_3\text{Na C}_6\text{H}_4\text{N}_2\text{SEt}$ When the alcoholic solution of this salt is boiled $\text{SO}_3\text{Na C}_6\text{H}_4\text{N}_2\text{SEt}$ and nitrogen are formed (Stadler, *B* 17, 2075)

Salts—All metallic mercaptides are decomposed by HCl —* KSEt From mercaptan and K Dull white granular mass, v sol water, sl sol alcohol Its aqueous solution gives a yellow pp with lead salts— NaSEt From mercaptan and Na or NaOEt Crystalline mass, v sol water and alcohol Its aqueous solution is alkaline in reaction, and gives off all its mercaptan on boiling In dry air it oxidises to $\text{C}_2\text{H}_5\text{SO}_2\text{Na}$ When oxygen is passed through its alcoholic solution Et_2S is formed— $\text{Zn}(\text{SEt})_2$ white, indistinctly crystalline pp obtained by adding mercaptan to aqueous zinc acetate Sol NH_4Ac and leppd on neutralisation— $\text{Cd}(\text{SEt})_2$ — $\text{Co}(\text{SEt})_2$ gummy pp Unlike the other mercaptides it is not attacked by fuming HNO_3 — $\text{Ni}(\text{SEt})_2$ chocolate brown powder, not decomposed by water—* $\text{Fe}(\text{SEt})_2$ from FeCl_3 , mercaptan, and ammonia Black slimy mass, resolved by heating into mercaptan and ferrous hydrate— $\text{EtSFe}(\text{NO})$, [78°] From $\text{KSEt}(\text{NO})$, and EtI (Pavel, *B* 15, 2607)— TiSEt yellow curdy pp , sl sol water— $\text{Cu}(\text{SEt})_2$ Obtained by adding mercaptan to a solution of CuSO_4 mixed with NaOAc Pale yellow amorphous powder (Klason, *B* 20, 3407)— $\text{Hg}(\text{SEt})_2$ [77°] S 7 in 85 p c alcohol Formed on shaking HgO with mercaptan or with an alcoholic solution of mercaptan, the action being attended with great evolution of heat It is from this body that mercaptan derives its name (*corpus mercurio aptum*) White unctuous scales When its alcoholic solution is heated to 190° it is decomposed into mercury and Et_2S (Otto, *B* 18, 1289, 15, 125) The same decomposition appears to occur when it is heated in the dry state It is not decomposed by aqueous KOH, but aqueous K_2S forms some HgS and KSEt H_2S forms HgS and mercaptan Nitric acid oxidises it to $(\text{C}_2\text{H}_5\text{SO}_3)_2\text{HgHgO}$ When melted with lead the products are mercury, PbS , and Et_2S Mercury mercaptide forms with iodoform needles of $(\text{EtS})\text{Hg}(\text{CHI}_3)$ [85.5°] (Jackson & Oppenheim, *B* 8, 1033)— EtSgHgCl White bulky pp obtained on mixing an alcoholic solution of mercaptan with HgCl_2 Changes after some time to crystalline laminae Sl sol. water and ether,

m sol boiling alcohol (Debus, *A* 72, 18) — $\text{Pb}(\text{SEt})_2$. Yellow pp, got by mixing alcoholic solutions of lead acetate and mercaptan. Decomposed by heating with alcohol at 190° into PbS and Et_2S — $\text{Bi}(\text{SEt})_3$, [79°] Obtained by mixing a solution of a salt of bismuth with mercaptan. Long flexible yellow needles, m sol alcohol and ether. Oxidises in the air. Resolved by heat into BiS and Et_2S — $\text{Sn}(\text{SEt})_4$. Formed by mixing solutions of mercaptan and SnCl_4 in CS_2 , and evaporating the solvent. Also from mercaptan and a conc aqueous solution of SnCl_4 . Oil, remaining liquid at -40° . May be distilled *in vacuo*, but when heated under atmospheric pressure it decomposes yielding Et_2S and metallic tin — $\text{Sn}(\text{SEt})_2$, yellow pp which quickly oxidises in air — EtSSbCl_4 . Oil, formed by evaporating a mixture of mercaptan and SbCl_5 . Decomposed by water with liberation of mercaptan — $\text{As}(\text{SEt})_3$. Obtained by adding NaSEt to an ethereal solution of AsCl_3 . Oil, with unpleasant odour. Decomposed on distillation into arsenic and Et_2S — $\text{ClAs}(\text{SEt})_2$. From AsCl_3 and mercaptan in the cold — EtSAu . White gelatinous pp obtained on mixing dilute alcoholic solutions of auric chloride and mercaptan — $\text{Pt}(\text{SEt})_2$, light yellow pp — $\text{Rh}_2(\text{SEt})_4$, yellow pp.

Hydrate $\text{EtSH} \cdot 18\text{H}_2\text{O}$. Solidifies as a crystalline mass when moist vapour of mercaptan is passed through a condenser cooled below 8° (H. Muller, *Ar Ph* [2] 150, 147). Mass of small needles, insol water and mercaptan. A compound containing 1.6 p.c. carbon, crystallising in prisms, is obtained by pouring mercaptan into a solution of H_2S at 0° . It perhaps contains H_2S as well as water. It melts and decomposes above 3° (Blakie, *Pr E* 10, 87).

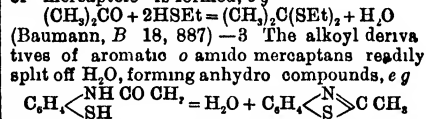
MERCAPTANS (Sulphydrates) Compounds of the formula RSH where R is a hydrocarbon radicle. They may be viewed as acid ethers of sulphydric acid or as alcohols in which O has been displaced by S . Just as H_2S is more acid than H_2O , the mercaptans are more acid than the alcohols. Thus they readily form salts by reacting with metallic oxides, and they derive their name from the ease with which they form mercuric salts. The salts of mercaptans may be called mercaptides. The mercaptans boil at a lower temperature than the corresponding alcohols, just as H_2S has a lower boiling point than H_2O . The mercaptans have an unpleasant odour. They are very readily oxidised, forming disulphides R_2S_2 , and finally sulphonic acids RSO_3H . The oxidation to sulphonic acids is best effected by nitric acid, and the chlorides of the sulphonic acids RSO_2Cl when reduced by tin and HCl yield the mercaptans again (Vogt, *A* 119, 152). By heating mercaptans with H_2SO_4 , disulphides R_2S_2 are formed (Erlenmeyer a Lisenko, *J* 1861, 590).

Preparation — 1 Fatty, but not aromatic, mercaptans are formed by heating a solution of KSH with alkyl chlorides or iodides or with potassium alkyl sulphates — 2 By heating alcohols or phenols with P_2S_5 (Kekulé, *A* 90, 311, *Z* 1867, 193). The yield is small — 3 By reducing chlorides of sulphonic acids with zinc or tin and dilute H_2SO_4 or HCl . This process is very convenient for the preparation of aromatic mercaptans. The product is distilled with steam.

4 Aromatic mercaptans may be prepared by treating diazo compounds with K_2S (Klason, *B*, 20, 350) — 5 Aromatic mercaptans may be obtained by reducing the corresponding disulphides with zinc and H_2SO_4 — 6 Aromatic mercaptans can be formed, together with the corresponding sulphides, by the action of aluminium chloride on a mixture of hydrocarbon and sulphur at 80° (Friedel a Crafts, *Bl* [2] 31, 464) — 7 When three atoms of hydrogen in the benzene nucleus have been displaced by chlorous radicles, and two of these are NO , and a halogen in ortho position to each other, alcoholic KSH displaces the halogen by SH (Beilstein a Kurbatoff, *A* 197, 75) — 8 By treating aromatic diazo compounds with sodium thiosulphate and reducing the resulting aromatic thiosulphates with zinc and H_2SO_4 (Leuckart, *G P* 1887, 45, 120).

Properties — Oils or crystalline solids with unpleasant odour. Insol water. The salts of the heavy metals are sparingly soluble in water. The mercuric salts can usually be recrystallised from alcohol. When a small quantity of a 1 p.c. solution of isatin in H_2SO_4 is mixed with a few c.c. of the strong acid and a small quantity of a fatty mercaptan, a green colouration is produced. Aldehydes and the higher alcohols interfere with the isatin reactions, and in this case the liquid may be shaken with a solution of KOH and then mixed with a little sodium nitroprusside, when a reddish violet colouration is produced (Demigès, *C R* 108, 350). The alkyl sulphides do not give this reaction. When sulphides are present they should first be pptd by an alkaline solution of PbO .

Reactions — 1 The ammoniacal solutions of aromatic mercaptans oxidise in the air with formation of disulphides — 2 When HCl is passed into a mixture of a ketone and a mercaptan condensation takes place and a thioketate or 'mercaptate' is formed, e.g.



These anhydro-compounds may even be formed by boiling the alkyl derivatives of aromatic amines with sulphur. They are also formed by the oxidation (by K_2FeCy_6) of the thioalkyl derivatives of aromatic amines. They are volatile liquids which exhibit feeble basic characters, and regenerate the amido mercaptans when fused with potash (Hofmann, *B* 13, 8, 1223, Jacob sen, *B* 19, 1069). When o-amido mercaptans are diazotised they produce characteristic stable compounds of the form $\text{C}_6\text{H}_4\text{--}\begin{array}{c} \text{N} \\ \text{S} \end{array}\text{N}$.

MERCAPTURIC ACIDS Acids obtained by placing chloro- or bromo benzene in food eaten by dogs. *V BROMO PHENYL* and *CHLORO PHENYL-MERCAPTURIC ACIDS*.

MERCURAMMONIUM COMPOUNDS (Ammonio mercury compounds Ammoniacal mercury bases Mercurammines) By the reactions of ammonia, or ammonium salts, on compounds of Hg , compounds are obtained, many of which may be represented by the empirical formulae $x\text{HgX} \cdot y\text{NH}_3$ and $x\text{HgX}_2 \cdot y\text{NH}_3$, where X = acidic radicle, some of the compounds, however, con-

tain N and H in the ratio NH_2 , and some are composed of N, Hg, and acidic radicles only. The mercurammonium compounds, as a class, do not react as loose compounds of Hg salts with NH_3 , but rather as compounds of Hg, N, H, and acidic radicles.

The mercurammonium compounds were regarded by Kane as compounds of NH_2Hg . Thus, the compound $HgONH_2$ was formulated as NH_2HgOH , and the compound $HgSO_2NH_2$ as $NH_2HgOSO_2NH_2$ (*A Ch* [2] 72, 215). Hirzel (*A* 84, 258) represented the mercurammonium compounds as compounds of mercuric nitride, N_2Hg_2 . Gerhardt suggested that these compounds are derived from NH_3 by replacing H by Hg, and after the ammonium hypothesis had been somewhat developed, Hofmann attempted to derive the compounds in question from different hypothetical mercurammoniums, i.e. compounds formed by replacing the H of NH_3 more or less completely by Hg (*v Handwörterbuch der Chemie* [2nd ed.] 2, 751, cf. Schmieder, *J pr* 75, 147, Nessler, *J* 1856 409). On this view, the compounds $HgCl_2 \cdot 2NH_3$ and $HgSO_4 \cdot 2NH_3$

are represented as $Hg \begin{smallmatrix} \text{NH}_2\text{Cl} \\ \text{NH}_2\text{Cl} \end{smallmatrix}$ and

$Hg \begin{smallmatrix} \text{NH}_2\text{SO}_4 \\ \text{NH}_2\text{SO}_4 \end{smallmatrix}$, derived from $\begin{smallmatrix} \text{NH}_2\text{Cl} \\ \text{NH}_2\text{Cl} \end{smallmatrix}$ and

$\begin{smallmatrix} \text{NH}_2\text{SO}_4 \\ \text{NH}_2\text{SO}_4 \end{smallmatrix}$. The mercurammonium compounds

may be divided, on this view, into classes according to the hypothetical Hg derivative of ammonium from which they are supposed to be derived. Thus we have the following N_2H_2Hg = mercurammonium, $N_2H_2Hg_2$ = mercurousammonium, $N_2H_2Hg_3$ = dimercurammonium, $N_2H_2Hg_4$ = dimercurousammonium, the mercurammonium compounds are those containing divalent Hg, and the compounds of monovalent Hg are called mercurousammonium derivatives. The compound $2HgONH_2$, obtained by the reaction of HgO with NH_4Aq , is represented on the ammonium hypothesis as hydrated mercurammonium hydrazide, $NH_2HgOH \cdot H_2O$.

A division is sometimes made between those mercurammonium compounds which contain N and H in the ratio NH_2 , and those which contain these elements in the ratio NH_4 , the latter are called mercuramide compounds.

Although some compounds readily give off NH_3 when heated, and others only when heated with alkali, yet it seems simpler to name all the compounds under consideration in accordance with the ammonium hypothesis. In this article the mercurammonium compounds are divided into two classes, mercurous and mercuric, the former being called mercurous and the latter mercuric compounds, the name given to each compound is intended to indicate the number of NH_2 groups from which the compound is derived, and the number of Hg atoms (mercurous or mercuric) in the formula. Several compounds are known whose reactions seem well expressed by supposing them to contain the group $HgOH$, such compounds are called mercurous ammonium derivatives in this article.

The following table presents the classification and nomenclature adopted in the present article --

Class I Mercurous compounds

Series (i) Mercurous-ammonium compounds, e.g. $(NH_2Hg)Cl$

„ (ii) Dimercurous-ammonium compounds, e.g. $(NH_2Hg_2)Cl$

Class II Mercuric compounds

Series (i) Mercuric-ammonium compounds, e.g. $(NH_2Hg)Cl$

„ (ii) Dimercuric-ammonium compounds, e.g. $(NH_2Hg_2)O$

„ (iii) Mercuric-diammonium compounds, e.g. $(N_2H_2Hg)I_2$

„ (iv) Dimercuric-diammonium compounds, e.g.

$(N_2H_2Hg_2)SO_4 \cdot H_2O$

„ (v) Trimercuric-diammonium compounds, e.g.

$(N_2H_2Hg_3)SO_4 \cdot 2H_2O$

Class III Mercurous ammonium compounds, e.g. $(NH_2Hg_2O)OH$, $(NH_2Hg_2O)SO_4$

Dimercurous ammonium chloride $(NH_2Hg_2)Cl$ is sometimes called mercurous chloramide or amido mercurous chloride, to express this view of its constitution the formula is written $Hg(NH_2)Cl$. So also mercurous ammonium chloride $(NH_2Hg)Cl$ is sometimes called ammonio mercurous chloride, and the formula is written, empirically, $HgClNH_2$, the formula of this compound is frequently doubled, and the name mercurousammonium chloride— $(N_2H_2Hg_2)Cl_2$ —is given to it. The salts called in this article mercurous ammonium compounds may be regarded as hydrated dimercurous ammonium salts, thus, the chloride $(NH_2Hg_2O)Cl$ may be written $(NH_2Hg_2O)Cl \cdot H_2O$, and the carbonate $(NH_2Hg_2O)_2CO$, may be written $(NH_2Hg_2O)_2CO \cdot 2H_2O$.

Class I Mercurous compounds $NH_2 \cdot 2Hg \cdot X$

The experiments of Barfoed (*J pr* [2] 39, 201) make it probable that the so called mercurous ammonium compounds are really mixtures of mercuric compounds and Hg. B asserts that exactly half the Hg in the black pps formed by NH_4Aq in solutions of mercurous salts exists uncombined, that nearly the whole of this Hg disappears as vapour when the pps are freely exposed to air, and that the light coloured compounds remaining are the same as those produced by adding NH_4Aq to mercuric salts. To the pp produced from $HgNO_3$, Barfoed gives the composition $3Hg + 2(HgNH_2NO_3)HgO$, that produced from Hg_2SO_4 is $4Hg + (NH_2Hg_2)SO_4 \cdot 2HgO$, that from $HgCl$ is $Hg + NH_2Hg_2Cl$, and that from $Hg_2C_2O_4$ is $4Hg + (NH_2Hg_2)_2C_2O_4 \cdot 2HgO$.

Series (i) Mercurous ammonium compounds; NH_2HgX

Mercurous-ammonium chloride, $(NH_2Hg)Cl$. (*Ammonio mercurous chloride* ($HgClNH_2$), *Mercurous ammonium chloride* ($N_2H_2Hg_2Cl_2$)). A black powder, obtained by saturating $HgCl$, prepared by ppn, with NH_3 gas, on gently warming all NH_3 is given off (*H Rose*, P 20, 159). The dissociation of this compound has been studied by Isambert (*C R* 66, 1259, v *Disso* *ciation*, vol II p 397). According to Barfoed (*J pr* [2] 39, 201) the compound formulated as above is really a mixture of Hg, mercuric ammonium chloride (NH_2Hg_2Cl) , and NH_4Cl .

Series (ii) Dimercurous-ammonium compounds, NH_2Hg_2X

Dimercur ammonium chloride, $(\text{NH}_4\text{Hg})\text{Cl}$ (*Mercurchloramide* or *amido-mercurous chloride* (HgNH_2Cl) *Tetramercurio - diammonium chloride* $(\text{N}_2\text{H}_4\text{Hg}_2\text{Cl}_2)$) A black solid, obtained by digesting HgCl with NH_4Aq ($2\text{HgCl} + 2\text{NH}_4\text{Aq} = \text{NH}_4\text{Hg}_2\text{Cl} + \text{NH}_4\text{ClAq}$) Becomes dark grey when dried, heated, gives off NH_3 and N , and at a higher temperature yields a sublimate of HgCl and Hg , treated with HCl gas, gives HgCl and NH_4Cl (Kane, *A Ch* [2] 72, 215, Ullgren, *P* 42, 395) According to Barfoed (*J pr* [2] 39, 201) the black solid obtained as described is a mixture of Hg and NH_4HgCl

Mercur ammonium nitrates Various compounds of the form $x\text{Hg}_2\text{O} \cdot y\text{N}_2\text{O}_5 \cdot z\text{NH}_3$ have been described by Kane (*lc*) and Mitscherlich (*P* 9, 387, 16, 41), but Barfoed's experiments make it very probable that these bodies are mixtures of Hg and mercuri ammonium salts (*vsupra*)

Class II. MERCURIC COMPOUNDS $\text{N}_2\text{H}_4 \cdot \text{Hg}_2\text{X}_2$

Series (1) Mercuri ammonium compounds, NH_4HgX Obtained by interaction of excess of NH_4Aq with mercuric salts in solution

Mercuri-ammonium chloride, $(\text{NH}_4\text{Hg})\text{Cl}$ (*Mercuri-chloramide*, or *amido mercuric chloride* (HgNH_2Cl) *Dimercuri diammonium chloride* $(\text{N}_2\text{H}_4\text{Hg}_2\text{Cl}_2)$ *Infusible white precipitate*) According to Rammelsberg (*J pr* [2] 38, 558) this salt is a double compound of dimercuri ammonium chloride— NH_4HgCl —and NH_4Cl , R assigns to it the formula $\text{NH}_4\text{HgCl} \cdot \text{NH}_4\text{Cl}$ (*vsupra*)

When excess of NH_4Aq is added to HgCl_2Aq , or when HgCl_2Aq is dropped into NH_4Aq , a white pp is obtained, this pp was long considered identical with that obtained by adding $\text{Na}_2\text{CO}_3\text{Aq}$ to HgCl_2Aq mixed with NH_4ClAq , and known in pharmacy as *Mercurius precipitatus albus* Wöhler (*P* 26, 203) found that the pp formed by NH_4Aq was volatilised without fusion when heated in a Pt dish, but that the pp formed by $\text{Na}_2\text{CO}_3\text{Aq}$ melted before it volatilised. The pp by NH_4Aq was then called *infusible white precipitate*, and to that produced by $\text{Na}_2\text{CO}_3\text{Aq}$ the name *fusible white precipitate* was given

Preparation— HgCl_2Aq is added to excess of cold NH_4Aq , the pp is collected at once, rapidly washed with a little cold water, and dried at 110° (André, *C R* 108, 293, 290) André (*lc*) has examined the compounds obtained by the interaction of NH_4Aq with HgCl_2Aq , varying the relative masses of the reacting bodies and the time of contact Using equal volumes of HgCl_2Aq , containing 33.875 g HgCl_2 per litre, and NH_4Aq containing 4.25 g NH_3 per litre, the pp after drying at 110° was not quite pure NH_4HgCl , but contained a little mercurous ammonium chloride $(\text{NH}_4\text{Hg}_2\text{O})\text{Cl}$ The quantity of $(\text{NH}_4\text{Hg}_2\text{O})\text{Cl}$ increased by allowing the reacting bodies to remain in contact, and also by increasing the quantity of NH_4Aq , when the NH_3 and HgCl_2 were present in the ratio $6\text{NH}_3 \cdot \text{HgCl}_2$, the pp contained equal molecular proportions of NH_4HgCl and $(\text{NH}_4\text{Hg}_2\text{O})\text{Cl}$, and by further increasing the NH_3 , only $(\text{NH}_4\text{Hg}_2\text{O})\text{Cl}$ was obtained André found that NH_4Cl tends to change $(\text{NH}_4\text{Hg}_2\text{O})\text{Cl}$ into NH_4HgCl , by adding 1 vol HgCl_2Aq to 5 vols NH_4Aq (strength as given above), decanting after 24 hours, and agitating the pp for 2 days with 2 vols NH_4ClAq (containing 13.4 g.

per litre), pure NH_4HgCl was obtained André has also examined the pps obtained by adding NH_4Aq , and also NH_4ClAq , to HgCl_2Aq in presence of KOH (*C R* 108, 1108, 1164) The pps are compounds of NH_4HgCl , $\text{NH}_4\text{Hg}_2\text{O}$, and $(\text{NH}_4\text{Hg}_2\text{O})\text{Cl}$ in varying proportions

Properties—A white solid, which volatilises completely, when strongly heated, without melting Readily sol in HClAq , not blackened by NH_4Aq , dissolved by acids, also by hot solutions of NH_3 salts

Reactions—1 When heated, in an open vessel, does not melt but yields a sublimate of HgCl and a mixture of 1 vol N and 2 vols NH_3 , $(6\text{NH}_4\text{HgCl} = 6\text{HgCl} + 4\text{NH}_3 + \text{N}_2)$, Kane, *A Ch* [2] 72, 215) When slowly heated in a retort to c 340° , HgCl , and NH_3 are given off, and a red crystalline compound, $2\text{NH}_4\text{HgCl} \cdot \text{HgCl}_2$, remains —2 Decomposed by water, slowly by cold, more rapidly by hot, eventually with formation of yellow mercurous ammonium chloride, $(\text{NH}_4\text{Hg}_2\text{O})\text{Cl}$, and NH_4Cl ($2\text{NH}_4\text{HgCl} + \text{H}_2\text{O} + \text{Aq} = (\text{NH}_4\text{Hg}_2\text{O})\text{Cl} + \text{NH}_4\text{ClAq}$) —3 Potash or soda separates $(\text{NH}_4\text{Hg}_2\text{O})\text{Cl}$ and evolves NH_3 Only half the N of the original NH_4HgCl is evolved as NH_3 (Kane, *lc*), but by boiling for some days with KOH , renewed from time to time, all the N is removed as NH_3 , and HgO remains (Schmieder, *J pr* 75, 147) The fact that only half the N is readily evolved as NH_3 , by the action of alkalis on NH_4HgCl , has induced Rammelsberg (*J pr* [2] 38, 558) to regard this body as a compound of dimercuri ammonium chloride and NH_4Cl , and to give it the formula $\text{NH}_4\text{HgCl} \cdot \text{NH}_4\text{Cl}$ But it is to be remembered, as pointed out by Kane, that water readily separates NH_4HgCl into $(\text{NH}_4\text{Hg}_2\text{O})\text{Cl}$ and NH_4Cl , hence the NH_3 obtained by the action of aqueous alkali is probably due to the direct decomposition of NH_4Cl by the alkali —4 Boiled with very dilute sulphuric acid till dissolved, the compound $\text{HgCl}_2 \cdot 2\text{NH}_4\text{Cl} \cdot 2\text{H}_2\text{SO}_4$ forms on cooling (Kosman, *A Ch* [3] 27, 238) —5 Heated in dry hydrogen chloride, HgCl_2 and NH_4Cl are produced (Ullgren, *A* 26, 203) —6 Heated with solutions of sodium chloride potassium iodide, barium sulphide, &c, NH_3 is evolved and HgCl_2 , HgI_2 , HgS , &c, formed, e.g. $\text{NH}_4\text{HgCl} + 2\text{KI} + \text{H}_2\text{O} = \text{HgI}_2 + \text{KClAq} + \text{KOH} + \text{NH}_3$ (Kane, *lc*, Rammelsberg, *P* 48, 182) —7 Boiled with ammonium chloride solution, mercuri diammonium chloride $\text{N}_2\text{H}_4\text{Hg}_2\text{Cl}_2$ (*fusible white precipitate*) is produced —8 Decomposed by heating with sulphur, with formation of a yellow sublimate of a chlorosulphide of N —9 Rubbed with iodine and a little alcohol, HgI_2 is formed and then an explosion occurs, probably from production and decomposition of N iodide (Rice, *Ph* [3] 6, 765, cf. Schwarzenbach, *B* 8, 1231, Flückiger, *B* 8, 619) —10 Bromine and chlorine cause evolution of N and formation of HgBr_2 , or HgCl_2 (Schwarzenbach, *B* 8, 1231) —11 Decomposed by alcoholic iodides, e.g. EtI (*v* Sonnenschein, *A* 101, 20) —12 For reaction with amylic mercaptan *v* Wagner, *J pr* 53, 378

Combinations—1 Forms various compounds with NH_4HgCl and $(\text{NH}_4\text{Hg}_2\text{O})\text{Cl}$; obtained by adding NH_4Aq or NH_4ClAq to solutions of HgCl_2Aq containing KOH (André, *C R* 108, 1108, 1164) —2 Combines with HgCl_2 to form

NH_4HgCl HgCl_2 ; produced by slowly heating to $c\ 340^\circ$ in a retort (Millon, *A Ch* [3] 18, 392)

Mercuri-ammonium bromide, $(\text{NH}_4\text{Hg})\text{Br}$ (*Mercuri bromide*, or *amido-mercuric bromide* ($\text{Hg NH}_2\text{Br}$)) *Dimercuri diammonium bromide* ($\text{N}_2\text{H}_4\text{Hg}_2\text{Br}_2$)) A yellow powder, obtained by ppg HgBr_2Aq with excess of NH_4Aq (*cf Mercuri ammonium chloride, supra*). Insol water or alcohol, slowly changed by cold water, more quickly by hot water, to NH_2HgBr (Pesci, *G* 19, 509) E sol NH_4Aq Heated gives N , NH_3 , and HgBr , when very slowly heated to $c\ 340^\circ$ gives NH_3 , HgBr , NH_3 , and NH_2HgBr (Mitscherlich, *J pr* 19, 455)

Series (ii) Dimercuri ammonium compounds, NH_2X According to Pesci (*G* 19, 509) many and probably all the dimercuri ammonium salts, when digested with NH_4Cl , NH_4Br , or NH_4I solution, evolve NH_3 in the proportion of 4NH_3 for each NH_2 group in the dimercuri-ammonium salt This reaction may be applied to estimate dimercuri ammonium salts, by conducting the digestion in a closed vessel containing a dish with a measured quantity of normal oxalic acid

Dimercuri-ammonium hydroxide, NH_2OH (*Mercurammonium hydroxide*) Prepared by saturating yellow HgO with NH_3 under increased pressure, finely powdering, and again saturating with NH_3 , or by shaking HgO with alcoholic NH_3 for some hours and drying at ordinary temp in NH_3 gas, the product, which is $\text{NH}_2\text{OH} \cdot \text{H}_2\text{O}$, is then heated in dry NH_3 to $80^\circ\text{--}85^\circ$ The operation should be conducted in the dark (Weyl, *P* 121, 601, 131, 539) A brown powder, explodes when strongly heated, decomposed by water, with evolution of NH_3 , absorbs CO from air, evolving NH_3 at the same time, dissolves in warm HClAq or HNO_3Aq with formation of salts of Hg and NH_4

Heated to 100° in absence of air, gives *dimercuri ammonium oxide* $(\text{NH}_2)_2\text{O}$ A dark brown powder which explodes when heated, struck, or rubbed in a mortar, readily combines with water to form $\text{NH}_2\text{OH} \cdot \text{H}_2\text{O}$, soluble in HClAq and HNO_3Aq , reacts with HCl gas to form HgCl_2 and NH_4Cl This oxide is also formed by the action of liquid NH_3 on HgO

The hydrate of dimercuri-ammonium hydroxide $\text{NH}_2\text{OH} \cdot \text{H}_2\text{O}$, is a yellow powder, obtained as described above It appears to be isomeric with mercurous ammonium hydroxide $(\text{NH}_4\text{Hg}_2\text{O})\text{OH}$ (p 211)

Dimercuri-ammonium chloride, NH_2Cl (*Mercurammonium chloride*) Obtained by treating the hydroxide (*v supra*) with alcoholic solution of HCl , also by treating the oxychloride formed by fusing together HgO and HgCl_2 in the ratio $3\text{HgO} \cdot \text{HgCl}_2$ with liquid NH_3 , and removing excess of NH_3 by warming in a stream of dry air at 150° Forms a yellow powder, decomposed suddenly at 300° to HgCl_2 , Hg , and N (Weyl, *P* 121, 601, 131, 529). Not acted on by boiling water, decomposed by hot KOH Aq with production of NH_3 and HgO ; slowly dissolved by cold HClAq

Forms a double compound with mercuric chloride $2\text{NH}_2\text{Cl} \cdot \text{HgCl}_2$ (*dimercuri-ammonium-mercuric chloride*). A red crystalline solid, resembling HgO , produced by very slowly

heating mercuri ammonium chloride, $\text{NH}_4\text{Hg Cl}$, in a retort until HgCl begins to sublime This compound reacts with hot HClAq to form NH_4Cl and HgCl_2 , it is said to be unchanged by conc HNO_3Aq , dilute $\text{H}_2\text{SO}_4\text{Aq}$, H_2O , or boiling alkali solutions (Mitscherlich, *J pr* 19, 453) Heated to 360° it separates into N , HgCl , and Hg

Dimercuri-ammonium bromide, NH_2Br (*Mercurammonium bromide*) Obtained by ppg HgBr_2Aq by excess of NH_4Aq and digesting the pp (NH_4HgBr) with water, better, by adding excess of NH_3 carbonate to HgBr_2Aq , washing the pp of $4\text{NH}_4\text{Br} \cdot 5\text{NH}_3\text{Br}$ with NH_4 carbonate, warming with KOH Aq , and washing with cold water (Pesci, *G* 19, 509) Also obtained, according to Pesci, by the action of dilute HBr Aq on Millon's base $(\text{NH}_4\text{Hg}_2\text{O})\text{OH}$ A yellow powder, insol water, sol HClAq , insol HNO_3Aq , decomposed by heat, without melting, yielding NH_3 and a sublimate which contains Hg (Pesci, *lc*) Digested for a short time with boiling $\text{NH}_4\text{Br Aq}$, and filtered, small crystals of $\text{NH}_4\text{Br} \cdot 5\text{NH}_3\text{Br}$ are obtained This compound seems similar to *fusible white precipitate*, it is described on p 210 as *mercuri diammonium bromide* The double compound $4\text{NH}_2\text{Br} \cdot 5\text{NH}_3\text{Br}$ is obtained, in clear microscopic needles, by adding excess of NH_3 carbonate to HgBr_2Aq (Pesci, *lc*)

Dimercuri ammonium iodide, NH_2I , is obtained by the action of liquid NH_3 on the oxy iodide formed by fusing together HgO and HgI_2 in the ratio $3\text{HgO} \cdot \text{HgI}_2$ (Weyl, *P* 121, 601, 131, 539)

Dimercuri ammonium selenate, $(\text{NH}_2)_2\text{SeO}_4 \cdot 2\text{H}_2\text{O}$ A white pp which darkens in light and is decomposed by heat; obtained by dissolving the basic selenate $\text{HgSeO}_4 \cdot 2\text{HgO}$ in conc NH_3Aq , and ppg by much water (Cameron a Davy, *C N* 44, 63)

Series (iii) Mercuri diammonium compounds, $\text{N}_2\text{H}_4\text{HgX}_2$

Mercuri-diammonium chloride, $\text{N}_2\text{H}_4\text{Hg Cl}_2$ (*Mercurammonium chloride*) *Fusible white precipitate* *Ammonio mercuric chloride*, $\text{HgCl}_2 \cdot 2\text{NH}_3$ Rammelsberg (*J pr* [2] 38, 558) regards this salt as a double compound of dimercuri ammonium chloride— NH_2Cl —and NH_4Cl He formulates it as $\text{NH}_2\text{Cl} \cdot 3\text{NH}_4\text{Cl}$ (*v infra*)

When an alkali carbonate is added to HgCl_2Aq containing NH_4Cl , a white pp is obtained, this pp was called in pharmacy *mercurius precipitatus albus*, and for long was supposed to be the same as that formed by adding NH_4Aq to HgCl_2Aq Wöhler (*P* 26, 203) found that the pp formed by alkali carbonate, in presence of sal ammoniac, melted when heated, before volatilising, but that the pp formed by ammonia volatilised without melting The former pp was then called *fusible white precipitate*, to distinguish it from the latter to which the name *infusible white precipitate* was given

The researches of Krug (*Ar Ph* 42, 1) have shown that pure mercuri diammonium chloride cannot be obtained by addition of alkali carbonate to HgCl_2Aq containing NH_4Cl at the ordinary temperature, the pp consists of a mixture of mercuri ammonium chloride (*infusible white precipitate*, $\text{NH}_2\text{Hg Cl}$) and mercuri-diam-

monium chloride (*fusible white precipitate*, $N_2H_5HgCl_2$), the longer the pp remains in contact with the supernatant liquor, and the higher the temperature, the greater is the amount of *fusible white precipitate* formed. Krug found that almost pure NH_4HgCl (*infusible white precipitate*) was obtained by adding Na_2CO_3 to $HgCl_2$ mixed with NH_4Cl at 0° , and filtering at once (about $4Na_2CO_3$, $3HgCl_2$, $9NH_4Cl$). The first pp formed by the alkali carbonate therefore probably always consists mostly of mercuri ammonium chloride, NH_4HgCl , but this is acted on by the NH_4Cl present, and thus more or less $N_2H_5HgCl_2$ is produced, CO_2 is also given off from the alkali carbonate used, and this CO_2 , according to Krug's experiments, seems to change some of the NH_4HgCl to $HgCl_2 \cdot 2HgO$ (with simultaneous formation of NH_4Cl and NH_4HCO_3), which oxychloride then probably reacts with NH_4Cl to produce $N_2H_5HgCl_2$.

Preparation— $HgCl_2$ is dropped into boiling NH_4Cl containing NH_3 , as long as the pp which forms is redissolved, and the liquid is allowed to cool. The compound crystallises out in garnet red dodecahedra. Melts at $c 300^\circ$ with decomposition.

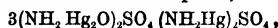
Properties and Reactions—Small red dodecahedra, also formed by ppn with Na_2CO_3 as a white powder. Gently heated, gives off half its NH_3 , leaving $HgCl_2$, (*v infra*), melts at $c 300^\circ$, evolving N and NH_3 and yielding a sublimate which reacts with water to form NH_4Cl , $HgCl_2$, and $HgCl$. Boiling water reacts to form mercurous ammonium chloride, $(NH_2HgO)Cl$, and NH_4Cl . Alkalis evolve NH_3 . According to Rammelsberg (*J pr* [2] 38, 558) three fourths of the N of mercuri ammonium chloride is evolved by the action of hot alkali solution, hence B. assigns to this compound the formula $NH_4HgCl \cdot 3NH_4Cl$, but as the action of boiling water on $N_2H_5HgCl_2$ is to produce $(NH_2HgO)Cl$ and NH_4Cl in the ratio $(NH_2HgO)Cl : 3NH_4Cl$, it is easy to account for the action of boiling alkali without supposing fusible white precipitate to be a double compound of NH_4HgCl with NH_4Cl . Iodine reacts energetically, according to Fluckiger (*B* 8, 1619) the reaction is expressed thus $6(N_2H_5HgCl_2) + 2I_2 = N_2 + 6NH_4Cl + 4NH_3 + 2HgCl_2 + 2HgI_2 + 2HgCl$. Liquid ammonia dissolves $N_2H_5HgCl_2$, after prolonged action, and evaporation of the excess of NH_3 , a white crystalline mass remains which has the same composition as the original, according to Weyl (*P* 1, 547).

Combination—The compound $HgCl_2 \cdot NH_3$ may be regarded as a compound of $N_2H_5HgCl_2$ with $HgCl_2$ [N_2H_5HgCl , $HgCl_2 = 2(HgCl_2 \cdot NH_3)$]. This compound is formed by heating $HgCl_2$ in dry NH_3 , or by distilling HgO with NH_4Cl , it melts when heated and distils without much change, it is decomposed by water, forming NH_4HgCl and $HgCl_2 \cdot NH_4Cl$ (Kane, *A Ch* [2] 72, 215).

Mercuri-diammonium bromide, N_2H_5HgBr , Small microscopic, rhombohedral, transparent crystals, obtained by adding excess of NH_3 carbonate to $HgBr_2$, boiling the pp of $4NH_3$, Br 5 NH_3 , Br for a short time with NH_4Br , filtering, and allowing to cool (Pesci, *G* 19, 509). Also formed by treating dimercuri ammonium bromide (NH_4HgBr) with boiling NH_4Br , by

the action of NH_3 on solution $HgBr_2$, NH_3 , Br ; also by adding alcoholic NH_3 to $HgBr_2$ dissolved in alcohol (*P*, *lc*). Melts at $c 180^\circ$ with evolution of NH_3 , more strongly heated, yields two distinct sublimates. Easily sol. HCl and H_2SO_4 forms $HgBr_2$, alkalis produce NH_4Br (*P*, *lc*). Forms a compound with $HgBr_2$, viz. $N_2H_5HgBr \cdot HgBr_2 = (HgBr_2 \cdot NH_3)$, similar to the chloro compound already described (*H Rose*, *P* 20, 160).

Mercuri-diammonium sulphate, $N_2H_5HgSO_4 \cdot H_2O$. Lustrous, orthorhombic, crystals. Obtained by adding, little by little, yellow HgO to pure, saturated, cold $(NH_4)_2SO_4$, each portion of HgO being allowed to dissolve before another portion is added, the liquid being kept cold, addition of HgO is continued till the liquid begins to grow turbid, the clear liquid is decanted and allowed to evaporate in the air or *in vacuo*. This compound is decomposed by the smallest quantity of water with production of a compound of mercurous ammonium sulphate and mercuri ammonium sulphate,



boiling water removes more H_2SO_4 , finally forming $(NH_2HgO)_2SO_4$ (*v Mercurous ammonium sulphate*, p 211, *cf* Millon, *A Ch* [3] 18, 410, Schmieder, *J pr* 75, 147). It is soluble in $(NH_4)_2SO_4$, NH_4Cl , HCl , dilute H_2SO_4 , and HNO_3 , insol. conc. HNO_3 . Boiled with conc. H_2SO_4 , it gives $(NH_4)_2SO_4$ and $HgSO_4$, decomposed by KOH , forming a basic compound, with boiling conc. KOH , NH_3 is evolved and HgO formed. Heated to 115° water is evolved, and at a higher temperature complete decomposition results.

Mercuri-diammonium iodide, N_2H_5HgI , *Mercuri-diammonium fluoride*, N_2H_5HgF , and the double compound $N_2H_5HgI \cdot HgI_2$, have been described (*H Rose*, *P* 20, 160, Nessler, *C C* 1856 530, Finkener, *P* 110, 147, Rammelsberg, *P* 48, 170, Caillot & Carriol, *J. Ph* 9, 381).

Series (iv) Dimercuri-diammonium compounds, $N_2H_5Hg_2X_2$.

Dimercuri-diammonium sulphate, $N_2H_5Hg_2SO_4 \cdot H_2O (= SO_4 \cdot 2HgO \cdot 2NH_3)$. Obtained, according to Millon (*A Ch* [3] 18, 410), by saturating 70 cc cold conc. NH_4 with $HgSO_4$, allowing to stand in an atmosphere of NH_3 over CaO for some months, powdering the crystals which separate, and drying over CaO .

Series (v) Trimercuri-diammonium compounds, $N_2H_5Hg_3X_2$.

Trimercuri-diammonium sulphate, $N_2H_5Hg_3SO_4 \cdot 2H_2O (= SO_4 \cdot 3HgO \cdot 2NH_3)$. Prepared similarly to preceding salt, but using 90 cc NH_4 (Millon, *lc*).

Class III MERCUROXY AMMONIUM COMPOUNDS (NH_2HgO)X.

The name *mercurous* is here given to the group HgO , which is supposed to act as a dyad radicle in these compounds. The salts of this series may be regarded as *hydrated tetramercuri-diammonium compounds*, thus the carbonate $(NH_2HgO)_2CO$ may be written $N_2Hg_4O_4 \cdot CO_2 \cdot 2H_2O$, and the chloride $(NH_2HgO)Cl$ may be written $N_2Hg_4O_4 \cdot Cl_2$. Some of the salts of this series may also be regarded as hydrated *dimercuri-*

ammonium compounds, thus the chloride ($\text{NH}_2\text{Hg}_2\text{O}$)Cl loses water at 200° , and is therefore regarded by Rammelsberg as $\text{NH}_2\text{Hg}_2\text{O}$.

Mercurioxy-ammonium hydroxide, ($\text{NH}_2\text{Hg}_2\text{O}$)OH (*Millon's base*). This is the final product of the reaction of NH_4Aq on HgO , the hydroxide contains H_2O . Millon gave the formula $\text{NH}_2\text{Hg}_2\text{O} \cdot \text{OH} \cdot \text{H}_2\text{O}$, Gerresheim, the formula $2(\text{NH}_2\text{Hg}_2\text{O} \cdot \text{OH}) \cdot \text{H}_2\text{O}$ (*A* 195, 373), and Rammelsberg, more recently (*J pr* [3] 38, 558), assigns the composition $3(\text{NH}_2\text{Hg}_2\text{O} \cdot \text{OH}) \cdot 2\text{H}_2\text{O}$. The empirical formula for the base is $2\text{HgO} \cdot \text{NH}_2 \cdot \text{H}_2\text{O}$.

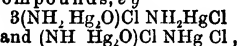
Preparation—Yellow HgO is agitated with NH_4Aq , quite free from carbonate, so long as any change occurs, if red HgO is used the change is not completed for some days. The yellow brown product is washed with cold water, crystallised from hot water, and dried (*Millon, A Ch* [3] 18, 392).

Properties—Yellow, microscopic crystals, probably rhombic. Appears to be isomeric with hydrated dimercuri ammonium hydroxide $\text{NH}_2\text{Hg}_2\text{OH} \cdot \text{H}_2\text{O}$ (*p* 209). Insol alcohol or ether, somewhat sol water. *S* 007 at 17° , 06 at 80° . Acts as a very strong base, decomposes NH_4 salts energetically with evolution of NH_3 , and rapidly combines with CO_2 when exposed to air. Several salts may be obtained by dissolving the base in excess of acetic acid, and ppg by alkali permanganate, chromate, &c. Over H_2SO_4 loses H_2O and becomes brown, heated to 130° further loses $\frac{1}{2}\text{H}_2\text{O}$, leaving *mercurioxy ammonium oxide* ($\text{NH}_2\text{Hg}_2\text{O}$).

Reactions—1 The hydroxide base is very slowly decomposed by boiling *alkali* solution, the dehydrated base is only decomposed by molten alkali—2 Decomposes *ammonium salts* rapidly, with evolution of NH_3 —3 Combines so rapidly with *carbon dioxide* that it is almost impossible to obtain the base quite free from carbonate—4 With acids forms salts ($\text{NH}_2\text{Hg}_2\text{O}$)X (X = NO_3 , $\frac{1}{2}\text{SO}_4$, &c) (*v* Rammelsberg, *J pr* [2] 38, 558)—5 Decomposed by *heat* with more or less explosion (*Millon, l c*, *cf* *Hirzel, J* 1852 419, 1853 381)—6 Many salts, *e.g.* sulphates and chlorides, are decomposed by shaking with *Millon's base*, with complete removal of the acid of the salt used (*v* Gerresheim, *A* 195, 873)—7 Heated with *ethylic iodide* to 100° yields crystals of $2\text{NEtI} \cdot 3\text{HgI}_2$, ethylic bromide reacts similarly—8 A thio carbonate of the base is obtained by heating with *carbon disulphide* (*Gerresheim, l c*).

Mercurioxy ammonium chloride, ($\text{NH}_2\text{Hg}_2\text{O}$)Cl (*Hydrated tetramercurammonium chloride* $\text{N}_2\text{Hg}_4\text{Cl}_2 \cdot 2\text{H}_2\text{O}$ *Hydrated dimercuri ammonium chloride* $\text{NH}_2\text{Hg}_2\text{Cl} \cdot \text{H}_2\text{O}$). This compound is formed by the reaction of boiling water with either fusible white pp $\text{N}_2\text{H}_4\text{HgCl}_2$, or infusible white pp NH_2HgCl . It is also produced by passing dry NH_3 over the oxychloride $8\text{HgO} \cdot \text{HgCl}_2$ at 150° (*Ullgren, P* 42, 395). A heavy, yellow powder, *v s* sol water, *e* sol HClAq and HNO_3Aq (*Kane, P* 42, 367). Decomposed by heat, giving NH_3 , N , H_2O , HgCl , and Hg . Decomposed by excess of solutions of KCl , NaCl , and KI , with evolution of NH_3 . Loses water at 200° , and is therefore regarded by Rammelsberg as $\text{NH}_2\text{Hg}_2\text{O} \cdot \text{H}_2\text{O}$ (*J. pr.* [2] 38, 558).

Mercurioxy-ammonium chloride combines with NH_2HgCl and $\text{NH}_2\text{Hg}_2\text{Cl}$ to form various double compounds, *e.g.*



these compounds were obtained by André by ppg HgCl_2Aq with NH_4Aq , some of them requiring the presence of KOH for their production (*C R* 108, 233, 290, 1108, 1164).

Mercurioxy ammonium iodide, ($\text{NH}_2\text{Hg}_2\text{O}$)I (*Hydrated tetramercurammonium iodide* $\text{N}_2\text{Hg}_4\text{I}_2 \cdot 2\text{H}_2\text{O}$ *Hydrated dimercuri ammonium iodide* $\text{NH}_2\text{Hg}_2\text{I} \cdot \text{H}_2\text{O}$). This compound is formed by adding NH_4Aq to *Nessler's reagent* (alkaline solution of HgI_2 in KIAq), the composition of the pp thus produced is often represented as $\text{HgNH}_2\text{I} + \text{HgO}$, or as $\text{HgNH}_2\text{I} + \text{H}_2\text{O}$. The compound is best prepared by dissolving HgI_2 , 2KI in KOH Aq , adding NH_4Aq , washing thoroughly with cold water, and drying at 160° ($2\text{K} \cdot \text{HgI}_2\text{Aq} + 3\text{KOH Aq} + \text{NH}_4\text{Aq} = (\text{NH}_2\text{Hg}_2\text{O})\text{I} + 7\text{KIAq} + 2\text{H}_2\text{O}$, *v* Rammelsberg, *P* 48, 170). ($\text{NH}_2\text{Hg}_2\text{O})\text{I}$ is also formed by heating $3\text{HgO} \cdot \text{HgI}_2$ to 180° in NH_3 (*Rammelsberg*), and by boiling HgI_2 with excess of conc NH_4Aq .

Mercurioxy ammonium iodide is a brown powder with a purple red tint. Begins to give off water at 128° , heated more highly, out of contact with air, it melts to a dark brown liquid, and then decomposes violently, giving H_2O , Hg , HgI , NH_3 , and N . Sol warm KIAq , with formation of K HgI , and KOH and evolution of NH_3 . Sol warm HClAq , HgI_2 , and HgICl are deposited on cooling (*Rammelsberg*). Decomposed by hot BaSAq , with evolution of all N as NH_3 . Heated in HCl gas, gives a sublimate of HgCl_2 , NH_4Cl , NH_4I , and (NH_4) HgI_2 .

Mercurioxy ammonium nitrates. Several nitrates, and some double nitrates, of the mercurioxy ammonium compounds seem to exist.

The salt often called *Soubiran's ammoniacal salt* is probably the *normal nitrate* ($\text{NH}_2\text{Hg}_2\text{O})\text{NO}_3$ (*Soubiran, J Ph* 12, 465). This compound is obtained by ppg dilute $\text{Hg}(\text{NO}_3)_2\text{Aq}$, containing only a little free acid, with very dilute NH_4Aq , boiling with water till the pp becomes compact and curdy, washing, and drying.

The loose flocculent pp produced by adding very dilute NH_4Aq to dilute $\text{Hg}(\text{NO}_3)_2\text{Aq}$, sometimes known as *Mitscherlich's ammoniacal salt*, is probably a *double compound of mercurioxy ammonium nitrate and mercuri ammonium nitrate*, ($\text{NH}_2\text{Hg}_2\text{O})\text{NO}_3 \cdot (\text{NH}_2\text{Hg})\text{NO}_3 \cdot \text{H}_2\text{O}$.

Other compounds are obtained by boiling *Mitscherlich's salt* with excess of NH_4Aq with addition of $\text{NH}_4\text{NO}_3\text{Aq}$, and by dissolving *Soubiran's salt* in hot conc $\text{NH}_4\text{NO}_3\text{Aq}$, the salts thus formed are probably *double compounds of mercurioxy ammonium nitrate and ammonium nitrate*, but their exact composition and constitution are not yet decided (*v* *Mitscherlich, P* 9, 387, 16, 41, *Millon, A Ch* [3] 18, 392, *Kane, A Ch* [2] 72, 215, *Nessler, J* 1856 409).

Mercurioxy-ammonium sulphate, ($\text{NH}_2\text{Hg}_2\text{O})\text{SO}_4$ (*Hydrated tetramercurammonium sulphate*, $\text{N}_2\text{Hg}_4\text{SO}_4 \cdot 2\text{H}_2\text{O}$ *Hydrated dimercuri-ammonium sulphate*,

(NH_4), SO , $2\text{H}_2\text{O}$ *Ammoniacal turpethum*) Obtained by saturating conc NH_4Aq with HgSO_4 , and allowing to evaporate in air, or over H_2SO_4 , also by digesting the basic sulphate $\text{HgSO}_4 \cdot \text{HgO}$ (*mineral turpethum*) with NH_4Aq , also by digesting HgO with hot (NH_4), SO_4Aq (Millon, *A Ch* [3] 18, 392, Ullgren, *P* 42, 395, Hirzel, *J* 1852 419, 1853 381) Schmieder (*J pr* 75, 147) obtained the salt by repeatedly treating mercuri-diammonium sulphate, $\text{N}_2\text{H}_5\text{HgSO}_4 \cdot \text{H}_2\text{O}$ (*v p* 210), with boiling water, till the filtrate was free from sulphates. Forms a heavy yellow white powder, sl sol water, easily sol HClAq or HNO_3Aq . Decomposed by heat to H_2O , N , NH_3 , and Hg_2SO_4 . Treated with KOH Aq , gives NH_3 and a *basic salt* (NH_4HgO), SO_4HgO (perhaps $\text{NH}_4\text{Hg}_2\text{ONH}_2\text{Hg}_2\text{O SO}_4$), and on further treatment with KOH Aq yields NH_3 , HgO , and K_2SO_4 .

Besides the mercurous ammonium salts described above, the following have been prepared —

Mercurous ammonium bromate ($\text{NH}_4\text{Hg}_2\text{OBrO}_3$, (Rammelsberg, *P* 55, 82)

Mercurous ammonium carbonate ($\text{NH}_4\text{Hg}_2\text{OCO}_3$, (Millon, *A Ch* [3] 18, 392, Hirzel, *J* 1852 419, 1853 381)

Mercurous ammonium chromate ($\text{NH}_4\text{Hg}_2\text{OCrO}_4$, and a *double basic salt* ($\text{NH}_4\text{Hg}_2\text{OCrO}_4 \cdot 3\text{HgCrO}_4$, HgO (Hirzel, *lc*))

Acid mercurous ammonium fluoride ($\text{NH}_4\text{Hg}_2\text{OFHF}$ (Finkener, *P* 110, 632)

Mercurous ammonium - ammonium iodate ($\text{NH}_4\text{Hg}_2\text{OIO}_3 \cdot 2\text{NH}_4\text{IO}_3$, (Millon, *A Ch* [3] 18, 410)

Mercurous ammonium - mercuric phosphate ($\text{NH}_4\text{Hg}_2\text{O}$) Hg PO_4 , (Hirzel, *J* 1852 419, 1853 381)

Mercurous - ammonium - mercuric sulphite ($\text{NH}_4\text{Hg}_2\text{O}$), SO_3 , HgSO_3 , (Hirzel, *lc*)

M M P M

MERCURIALINE The volatile base obtained by distilling the seeds of *Mercurialis annua* or *M perennis* with lime and water, and formerly called 'mercurialine,' is identical with methylamine (E Schmidt, *B* 10, 2226, A 193, 73, cf Reichardt, *J pr* 104, 301)

MERCURY (*Quicksilver*) Hg At w 199.8 Mol w 199.8 [-38.5°] (Regnault, *Acad* 26, 525) [-38.85°] (Mallet, *P M* [5] 4, 145) (357.25° at 760 mm) (Regnault, *lc*), for table of B P of Hg at pressures from 123 to 798 mm *v* Ramsay & Young, *C J* 47, 656) S G $\frac{1}{4}$ 13.5958 to 13.596 (Regnault), $\frac{10}{15}$ 13.546 (Volkman, *W* 13, 209, a very full table of S G and volume of Hg from 0° to 360° is given) S G solid Hg , $^{-28.5^\circ}$ = 14.1932 (Mallet, *P M* [5] 4, 145) V D 99.3 at 440° to 1565° (V Meyer, *B* 12, 1426) S H 03312 from 20° to 50° , 03278 from 25° to 142° (Winkelmann, *P* 159, 152, *v* also Pettersson, *B* 12, 1718)

S H p

$\frac{\text{S H v}}{\text{S H v}}$ 275° to 355° = 1.666 (Kundt & Warburg, *P* 157, 353, experimentally determined) C E 00018153 from 0° to 100° (Regnault, *Acad* 21, 271, *v* also Wüllner, *P* 153, 440) T C (Ag = 100) 5.3 (Calvert & Johnson, *P M* [4] 16, 381, cf Weber, *W* 10, 490) E C c 196 (Cu = 100) For electrical resistance of Hg *v*

Kohlrausch, *W* 35, 700, (cf Guinmach, *W* 37, 508) For table of vapour pressures of Hg from 135° to 520° *v* Ramsay & Young, *C J* 49, 37 Heat required to change 200 grm Hg at 358° to gas = 12,400 gram calories, heat required to melt 200 grm solid Hg = 564 (Person, *A Ch* [3] 24, 257, *ibid* 21, 295) Chief lines in emission spectrum 6151, 5460.5, 4358 (Thalén, 1868, for table of lines of high refrangibility *v* Hartley & Adeney, *T* 1884 136) S V S c 14 Crystal luses in regular octahedra

Occurrence—Small quantities of Hg are found as globules disseminated through ores of Hg , or in the rocks in which these ores occur. The chief ore of Hg is *cinnabar*, HgS , found in limited quantities in Illyria, Spain, Bohemia, Peru, China, California, the Ural, and one or two other localities. An amalgam of Hg and Ag is sometimes found native, iodide, chloride, and minute quantities of selenide, of Hg are also known to occur

Preparation—1 By distilling *cinnabar* in a regulated supply of air, condensing the Hg in a suitable apparatus, and allowing the SO_2 to escape—2 By heating *cinnabar* with lime, CaS and CaSO_4 are formed, and the Hg distils and is condensed—3 By heating *cinnabar* with iron (smithy scales), Hg , FeS , and SO_2 are produced, the Hg is condensed in long chambers, or by downward distillation in a trough filled with water

To purify Hg from dust, &c, it may be filtered through a paper cone having a minute hole at the apex, or pressed through leather. Traces of metals in solution may be removed by leaving the Hg in contact with cold conc H_2SO_4 for some weeks, with frequent agitation, and then digesting with very dilute HNO_3Aq , or with HgNO_3Aq (Branchi, *R P* 6, 77, Wittstein, *R P* 65, 362) Karsten and Ulex (*A* 60, 210) recommend to rub the Hg in a basin for ten minutes, with 1/60th of its weight of FeCl_3Aq , S G 1.48 diluted with an equal weight of water, to pour off the liquid, which contains the foreign metals and some HgCl_2 , to wash with water, and then to dry the Hg by heating it. L Meyer (*B* 12, 437) has described a convenient apparatus for purifying Hg by bringing a fine stream of it into contact with FeCl_3Aq Bruhl (*B* 12, 204) shakes Hg with an equal volume of chromate solution, made by dissolving 5 grams $\text{K}_2\text{Cr}_2\text{O}_7$ in 1 litre water, and adding a few c.c. of H_2SO_4 , the shaking is continued until the red HgCrO_4 at first produced has disappeared, and the liquid is coloured green by $\text{Cr}_2(\text{SO}_4)_3$, the fine greyish powder which is formed, and which consists of oxides of foreign metals, is washed away by a rapid stream of water. The process is repeated, and the Hg is then shaken with pure water until no more greyish powder is formed. About half p.c. of Hg is lost. Crafts (*B* [2] 49, 856) removes Pb, Zn, Sn, and most other metallic impurities from Hg by placing the Hg in a slightly inclined glass tube, and aspirating a gentle current of air through the apparatus for about forty eight hours. The oxides of the foreign metals collect at the top. Ag cannot be removed by this method

Pure Hg may be obtained by distilling equal parts of *cinnabar* and burnt lime or Fe filings. Also by boiling pure HgCl_2Aq with pure Fe .

Also by heating HgO in a retort, and digesting the residual Hg , which contains a little HgO , with dilute HNO_3Aq or H_2SO_4 . Millon (*B J* 27, 110) prepares HgO (from which to make Hg) by shaking Hg for some time with dilute HNO_3Aq to remove foreign metals, washing, dissolving in such a quantity of HNO_3Aq that 1/10th of the Hg remains undissolved, evaporating to dryness, and heating.

After purifying Hg by one of the foregoing methods it is advisable to distil it. A very convenient apparatus for distilling Hg under reduced pressure is described by Wright (*Am S* [3] 22, 479, v also Weber, *Carl Rep* 15, 52, Weinhold, *Carl Rep* 15, 1).

Properties—A very lustrous metal, white, with the slightest tinge of blue. Exists as a very mobile liquid at ordinary temperatures, and does not solidify above -38.5° . Hg boils at 360° , but it is very easily volatilised, even at -13° sufficient Hg is vaporised to produce a daguerreotype by twenty four hours' exposure (Regnault, *C R* 73, 141/2). According to Merget (*C R* 73, 1386) solid Hg at -44° is very slowly vaporised. Solid Hg is a tin-white, ductile mass of octahedral and needle shaped crystals. It may be cut with a knife. Hg is a very coherent liquid. By trituration with sugar, grease, sulphur, chalk, &c., or by agitation with ether, turpentine, and some other liquids, Hg appears as a grey powder, which consists of minute globules of Hg separated by the foreign matter present (*extinction or deadening of mercury*). Finely divided Hg is also obtained by mixing about equal parts of SnCl_2 , dissolved in HClAq , and HgCl_2 , dissolved in hot water ($\text{SnCl}_2\text{Aq} + \text{HgCl}_2\text{Aq} = \text{SnCl}_4\text{Aq} + \text{Hg}$). In this state of fine division, Hg is slowly oxidised by contact with air. Pure Hg adheres very slightly to glass. When foreign metals are present the Hg leaves a blackish film on glass. A globule of pure Hg should roll down a slightly inclined surface without losing its round form, and without leaving any streak behind it. When shaken in a bottle with dry air it should not form any black powder.

Pure Hg is not oxidised by exposure to air at ordinary temperatures, when heated to near its B.P. HgO is slowly formed [Berthelot (*C R* 91, 871) asserts that pure Hg is very slightly oxidised in air]. Hg obtained in a state of fine division by trituration, e.g. with chalk, is slowly oxidised at the ordinary temperature. Ozone oxidises Hg at ordinary temperatures (Volta, *G* 9, 521, cf *Reactions*, No 2). Hg combines directly with O , S , and the halogens, it alloys with most of the metals, in some cases forming definite compounds (v *Mercury, amalgams of*). Hg reacts with hot conc H_2SO_4 to produce HgSO_4 and SO_2 , with HNO_3Aq it produces HgNO_2 , $\text{Hg(NO}_3)_2$, or a basic nitrate, according to the temperature and the quantity of acid. HClAq does not react with Hg .

The atomic weight of Hg has been determined (1) by analysing HgO^f (Sefström, *S* 23, 328, Turner, *A* 13, 18, Erdmann & Marchand, *J pr* 31, 395), (2) by determining Cl in HgCl and HgCl_2 (Turner, *lc*, Svanberg, *J pr* 45, 468, Millon, *A Ch* [3] 18, 345), (3) by estimating Hg in HgS (Erdmann & Marchand, *J pr* 31, 400), (4) by determining S in Hg .

Hg is distinctly a metallic element. It forms salts by replacing the H of most acids. These salts belong to two series, HgX and HgX_2 , where $\text{X} = \text{NO}_2$, ClO_2 , $\frac{1}{2}\text{SO}_3$, PO_3 , &c. Both series are well represented by definite and stable salts. HgO dissolves in molten KOH , and crystals of $\text{K}_2\text{O HgO}$ are obtained on cooling. The compound $\text{Na}_2\text{O HgO}$ is said also to exist. HgS dissolves in NaHSaAq and conc KHSaAq . The compound $\text{K}_2\text{S HgS} \cdot 5\text{H}_2\text{O}$ has been isolated. The existence of these compounds shows that HgO and HgS are slightly acidic towards K_2O and KHS .

The molecular weight of Hg has been directly determined, the molecule is monatomic. The atom of Hg is divalent in the gaseous molecules HgCl_2 , HgBr_2 , and HgI_2 . The molecular formula of calomel is probably HgCl , if this is so, the atom of Hg is monovalent in this molecule.

Hg is related to Mg and Zn , and more distantly to Be , Ca , Sr , and Ba (v **MAGNESIUM GROUP OF ELEMENTS**, p 165, and cf **CLASSIFICATION**, vol II pp 204, 207).

Reactions—1 Heated in *air* or *oxygen*, to near its B.P., Hg slowly forms HgO .—2 *Ozone* produces some Hg_2O at ordinary temperatures, perfectly dry ozone does not react with dry Hg (Shenstone & Cundall, *C J* 51, 619).—3 Hg is oxidised by agitation with solution of *potassium permanganate*, Hg_2O is formed if the solution is cold, and HgO if hot (Kirchmann, *Ar Ph* [2] 150, 203).—4 Hg does not react with *hydrochloric acid*.—5 *Sulphuric acid*, when hot and conc, forms HgSO_4 , or Hg_2SO_4 , if there be an excess of Hg and the acid is not very hot, and SO_2 , dilute $\text{H}_2\text{SO}_4\text{Aq}$ scarcely acts on Hg .—6 *Nitric acid*, cold and dilute, forms HgNO_2 , with hot HNO_3Aq and excess of Hg , basic mercurous nitrates, $x\text{Hg}_2\text{O} \cdot y\text{N}_2\text{O}_5$, are formed, excess of hot HNO_3Aq produces $\text{Hg(NO}_3)_2$.—7 Warm conc *hydrobromic* and *hydrochloric* acids produce HgBr_2 and HgI_2 , respectively (cf *Berthelot*, *A Ch* [6] 16, 433).—8 *Hydrogen sulphide*, or *alkaline polysulphides*, produce HgS .—9 With *nitrogen tetroxide*, forms HgNO , and NO (Ramsay, *B* 13, 3154).

Combinations—1 Hg combines with the *halogens*, forming HgX or HgX_2 , according to the relative quantities of the reacting bodies.—2 With *oxygen*, Hg forms Hg_2O and HgO .—3 With *sulphur* and *selenium*, HgS and HgSe are produced.—4 Hg probably combines with *phosphorus*, but no definite compound has been isolated with certainty.

Detection and Estimation—Solid Hg compounds are decomposed by drying, mixing well with dry Na_2CO_3 , and heating strongly in a small tube closed at one end, Hg is volatilised, and condenses on the cold part of the tube in minute globules. Mercurous salts in solution give a white pp (HgCl) with HClAq , or a soluble chloride, this pp is blackened by NH_3Aq with formation of $\text{NH}_2\text{Hg}_2\text{Cl}$. Mercurous salts are oxidised to mercuric salts by boiling with HNO_3Aq . Mercuric salts are not pptd by HClAq , H_2SAq , or a soluble sulphide, ppts black HgS , addition of a small quantity of the reagent causes the formation of a compound of HgS with the original salt present, this pp is white, yellow, or brown, according to the quantity of reagent used.

Hg is sometimes estimated by heating its dry compounds with powdered CaO, condensing the Hg under water, transferring to a weighed crucible, drying by blotting paper and then over H_2SO_4 , the metal is sometimes pptd by SnCl_2 or H_3PO_4 at c 60° , dried, and weighed. Hg is also estimated as HgCl_2 by adding HClAq , then nearly neutralising with KOH Aq , adding K or Na formate, digesting for some days at 60° – 70° (Hg is pptd at boiling temperature), collecting the HgCl_2 on a weighed filter, washing, drying at a low temperature and weighing. Hg is also determined as HgS , by pptn from HgCl_2Aq by H_2S , and collection on a weighed filter. If the ppt is formed from solutions containing HNO_3Aq or FeCl_2Aq &c, free S is pptd with the HgS , the ppt may then be heated with CaO and the Hg collected, or heated in Cl, the HgCl_2 passed into water, and Hg pptd by SnCl_2 . Hg may also be determined by depositing it by electrolysis on a weighed Pt dish. A mercurous salt, in presence of mercuric, may be estimated by ppg by HClAq , from a cold very dilute solution, not more than a very small quantity of HNO_3 being present.

There are several methods for the volumetric determination of Hg, that described by Hannay, which is based on the fact that the turbidity produced in mercuric salt solutions by a few drops of NH_3Aq is removed by a definite quantity of KCN Aq , is said to give satisfactory results. (For details of the methods *see Manuals of Analysis*.)

Mercury, alloys of, *v* next line

Mercury, amalgams of The alloys of Hg are usually called *amalgams*, the word is supposed to be an alchemist's anagram of $\mu\lambda\alpha\gamma\mu\alpha$ = a soft mass. Amalgams are known of most of those metals which have been fairly well studied. Amalgams are formed by mixing the other metal with Hg, by ppg the other metal on Hg (or Hg on the other metal), and by placing the other metal in contact with Hg and a dilute acid. Many amalgams are definite compounds, others seem to be merely solutions of metals in Hg, and others are probably solutions of definite compounds in excess of Hg. Amalgams of Bi, Pb, Sn, and Zn, for instance, retain Hg at 360° (B P of Hg), but not at 410° . Crystalline amalgams of K and Na, K₂Hg and Na₂Hg, are obtained by heating solutions of K and Na in Hg to 440° .

The best known amalgams are those of Al, Ba, Bi, Cd, Cs, Ca, Cr, Co, Cu, Au, Fe, Pb, Mg, Mn, Ni, Os, Pd, Pt, K, Ag, Na, Sr, Ti, Sn, and Zn. For brief accounts of these amalgams, with the exception of those of Cu which are described below, *v* the various metals, **ALUMINUM, BARIUM, &c**, the articles, **AMALGAMS**, vol 1 p 149, and **ALLOYS**, vol 1 p 132, should be consulted. An interesting paper on *Amalgams*, by Dudley, will be found in *Proc Amer Ass for Advancement of Science*, 1890 145.

COPPER AMALGAMS (should have been described under **COPPER, ALLOYS OF**, vol 1 p 254). Amalgams of Cu are formed by placing Cu foil in HgNO_3Aq , by triturating 3 parts Hg with a mixture of 1 part very finely divided Cu and a few drops of HgNO_3Aq , by triturating Hg with NaCl and verdigris, by making Hg the negative electrode during electrolysis of CuSO_4Aq , by immersing Na-amalgam, or Zn amalgam, in

CuSO_4Aq , and in some other ways. By dissolving Cu in Hg, and removing excess of Hg by pressure, Joule obtained an amalgam having the composition CuHg (*Chem Gazette*, 1850 339). By heating amalgams rich in Hg, De Souza obtained Cu_2Hg at c 410° , and Cu_3Hg at c 360° (*B* 9, 1050). By ppg Cu from CuSO_4Aq by Fe, washing well, moistening with HgNO_3Aq , and rubbing with Hg, amalgams are obtained which harden after a time. The amalgam with from 25 to 33 p.c. Cu, may be obtained soft and plastic by heating nearly to 360° , and kneading for some time in a warm mortar, this amalgam slowly hardens without either contracting or expanding, it is used for stopping teeth (*v* Von Gersheim, *A* 70, 344).

Mercury, ammonio compounds of; *v* **MERCURAMMONIUM COMPOUNDS**, p 206

Mercury, antimonides of, *v* vol 1 pp 285–6

Mercury, antimonide of A substance of the nature of an amalgam is formed by triturating together Sb and Hg, little is known of its composition and properties, it is very easily decomposed.

Mercury, arsenates of, *v* vol 1 p 309

Mercury, arsenide of Bergmann obtained what he thought was a compound of 1 pt As with 5 pts Hg by heating Hg and As together.

Mercury, arsenites of, *v* vol 1 p 306

Mercury, bromides of Two bromides of Hg are known, HgBr and HgBr_2 . Mercuric bromide has been vapourised, its mol w is 359.3 = HgBr_2 . Mercurous bromide has also been vapourised, the results indicate the mol w 279.55 = HgBr , but it is not certain whether partial dissociation into HgBr , and Hg occurred during vapourisation (*v* *infra*, also *Mercurous chloride*, p 215). Both HgBr and HgBr_2 form double compounds with the bromides of metals more positive than Hg.

MERCUROUS BROMIDE HgBr (*Mercury proto or mono bromide*). Mol w 279.55 (?). H F [HgBr] = 25.475 (Thomsen, *Z P C* 2, 21).

Preparation—1 By subliming an intimate mixture of 80 pts Br and 200 pts Hg—2 By adding KBrAq to HgNO_3Aq —3 By heating saturated HgNO_3Aq , as free as possible from HgO and slightly acidified with HNO_3Aq , with Br, decanting, and allowing to cool in the dark (Stroman, *B* 20, 2818)—4 By adding alcoholic solution of Br to cold conc HgNO_3Aq (Stroman, *l c*).

Properties and Reactions—As prepared by method 3 (*supra*), forms small, white, nacreous, tetragonal, scales, prepared by method 2, appears as yellow, crystalline, spangles (Stroman, *l c*). SG 7.307 (Karsten, *S* 65, 394). Sublimes at 340° – 350° . Melts c 405° (?) (Carnelley, *C J* 33, 277). VD 146.3 (according to Mitscherlich (*P* 29, 193)). Tasteless and odourless. Insol in water. Hot conc H_2SO_4 dissolves HgBr with evolution of SO_2 , dissolves slowly in hot HNO_3Aq SG 1.42, decomposed by hot HClAq dil or conc, decomposed by NH_3Aq or KOH Aq , forming NH_4Br or KBr , and leaving a black residue which contains Hg (Stroman, *l c*). Slowly decomposed by boiling KBrAq , giving HgBr_2 and Hg. Soluble in some NH_4 salt solutions.

Combinations—Two compounds with *strontium chloride* are described by Löwig (*P* 14, 485). By dissolving HgBr in boiling SrBr_2Aq , allowing

to cool, pouring off from separated HgBr , and evaporating, crystals of $6\text{HgBr}\cdot\text{SrBr}$, separate, by dissolving these crystals in water, filtering from HgBr which separates, and evaporating, a very soluble crystalline compound $2\text{HgBr}\cdot\text{SrBr}$, is obtained

MERCURIC BROMIDE HgBr_2 (*Dibromide of Mercury*) Mol w 359.3 HF [HgBr_2] = 41.880 (Thomsen, *Z P C* 2, 21)

Preparation — 1 By dissolving HgO in HBrAq , and crystallising — 2 By bringing Hg into slight excess of Br , warming to remove uncombined Br , and subliming — 3 By heating an intimate mixture of HgSO_4 with rather more than its own weight of KBr , the HgBr_2 , which sublimes is said to contain HgBr — 4 By adding KBrAq to $\text{Hg}(\text{NO}_3)_2\text{Aq}$, evaporating as long as HgBr_2 crystallises out, and crystallising from alcohol — 5 By agitating Hg with water, adding Br as long as its colour is removed, boiling, filtering, allowing to crystallise, and drying at $c\ 160^\circ$ — 6 By adding excess of Br to a slightly acid solution (S G 1 197) of $\text{Hg}(\text{NO}_3)_2$, HBrOAq is formed in the reaction (Sievers, *B* 21, 647)

Properties — White rhombic prisms, isomorphous with HgCl_2 , $a\ b\ c = 6817\ 1\ 9975$ (Hjorthal, *Z K* 3, 362) S G 5 9202 (Karsten, *S* 65, 394), 5 7298 at 16° , 5 7461 at 18° (Clarke's *Specific Gravity Tables*, new ed 32) V D 175.5, Mitscherlich (*P* 29, 193) Melts at 244° (Carnelley a Williams, *C J* 37, 127) Can be sublimed unchanged S 1 06 at 9° , 20–25 at 100° (Lassaigne) Very sol in alcohol and ether HgBr_2Aq reddens litmus

Reactions — 1 Heated with phosphorus, arsenic or antimony, forms bromide of P, As, or Sb — 2 Decomposed, yielding HgBr , by sunlight, or contact with copper or mercury, or by mixing with cuprous bromide dissolved in HBrAq — 3 HgBr_2Aq with ammonia gives a white pp of $\text{Hg}_2\text{Br}_2\text{NH}_3$ (Mitscherlich, *J pr* 19, 455) — 4 With sodium hypochlorite, HgBr_2Aq gives oxychlorides $x\text{HgO}\cdot y\text{HgCl}_2$ (Rammel-berg) — 5 With hypochlorous acid, HgCl_2 and $\text{Hg}(\text{BrO})_2$, are formed (Balard) — 6 Decomposed by nitric or sulphuric acid, giving off Br — 7 Oxybromides, $x\text{HgO}\cdot y\text{HgBr}_2$, are obtained by boiling HgBr_2Aq with mercuric oxide — 8 Mixed with mercuric iodide and crystallised from ether or acetone, HgI_2Br is obtained, this compound is also formed by the action of an alkyl iodide, e.g. EtI , on HgBr_2 dissolved in acetone (Oppenheim, *B* 2, 571) (*v Mercury, iodobromide of*, p 221)

Combinations — 1 With hydrobromic acid to form bromomercuric acid HHgBr_2 (*v infra*) — 2 Combines with many bromides of more positive metals than Hg , to form salts regarded by Von Bonsdorff as bromomercurates (*P* 19, 339) These salts are obtained by evaporating solutions of the component bromides, the more important are $\text{HgBr}\cdot\text{KBr}$ and $\text{HgBr}\cdot 2\text{KBr}$, $\text{HgBr}\cdot\text{MgBr}$, and $2\text{HgBr}\cdot\text{MgBr}_2$, $\text{HgBr}_2\cdot\text{SrBr}$, and $2\text{HgBr}_2\cdot\text{SrBr}_2$ (Löwig, *P* 14, 485) Salts containing NaBr , CaBr , BaBr , and ZnBr , were also obtained by Von Bonsdorff (*loc*) Bromomercuric acid, HHgBr_2 , was obtained by Neumann (*M* 10, 286), in long transparent needles, by slightly warming excess of HgBr_2 with HBrAq , filtering through asbestos, and cooling to between 0° and -4° . This acid is

readily decomposed by moisture or heat Solution of HgBr_2 in HBrAq reacts with alkalis to give $\text{HgBr}_2\cdot 2\text{MX}$ ($\text{M} = \text{alkali metal}$), the heat produced is $c\ 27,200$ for 2MOH , the solution of HgBr_2 in HBrAq probably contains the acid H_2HgBr_4 — 3 With mercuric cyanide and potassium cyanide, forms lustrous tablets of $\text{HgBr}_2\cdot\text{HgCy}_2\cdot 2\text{KC}_y\ 4\text{H}_2\text{O}$ (Geuther, *A* 29, 325) With mercuric cyanide and zinc cyanide, forms transparent prisms of $\text{HgBr}_2\cdot\text{HgCy}_2\cdot\text{ZnCy}_2\cdot 8\text{H}_2\text{O}$ (Varet, *C R* 109, 809) — 4 Heated with mercuric sulphide, forms $\text{HgBr}_2\cdot 2\text{HgS}$ (Schneider, *P* 115, 167), this salt is also obtained by boiling HgBr_2Aq with freshly ppd HgS (H Rose, *P* 13, 59) (*v Mercuric sulphide, Combinations*, No 4, p 225)

Mercury, bromide of, *v Mercury, iodobromide of*, p 221

Mercury, bromosulphide of, *v Mercuric sulphide, Combinations*, No 4, p 225

Mercury, chlorides of Two chlorides are known, HgCl and HgCl_2 . The molecular weight of the latter has been determined from the V D of the compound The numbers obtained for the V D of the former agreed with the formula HgCl , but it was shown that the vapour produced by heating mercurous chloride contained Hg and HgCl_2 , hence the V D could not decide between the possible formulæ HgCl and HgCl_2 . More recently it has been found that the V D calculated from observations of V D of mercurous chloride vapourised into an atmosphere of HgCl_2 agrees with that required by HgCl (*v infra*)

MERCUROUS CHLORIDE HgCl (*Protochloride of mercury Calomel*) Mol w very probably 235.17 Sublimes at 400° – 500° without melting S G 6 993 (Karsten, *S* 65, 394), to 7 176 (Hassenfratz, *A Ch* 28, 3) V D 118.6 at 440° (Deville a Troost, *C R* 45, 821), 120.8 (Mitscherlich, *P* 29, 193) V D calculated for $\text{HgCl} = 117.58$ Odling showed that gold leaf is amalgamated when exposed to calomel vapour at $c\ 400^\circ$, hence he concluded that the vapour contained Hg , and that therefore the observed V D did not prove the formula HgCl (*J* 1864 280) Debray found that only a limited portion of the calomel vapourised at 440° is decomposed with separation of Hg (*C R* 83, 330), hence it appears impossible that calomel should have the molecular formula HgCl , else the observed V D at $c\ 440^\circ$ would be considerably greater than 120 (calc for $\text{HgCl}_2 = 235.16$) In 1881, Fielet showed that when a mixture of mercurous and mercuric chlorides is heated to $c\ 400^\circ$ in a Pt tube, containing a tube of silver-gilt traversed by a stream of cold water, not a trace of Hg is deposited on the gilt tube He then determined the V D of a mixture of mercurous and mercuric chlorides, and calculated the V D of the mercurous chloride in the mixed vapours, the results were 115.9 and 120.1, hence the molecular formula of calomel is most probably HgCl (*G* 1881 341) SH at 7° – 99° , 05205 (Regnault, *A Ch* [3] 1, 129) Crystallises in tetragonal forms, $a\ c = 1.17414$ HF [HgCl] = 32,600 (Thomsen, *Z P C* 2, 21) For T C *v* Von Lang, *P* 135, 29, for heat of vaporisation, *v Manganese, C R* 67, 877

Occurrence — As horn quicksilver, in tetragonal crystals. S G 6 482.

Formation.—1 By passing Cl over excess of Hg; action proceeds slowly at ordinary temperatures, rapidly at near B P of Hg.—2 By reducing HgCl_2Aq , saturated at 50° , by SO_2 (Wohler, *A* 90, 124, Sartorius, *A* 96, 325)—3 By reducing HgCl_2Aq by oxalic acid in sunshine (Eder, *B* 13, 166)—4 By shaking Hg with FeCl_2Aq —5 By ppg a mercurous salt by HClAq —6 By heating HgCl_2 with Hg—7 By heating HgSO_4 with Hg and NaCl —8 By triturating Hg with NaCl , $\text{Fe}_2(\text{SO}_4)_3$, and a little water, till the metal has lost its fluidity, and subliming $(\text{Fe}_2(\text{SO}_4)_3 + 6\text{NaCl} + 2\text{Hg} = 3\text{Na}_2\text{SO}_4 + 2\text{FeCl}_2 + 2\text{HgCl})$

Preparation.—1 An intimate mixture of 4 pts HgCl_2 with 3 pts Hg is made by moistening with alcohol and triturating until the Hg loses its fluidity, the mixture is gently heated for a few hours, again pulverised, and then slowly sublimed in a flask or retort, not more than a quarter filled with the mixture and placed on a thin layer of sand.—2 4 pts Hg, 9 pts dry HgSO_4 , and 3 pts water are very intimately mixed, a quantity of NaCl equal to the weight of the Hg and HgSO_4 , used is added, and the mixture is slowly heated till HgCl sublimes (Planche, *A Ch* 66, 168)—3 A well-pulverised mixture of 1 pt NaCl , $1\frac{1}{2}$ pts NaCl , and $2\frac{1}{2}$ pts Hg is heated with 2 pts conc H_2SO_4 , until HgCl sublimes.—4 A warm dilute solution of HgNO_3Aq , mixed with a little HNO_3 , to prevent formation of basic salts, is ppd by excess of dilute NaClAq containing a little HCl , the liquid is heated for some time in contact with the pp which is then thoroughly washed with cold water in the dark (Schæele, *cf* Frautwein, *R* P 11, 72, 12, 155, Mialhe, *J Ph* 22, 586)—5 Cl is passed into HgNO_3Aq , the pp of HgCl mixed with HgCl_2 , is washed with hot water till HgCl_2 is all dissolved, the HgCl is then crystallised from warm HgNO_3Aq (Sievers, *B* 21, 647)

Properties.—White, semi transparent, tetragonal, prisms, if sublimed quickly, forms a fibrous mass of small crystals. When powdered, calomel shows a slight lemon yellow colour. Highly refractive and dispersive. Tasteless and odorless. Sublimes without melting. Almost quite insol water, alcohol, ether, and dilute acids. Sl sol cold, more sol hot, HgNO_3Aq (*v* Debray, *O R* 70, 995, *cf* Dreehsel, *J pr* [2] 24, 44). Boiled with HClAq , HgCl_2 goes into solution, and Hg is separated, with conc hot H_2SO_4 , HgCl_2 and HgSO_4 are formed. HgCl is decomposed by solutions of alkalis, alkaline earths, and alkaline carbonates, with ppn of Hg_2O . For experiments on V D of calomel, *v supra*

Reactions.—1 Exposed to sunlight, HgCl darkens with separation of Hg (for action of heat on HgCl *v supra*)—2 Partly decomposed by aqueous vapour or boiling water, with production of HgCl_2 and Hg—3 Decomposed by many metals when heated with them in presence or absence of water, giving metallic chlorides and Hg—4 Triturated with iodine and water, HgCl_2 and HgI_2 are produced (Planche & Soubeiran, *J Ph* 12, 651)—5 Heated with a little sulphur, HgS and HgCl_2 are formed, with excess of S, the products are HgS and S_2Cl_2 —6 Heated in phosphorus vapour, HgCl yields PCl_2 and Hg phosphite (Davy).—7 By heating an intimate

mixture of 3 pts HgCl with 1 pt arsenic, a sublimate is obtained consisting in part of yellow tetrahedral crystals, according to Capitaine (*J pr* 18, 422) these crystals are *mercuro-arsenic chloride* AsHgCl . The compound is decomposed by hot water to As Hg amalgam, As_2O_3 and HCl , it is partly decomposed by heat, giving Hg, As, and AsCl_3 —8 Reduced by sulphurous acid, also by hot stannous chloride solution, with separation of Hg—9 Cold sulphuric acid has no action on HgCl , the hot conc acid forms HgCl_2 , HgSO_4 , and SO —10 Dissolved by hot nitric acid with evolution of NO , the solution contains $\text{Hg}(\text{NO}_3)_2$ and HgCl_2 —11 Boiling hydrochloric acid forms HgCl_2 and Hg, boiled in an open vessel with water and a little HCl , no Hg is separated, but HgCl_2 goes into solution—12 Decomposed by *hydrocyanic acid* solution, forming HgCy , Hg, and HCl —13 With solution of *alkali chlorides* forms HgCl_2 , which combines with the excess of alkali chloride to form double salts, some of which are soluble and some insoluble—14 Digested with solutions of *alkali iodides*, or *iodide of magnesium*, or *von*, HgI is separated, and metallic chloride goes into solution—15 Solutions of *alkalis* form Hg_2O and alkali chloride, heated with solid alkalis, Hg, O, and alkali chloride are produced—16 *Ammonia* produces black NH_4HgCl (*v* MERCURAMMONIUM COMPOUNDS)—17 Heated with *carbonates of calcium, barium, or magnesium*, Hg, Hg_2O , and HgO are separated, and CO is evolved (Buchner, *R P* 3, 31, 4, 289, Vogel, *R P* [3] 1, 34)—18 With *alkali sulphides* in solution, Hg separates, and a double sulphite of Hg and alkali metal goes into solution—19 Rubbed with *antimony tri* or *penta sulphide*, HgS and SbCl_3 are formed

Testing calomel for impurities.— HgCl some times contains HgCl_2 , HgNO_3 , or $\text{Hg}(\text{NO}_3)_2$, or Hg. HgCl_2 is detected by shaking with water, filtering, and adding SnCl_2Aq , when a white cry pp is produced. Bonnewyn (*Bl* [2] 4, 201) says that $\frac{1}{50000}$ pt of HgCl_2 in HgCl may be detected by immersing a clean knife blade in the calomel moistened with alcohol or ether, if HgCl_2 is present a black spot is formed on the blade. HgNO_3 or $\text{Hg}(\text{NO}_3)_2$ is detected by heating, when NO_2 is evolved. Hg may be detected by examination under the microscope. HgCl may be adulterated with powdered heavy spar, &c., this remains unchanged on sublimation. Adulteration with gum or other organic material is detected by the smell produced on heating.

Combinations.—1 With *sulphur chloride*, to form $2\text{HgCl}_2\text{SCL}_2$, produced by long continued warming HgCl with S_2Cl_2 , also by heating an intimate mixture of 31 pts HgCl_2 and 2 pts S in a dish covered with a funnel, on to which the double compound sublimes, also by heating As_2S_3 with HgCl_2 .

$(7\text{HgCl}_2 + \text{As}_2\text{S}_3 = 2\text{AsCl}_3 + 3\text{HgS} + 2\text{Hg}_2\text{SCL}_2)$
Forms rectangular needles, which melt to a brown liquid, and volatilise without decomposition, decomposed by water (Capitaine, *J pr* 18, 422)—2 With *stannous chloride* to form $2\text{HgCl}_2\text{SnCl}_2$. Produced by heating an amalgam of 8 pts Sn with 1 pt Hg mixed with 24 pts HgCl_2 , and subliming. Forms white dendritic crystals, partially decomposed by heat;

wholly decomposed by water, with formation of SnCl_2 which reduces the HgCl (Capitaine, *J pr* 18, 422)—3 H Rose (*P* 44, 325) states that HgCl combines with sulphur dioxide

MERCURIC CHLORIDE HgCl_2 (*Perchloride of mercury Corrosive sublimate*) Mol w 270.54 [288*] (Carnelley, *C J* 33, 276) (303*) (Carnelley & Williams, *C J* 33, 262) S G 6.223 (Playfair & Joule, *C S Mem* 2, 401), 5.418 (Schroder, *P* 107, 113) V D 136.3 (Mitscherlich, *P* 29, 193) S H 12°-45° 064 (Kopp, *Tr* 155, 71), 13°-98° 06889 (Regnault, *A Ch* [3] 1, 129) S 5.73 at 0°, 6.57 at 10°, 7.39 at 20°, 8.43 at 30°, 9.62 at 40°, 11.34 at 50°, 13.86 at 60°, 17.29 at 70°, 24.32 at 80°, 37.05 at 90°, 53.96 at 100° (Poggiale, *A Ch* [3] 8, 463) S cold alcohol 43.5, boiling alcohol 86.2, S ether 33, S glycerin 7 (Fairley, *M S* [3] 9, 685) Crystallises in two forms belonging to the trimetric system (i) crystals from alcoholic solution, $a:b:c = 7.254:1:1.0686$, (ii) crystals by sublimation, $a:b:c = 9.347:1:3.396$ (v Mitscherlich, *P* 28, 118, Von Lang, *W A B* 45, 119) H F [Hg_2Cl_2] = 64,490 (Thomsen, *Z P C* 2, 21)

Occurrence—In a volcanic district on Ternate, one of the Molucca islands (Frenzel, *Min Metall* 1877 305)

Formation—1 By heating Hg in excess of Cl 2 By dissolving HgO in hot HClAq , and evaporating 3 By dissolving Hg in hot HClAq containing some HNO_3 , evaporating, and recrystallising from hot water—4 By adding excess of Cl to a slightly acid solution of $\text{Hg}(\text{NO}_3)_2$ (Sievers, *B* 21, 617)

Preparation—1 An intimate mixture of equal parts NaCl and Hg_2SO (obtained by boiling 4 pts Hg with 5 pts conc H_2SO_4 till a dry mass is formed), to which a little MnO_2 has been added (to prevent formation of HgCl from Hg_2SO , which may be present), is slowly heated in a long necked flask or retort, HgCl_2 sublimes on the cooler parts of the apparatus. The operation must be conducted in a good draught cupboard because of the very poisonous nature of HgCl_2 —2 Boiling conc HgNO_3Aq is mixed with conc HClAq as long as a pp forms, the pp of HgCl is boiled with as much HClAq as was used in its formation ($\text{HgNO}_3\text{Aq} + 2\text{HCl} = \text{HgCl}_2\text{Aq} + \text{H}_2\text{O} + \text{NO}_2$), crystals of HgCl_2 separate on cooling, they may be recrystallised from hot water

Properties—Translucent crystals, forming a white powder when crushed HgCl_2 has a disagreeable metallic taste, it is extremely poisonous Melts at 288° (*v supra*) to a colourless liquid; according to Haass, HgCl_2 does not melt when heated if the pressure is less than 120 mm (*B* 13, 2203) Easily sol water, sol alcohol, ether, and glycerin (*v supra*) Schröder (*J R* 1886 18) gives the following table, showing the S G of aqueous and alcoholic solutions of HgCl_2 , and the percentage of HgCl_2 in these solutions—

Aqueous solutions of HgCl_2 .

P. HgCl_2 .	S G compared with water at 4°			
	0°	10°	20°	30°
1.22	1.01008	1.0099	1.00836	1.00575
2.434	1.02035	1.02018	1.01856	1.01585
3.578	1.0305	1.03022	1.02855	1.02577
4.725	1.0407	1.04038	1.03856	1.03566

Alcoholic solutions.

P. HgCl_2 .	S G compared with water at 4°				Coefficient of change of S G
	0°	10°	20°	30°	
0	8315	82286	81435	80594	0.00849
1.22	8397	8312	8228	8141	0.00854
2.38	8484	8399	8314	8227	0.00858
4.42	8635	8549	8403	8375	0.00867
8.56	8966	8877	8789	8698	0.00891
2.43	9308	9213	9119	9024	0.00946
15.91	9620	9523	9425	9329	0.00971
19.32	9951	9852	9753	9652	0.00997
22.46	1.0285	1.0184	1.0083	9982	0.01011

HgCl_2Aq reacts slightly acid to litmus On boiling HgCl_2Aq some HgCl passes off with the steam HgCl_2Aq is decomposed by light with evolution of O and formation of HCl and Hg oxychloride (*cf* V Meyer, *B* 20, 2970)

Reactions—1 Many metals decompose HgCl_2 when heated with it, forming HgCl or Hg , such metals are Sb, As, Bi, Cu, Fe, Pb, Ni, Sn Many metals also reduce HgCl_2 in aqueous solution, generally ppg HgCl and Hg , Cd, Fe, and Zn pp Hg only—2 Reducing agents, e.g. SnCl_2Aq , SO Aq , formic acid, oxalic acid, produce HgCl (*cf* I der, *B* 13, 166, also Wurtz's *Dictionnaire*, *Suppl* p 1066)—3 With many carbon compounds, HgCl_2 forms H_2Cl and HCl (the H being withdrawn from the C compound)—4 HgCl_2 is not decomposed by sulphuric or nitric acid, it dissolves fairly easily in HNO_3Aq —5 Dissolves in hot conc hydrochloric acid, on cooling, a white crystalline mass is obtained which has the composition 2HgCl HCl according to Boullay (*A Ch* 34, 243, *cf* *Combinations*, No 1)—6 According to Fairley (*B A* 1875 42), hypochlorites reduce HgCl_2 to HgCl —7 Albumen is rapidly coagulated and ppd by HgCl_2Aq , white of egg serves as an antidote to poisoning by corrosive sublimate—8 Boiled with mercuric oxide, oxychlorides are obtained (*v* Mercury, oxychlorides or, p 223)—9 Alkalis produce HgO and alkali chloride—10 Normal potassium or sodium carbonate ppts HgO , (*K* or *Na*) HCO_3 is also formed, and this reacts with the rest of the HgCl_2 to pp oxychlorides—11 Potassium (or sodium) hydrogen carbonate produces various oxychlorides (*q v* p 223)—12 Phosphorus trihydride produces a black pp when passed into HgCl_2 dissolved in water or alcohol, on continuing the passage of PH_3 , the pp becomes yellow, and then has the composition $\text{P.Hg}, 3\text{HgCl}_2, 3\text{H}_2\text{O} = 2(\text{PHg}_2\text{Cl HgCl}_2) 3\text{H}_2\text{O}$, the solution contains HCl This compo is known as *dimercuriophosphonium mercuric chloride*, it must be washed quickly with cold water, and dried over H_2SO_4 *in vacuo*, it is decomposed by hot water, giving Hg , HCl , and H_3PO_4 , heat produces HCl , Hg , and H_3PO_4 (H Rose, *P* 40, 75)—13 Arsenic trihydride produces a brown yellow pp in HgCl_2Aq This pp has the composition $\text{AsHg}_2\text{Cl HgCl}_2$, it is known as *dimercuriarsonium-mercuric chloride* The pp is washed with cold water and quickly dried over H_2SO_4 *in vacuo*, it is decomposed by water to Hg , As_2O_3 , and HCl (H Rose, *P* 51, 123)

Combinations—1 With hydrochloric acid, according to Boullay (*A Ch* 34, 243) the compound 2HgCl HCl is obtained by dissolving HgCl_2 in hot conc. HClAq , and cooling, this

compound forms a crystalline lustrous mass which melts by the heat of the hand (v also Ditte, *O. R.* 92, 853) Neumann (*M* 10, 236) obtained *chloromercuric acid* HHgCl_2 by gently warming excess of HgCl_2 with HClAq , filtering through asbestos, and cooling to between 0° and -4° . This acid is very readily decomposed by heat or moisture. Solutions of HgCl_2 in HClAq react with alkalis to form $\text{HgCl}_2 \cdot 2\text{MX}$ (M = alkali metal), the heat produced in the reaction is c 27,200 for 2MOH , the solution of HgCl_2 in HClAq probably contains the acid H_2HgCl_2 . Dry HCl passed over well-cooled HgCl_2 has no action (*N*, *lc*) — 2 With many chlorides and with some other salts. The chief compounds with other chlorides are the following (v Von Bonsdorff, *P* 17, 115) —

(i) With phosphorus pentachloride $3\text{HgCl}_2 \cdot 2\text{PCl}_5$. Pearl white needles, sublimes unchanged, decomposed by water, formed by heating the constituents together (Baudrimont, *A Ch* [4] 2, 45)

(ii) With alkali chlorides (a) With potassium chloride $2\text{KCl} \cdot \text{HgCl}_2 \cdot \text{H}_2\text{O}$, $\text{KCl} \cdot \text{HgCl}_2 \cdot \text{H}_2\text{O}$, $\text{KCl} \cdot 2\text{HgCl}_2 \cdot 2\text{H}_2\text{O}$. The first salt is formed by saturating KClAq with HgCl_2 at 30° , adding as much KCl as originally used, and evaporating. The second salt is produced by evaporating a solution of equal equivalents of KCl and HgCl_2 (Rammelsberg, *P* 90, 34). The third salt results by saturating KClAq at 60° with HgCl_2 and allowing to cool. The compound $6\text{KCl} \cdot 3\text{HgCl}_2 \cdot \text{CuCl}_2 \cdot 2\text{H}_2\text{O}$ was obtained by Von Bonsdorff (*P* 33, 81) by adding CuCl_2Aq to a dilute solution of $\text{KCl} \cdot \text{HgCl}_2$. (b) With sodium chloride $2(\text{NaCl} \cdot \text{HgCl}_2) \cdot 3\text{H}_2\text{O}$, $2\text{NaCl} \cdot \text{HgCl}_2$. The first salt is obtained by adding NaCl to NaClAq saturated with HgCl_2 . The second salt is produced by adding alcohol to saturated NaClAq shaken with powdered HgCl_2 . (c) With rubidium chloride $2\text{RbCl} \cdot \text{HgCl}_2$, $\text{RbCl} \cdot 2\text{HgCl}_2$. Obtained, respectively, by evaporating a neutral solution of RbCl and HgCl_2 , by evaporating a solution of equal parts of the constituent salts, and by heating the second salt to 50° (Godeffroy, *Ar Ph* [3] 12, 47). (d) With ammonium chloride $2\text{NH}_4\text{Cl} \cdot \text{HgCl}_2 \cdot \text{H}_2\text{O}$ (*sal alembrothi*) is obtained by evaporating a solution of 1 pt NH_4Cl and 2 pts HgCl_2 , loses H_2O in dry air, isomorphous with the corresponding K salt. The other salts obtained are $\text{NH}_4\text{Cl} \cdot \text{HgCl}_2$, $2\text{NH}_4\text{Cl} \cdot 2\text{HgCl}_2 \cdot 2\text{H}_2\text{O}$ (Kane), $2\text{NH}_4\text{Cl} \cdot 9\text{HgCl}_2$, and $2\text{NH}_4\text{Cl} \cdot 3\text{HgCl}_2 \cdot 4\text{H}_2\text{O}$ (Holmes, *O N* 5, 351).

(iii) With alkaline earth chlorides (a) With calcium chloride $\text{CaCl}_2 \cdot 5\text{HgCl}_2 \cdot 8\text{H}_2\text{O}$, by saturating warm CaCl_2Aq with HgCl_2 , and cooling, by evaporating the mother liquor the salt $\text{CaCl}_2 \cdot 2\text{HgCl}_2 \cdot 6\text{H}_2\text{O}$ is obtained. (b) With strontium chloride $\text{SrCl}_2 \cdot 2\text{HgCl}_2 \cdot 2\text{H}_2\text{O}$. (c) With barium chloride $\text{BaCl}_2 \cdot 2\text{HgCl}_2 \cdot 2\text{H}_2\text{O}$. (d) With magnesium chloride $\text{MgCl}_2 \cdot 3\text{HgCl}_2 \cdot 5\text{H}_2\text{O}$ and $\text{MgCl}_2 \cdot \text{HgCl}_2 \cdot 6\text{H}_2\text{O}$.

(iv) With chloride of zinc $\text{ZnCl}_2 \cdot \text{HgCl}_2 \cdot 4\text{H}_2\text{O}$, by evaporating a mixed solution of the constituents. Analogous salts with CuCl_2 , CoCl_2 , NiCl_2 , and FeCl_2 have been obtained.

The chief compounds with oxyacids are the following. —

(i) With ammonium sulphite

$3\text{HgCl}_2 \cdot 2(\text{NH}_4)_2\text{SO}_3$. By adding hot saturated HgCl_2Aq to cold $(\text{NH}_4)_2\text{SO}_3\text{Aq}$ (*P de Saint-Gilles*, *A* 84, 266, 269).

(ii) With potassium chromate, and dichromate (a) $\text{HgCl}_2 \cdot \text{K}_2\text{Cr}_2\text{O}_7$, by cooling a warm solution of equivalents of the two salts (Millon, *A Ch* [8] 18, 388, Darby, *C S Mem* 1, 24).

(b) $2\text{HgCl}_2 \cdot \text{K}_2\text{CrO}_4$, by mixing solutions of the constituent salts in the ratio $2\text{HgCl}_2 \cdot \text{K}_2\text{CrO}_4$, adding HClAq sufficient to dissolve ppd HgCrO_4 , and evaporating (Darby, *lc*).

(iii) With ammonium dichromate (a) $\text{HgCl}_2 \cdot (\text{NH}_4)_2\text{Cr}_2\text{O}_7 \cdot \text{H}_2\text{O}$, by evaporating a solution of the constituents (Abel & Richmond, *C J* 3, 202, cf Darby, *C S Mem* 1, 24, Zephorovich, *W A B* 39, 17). (b) $\text{HgCl}_2 \cdot 3(\text{NH}_4)_2\text{CrO}_4$, by evaporating the mother liquor from the first salt (*A A R*, *lc*; cf Clarke & Stern, *Am. Ch* 3, 351).

(iv) With copper acetate

$2\text{HgCl}_2 \cdot \text{Cu}(\text{C}_2\text{H}_3\text{O}_2)_2 \cdot \text{CuO}$, deposited from a mixture of cold saturated solutions of HgCl_2 and $\text{Cu}(\text{O}_2\text{H})_2$ (Hutteroth & Wöhler, *A* 53, 142).

(v) With chlorochrometetrammonium chloride $6\text{HgCl}_2 \cdot \text{Cl}_2\text{Cr}_2\text{S}_2\text{NH}_4 \cdot \text{Cl}_4$, by spontaneous evaporation of a mixed solution of the two salts (Clève, *J* 1862 150).

3 Compounds of HgCl_2 with PHgCl and AsHgCl are known (a) *Dimercuriphosphonium mercuric chloride* $2(\text{PHgCl} \cdot \text{HgCl}_2) \cdot 3\text{H}_2\text{O}$, a yellow solid formed by passing PH_3 into HgCl_2 dissolved in water or alcohol (v *Reactions*, No 12).

(b) *Dimercuriarsonum - mercuric chloride* $\text{AsHg}_2\text{Cl} \cdot \text{HgCl}_2$, a brown yellow pp by passing AsH_3 into HgCl_2Aq (v *Reactions*, No 13) — 4 A compound of HgCl_2 with *mercuric sulphocyanide* — $\text{HgCl}_2 \cdot \text{Hg}(\text{CSN})_2$ — is obtained by the reaction of SOCl_2 with $\text{Hg}(\text{SCN})_2$ (McMurtroy, *C J* 55, 50).

Mercury, chlorocyanide of, v Mercury, cyanochloride of, infra

Mercury, chlorosulphides of, v Mercuric sulphide, Combinations, No 4, p 225

Mercury, chloro-iodides of, v Mercury, iodochlorides of, p 221

Mercury, chromates of, v vol II p 155

Mercury, cyanides of, v vol II p 312

Mercury, cyanochloride of, H_2CyCl Quadratic prisms, stable in air, produced by evaporating an aqueous solution of equivalent weights of HgCl_2 and HgCy_2 (Liebig, *S* 49, 253, Poggiale, *C R* 23, 762, cf Weeren, *J pr* 64, 63).

Mercury, cyano-selenides and cyano-sulphides of, v Mercury, selenocyanides and sulphocyanides of, under CYANIDES, vol II pp 348 and 350.

Mercury, ferrocyanides of, v vol II p 335

Mercury, fluorides of Two are known, HgF and HgF_2 , the VD of neither has been determined. These fluorides are remarkably unstable both as regards the action of water and of heat.

MERCURIOS FLUORIDE HgF or Hg_2F_2 . Prepared by adding freshly $\text{ppd Hg}_2\text{CO}_3$ to HFAq so long as the salt is dissolved, and evaporating, also by adding recently prepared HgCl_2 to a solution of Ag_2CO_3 in HFAq , filtering from AgCl , and evaporating on a steam bath (Finkner, *A* 110, 142). Berzelius obtained HgF mixed with HgCl_2 by subliming a mixture of HgCl_2 with NaF . HgF is a light yellow, crystalline, powder. It is partly

dissolved, and partly decomposed to HF and Hg_2O , by water. Exposed to light when moist it is blackened. HgF may be heated to $c\ 200^\circ$ without change, above this temperature Hg sublimes and the glass vessel is corroded. HgF is decomposed by alkalis with separation of Hg_2O , the action of NH_4Aq is more complex. HgF absorbs NH_3 gas forming $\text{N}_2\text{H}_5\text{HgF}_2$ (*v. Mercurammonium compounds*, p 206). Combines with SiF_4 to form *Mercurus silicofluoride* $\text{Hg}_2\text{SiF}_6 \cdot 2\text{H}_2\text{O}$ (*v. Mercury, silicofluorides of*, p 224).

MERCURIC FLUORIDE HgF_2 . According to Fremy (*A Ch* [3] 47, 5), crystals of this compound are obtained by dissolving HgO in excess of HFAq , and evaporating the solution slowly over lime. Finkener (*P* 110, 628) obtained an oxyfluoride, $\text{HgF}_2 \cdot \text{HgO} \cdot \text{H}_2\text{O}$, by this method, by adding conc. HFAq (50 p c) to this oxyfluoride, it was suddenly changed to a white crystalline mass of $\text{HgF}_2 \cdot 2\text{H}_2\text{O}$ (Finkener, *Lc*). The hydrated fluoride is decomposed at 30° , giving H_2O , HF, and HgO . $\text{HgF}_2 \cdot \text{H}_2\text{O}$ (F) HgF_2 is decomposed by water with separation of HgO . Addition of slight excess of NH_4Aq to HgF_2 in HFAq ppts $\text{NH}_4\text{HgOF} \cdot \text{HF}$ (Finkener, *P* 110, 632). *Mercuric silicofluoride* $\text{HgSiF}_6 \cdot 6\text{H}_2\text{O}$ probably exists (*v. Mercury, silicofluorides of*, p 224). Berzelius obtained a compound of HgF_2 with NH_4F by treating HgF_2 with NH_4Aq . HgF_2 combines with HgS (*v. Mercuric sulphide, Combinations*, No 6, p 225).

Mercury, fluosulphide of, *v. Mercuric sulphide, Combinations*, No 6, p 225.

Mercury, fulminate of, *v. vol II* p 317.

Mercury, haloid compounds of. Two series of these compounds exist, HgX and HgX_2 . The molecular weights of HgCl , HgBr , and HgI , are known, the molecular weights of HgCl , HgBr , and HgI are probably as represented by these formulæ. The compound HgI HgI also exists, and a peroxide of Hg has been isolated. The mercurous compounds HgX are insoluble in water, except HgCl , which is partly dissolved, and partly decomposed, by water, the mercuric compounds HgX_2 are soluble in water, with the exception of HgF_2 , which is decomposed by water. The Hg haloid compounds combine with many other haloid compounds, and also with oxyalts, to form numerous double salts. The double salts of the type $\text{HgX}_2 \cdot \text{M}'\text{X}$ and $\text{HgX}_2 \cdot 2\text{M}'\text{X}$ are best regarded as salts of the mercur halogen acids HIIgX_2 and H HgX , respectively, the acids HIIgX_2 have been isolated, and probably also the acids H HgX . The heats of formation of HgX and HgX_2 decrease as the atomic weight of X increases. Thomsen gives the following data (*Z P C* 2, 21) —

X	$[\text{Hg}, \text{X}]$	X	$[\text{Hg}, \text{X}]$
Cl	32,605	Cl	54,490
Br	25,475	Br	41,880
I	15,550	I	25,640

An iodochloride HgICl , and an iodobromide HgIBr , have been isolated.

Mercury, hydrated oxide of. According to Carnelley & Walker (*C J* 53, 80), the hydrate $\text{HgO} \cdot \text{H}_2\text{O}$ is obtained by ppg $\text{HgCl}_2 \cdot \text{Aq}$ by NaOH Aq , and drying in air, it is stable to about 100° , and is completely dehydrated at $c\ 175^\circ$. According to Schaffner (*A* 51, 181) the yellow pp formed by adding KOH Aq to solution of a

mercuric salt is $\text{HgO} \cdot 3\text{H}_2\text{O}$ (*v. Mercuric oxide*, p 222). But according to Millon (*B J* 27, 112), Marchand (*J pr* 37, 277), Rammelsberg (*J pr* [2] 38, 559), and Wallace (*Chem Gazette*, 1858 345), the pp obtained from Hg salts by KOH Aq is HgO .

Mercury, iodides of. Besides the iodides HgI and HgI_2 , corresponding with the chlorides, bromides, and fluorides, of Hg , there is said to exist a definite mercurio mercurio iodide, HgI HgI_2 , and a hexa iodide HgI_6 . Hg and I combine directly with production of heat.

Mercurous iodide HgI or Hg_2I_2 . Mol w not known with certainty. Melts at $c\ 290^\circ$ (with partial decomposition according to Stroman, *B* 20, 2818). Boils at $c\ 310^\circ$ with partial decomposition (Yvon, *C R* 76, 1607). *SG* 7.75 (Boullay, *A Ch* [2] 43, 266). *SH* $17^\circ-99^\circ$ 03949 (Regnault, *A Ch* [3] 1, 129). Crystallises in tetragonal forms, $a = 1.6726$ (Des Cloizeaux, *C R* 84, 1418). *H F* $[\text{Hg}, \text{I}] = 15,550$ (Thomsen, *Z P C* 2, 21).

Formation —1 By rubbing together Hg and I , in the ratio $\text{Hg} : \text{I}$, moistened with a few drops of alcohol, and removing the small quantity of HgI , which is always formed by treatment with alcohol, in which HgI is soluble but HgI_2 is insoluble —2 By rubbing together HgI and Hg in the ratio $\text{HgI} : \text{Hg}$, and dissolving out unchanged HgI by alcohol —3 By heating 10 parts I with $1\frac{1}{2}$ parts Hg in a retort, on a sand bath, to not above 200° , HgI sublimes in red crystals, becoming yellow when cold (Yvon, *C R* 76, 1607). 4 By adding KIAq to solution of a mercuric salt, preferably the acetate (*v. Lefort, Ph* [3] 3, 823).

Preparation —A conc solution of HgNO_3 , containing a little HNO_3 , and free from basic nitrate, is heated to boiling with excess of I , when the I is partially dissolved the liquid is carefully decanted into a warm basin, on cooling, very lustrous transparent, yellow tablets of HgI separate. The crystals are brought on to a filter (best in the dark), washed with cold water containing a little HNO_3 , and then with pure cold water, and dried in the dark at ordinary summer temperature by placing them on filter paper which is frequently changed (Stroman, *B* 20, 2819). HgI is obtained as a flocculent pp by rapidly adding a fairly conc solution of I in alcohol to cold conc $\text{HgNO}_3 \cdot \text{Aq}$ (*S, lc*).

Properties and Reactions — HgI crystallises in yellow tetragonal tablets, isomorphous with HgCl . HgI is sometimes described as a green powder. Stroman (*lc*) says that green preparations are impure. When heated, the yellow crystals become dark yellow, then orange, and finally garnet red. Yvon (*C R* 76, 1607) says that the change of colour begins at 70° , and assigns a definite temperature for each change of colour. Stroman (*B* 20, 2818) could not connect the different colours with definite intervals of temperature, he says that the salt prepared by him was pure yellow at 100° . Sublimation begins at $c\ 190^\circ$ according to Yvon, at $c\ 110^\circ-120^\circ$ according to Stroman. HgI when moist is rapidly blackened by light. HgI is very slightly soluble in water, insoluble in alcohol. It is easily decomposed to Hg and HgI_2 , eg by HIAq , KIAq , and similar iodides. Heated rapidly HgI gives a sublimate of Hg and HgI HgI , HgI

is used in medicine. For medicinal purposes it must be quite free from HgI_2 , which is a violent poison.

MERCURIO MERCURIO IODIDE HgI HgI_2 . A compound of Hg and I of this composition was obtained by Boullay (*A Ch* [2] 34, 345) by adding KIAq containing I, in the ratio $\text{KI} \frac{1}{2} \text{I}$, to HgNO_3Aq . The same compound is said to be obtained by adding KIAq to HgNO_3Aq and digesting the pp for some time in the liquid, also by rubbing together HgI_2 and I in the ratio $3\text{HgI}_2, \text{Hg}$. Mercurio mercurio iodide is described as a yellow powder, insol water and alcohol, unchanged in the dark, but becoming dark when exposed to light. When heated, it turns red, then melts, and may be sublimed unchanged, yielding crystals which are red when hot but become yellow on cooling. HIAq , and various iodides in solution, dissolve HgI_2 , leaving HgI , and then decompose the HgI with separation of Hg.

MERCURIO IODIDE HgI_2 . Mol w 452.86 [238°] (Carnelley & Williams, *C J* 33, 283) (339°–359°) (*C a W, l c*). SG red variety 6.2941 to 6.3004 at 0°, 6.276 at 126°, yellow variety 6.225 at 126° (Rodwell & Elder, *Pr* 28, 284), 6.179 at 200° solid, 5.286 at 200° molten (*R a E, l c*). VD 225.7 to 234.4 (Mitscherlich, *P* 29, 193). SH 18°–99° 0.4197 (Regnault, *A Ch* [3] 1, 129). Crystallises in tetragonal forms (red), $a, c = 1.19955$, also in rhombic forms (yellow), $p, p = 114^\circ 30'$. HF [HgI_2] = 25,640 (Thomsen, *Z P C* 2, 21). Change of yellow to red crystals is accompanied by production of heat, 3000 gram units (Berthelot, *Bl* [2] 39, 17, cf Weber, *P* 100, 127). S 0.04 at 17.5°, 0.05 at 22°, S 286.50 p.c. alcohol at 18°, 1.186 absolute alcohol at 18° (Bourgoin, *Bl* [2] 42, 620).

Formation—1 By the action of I on Hg, Dublanc (*Ph C* 1849 656) says that pure HgI_2 is obtained by pouring 1,000 pts alcohol (93 p.c.) on to 100 pts Hg, and adding 124 pts I in portions of 10 pts at a time, after each addition of I, the whole is agitated until the alcohol becomes colourless, the addition of the last portion of I should leave the alcohol coloured, the HgI_2 is washed with alcohol—2 By adding KIAq or FeI_2Aq to HgCl_2Aq .

Preparation—1 8 pts HgCl_2 and 10 pts KI are dissolved, separately, in water, the solutions are mixed, and the pp is thoroughly washed with cold water. Any excess of KI dissolves some HgI_2 , and any excess of HgCl_2 produces a yellowish pp containing more or less HgCl_2 . Williams (*Ph* [3] 3, 1015) dissolves 8 pts HgCl_2 in 4 pts NH_4Cl in water, and then adds 10 pts KI in water—2 10 pts I are suspended in water, and Fe filings are added until all the I is changed to FeI_2 , the solution is at once added to a solution of 10½ pts HgCl_2 in water, the pp of HgI_2 is rapidly separated from the liquid (to prevent formation of Fe oxychlorides) and washed with cold water. By crystallising from hot KIAq , or, better, from hot conc HClAq (Köhler, *B* 12, 608), HgI_2 is obtained in very lustrous, red tetragonal prisms, with a greenish reflection.

Properties— HgI_2 obtained by ppn is a pure scarlet coloured, crystalline, heavy powder. It is sol. water, S = 66 (Wurtz). Sol hot alcohol; somewhat sol glycerin (in 340 pts, Fairley,

M. S [3] 9, 685); also in ether, and some oils. Sol KIAq , also in conc hot HClAq , from these solutions HgI_2 separates in well formed crystals. HgI_2 is dimorphous. Red (tetragonal) HgI_2 is changed to the yellow (rhombic) modification by heat, the change occurs at 126° (Rodwell & Elder, *Pr* 28, 284), it is accompanied by a sudden, and then by a regular, expansion. SG, red at 126° = 6.276, yellow at 126° = 6.225 (*R a E*). HgI_2 melts at 238° (Carnelley & Williams, *C J* 33, 283), at 253°–254° according to Köhler (*B* 12, 608), to a blood red liquid, melting is attended with considerable expansion. At c 339°–359° the liquid boils, and sublimes to yellow rhombic plates. The yellow variety passes into the red at the ordinary temperature, this change is hastened by rubbing or scratching the red crystals, heat is produced in the process, 3,000 gram units according to Berthelot (cf Weber, *P* 100, 127). The HgI_2 obtained by adding cold KIAq to cold HgCl_2Aq appears at first yellow, but it quickly changes to red, the change of crystalline form in this case may be observed under the microscope. Solutions of HgI_2 in alcohol, ether, or KIAq are colourless, if the HgI_2 is allowed to form slowly from these solutions, the crystals are red, but if the ppn is made rapid, e.g. by pouring an alcoholic solution into water, the crystals appear yellow for a moment, but they quickly change to red (cf Schiff, *A* 111, 371, Selmi, *J* 1855 417).

Reactions—1 HgI_2 dissolves in *hydriodic acid* solution with production of heat (v Berthelot, *Bl* 38, 369). This solution has about the same heat of neutralisation as HIAq (Thomson), with alkalis it forms salts HgI_2MI , the solution very probably contains the acid H_2HgI_2 (cf *Combinations*, No 2). An *iodomercuric acid*, HHgI_2 , has been obtained by slightly warming excess of HgI_2 with HIAq , filtering through asbestos, and cooling to between 0° and –4° (Neumann, *M* 10, 236), this acid is very readily decomposed by moisture and by heat. According to Boullay (*A Ch* [2] 34, 345) a solution of HgI_2 in hot HIAq deposits I on cooling, and then crystals of $2\text{HgI}_2, 3\text{HI}$ —2 Most heavy metals, e.g. Zn, Sn, when triturated with HgI_2 , remove I and leave Hg, *potassium* acts in the same way—3 HgI_2Aq is decomposed by alkalis, with separation of HgO and formation in solution of a double salt $x\text{HgI}_2, y\text{MI}$. *Baryta* and *strontia* react similarly to alkalis—4 *Lime*, *sodium carbonate*, and *potassium carbonate* do not decompose HgI_2Aq , but a solution of HgI_2 in alcohol is decomposed by these reagents—5 HgI_2 dissolves in hot *calcium hypochlorite* solution, on cooling Ca periodate is deposited and HgCl_2 remains in solution (Rammelsberg). 6 When HgI_2 is boiled with *nitric acid*, S HgI_2 1.5–1.4, $\text{Hg}(\text{IO}_3)_2$ is formed, with acid SG 1.3, white leaflets of $\text{HgI}_2, \text{Hg}(\text{NO}_3)_2$ separate, with acid SG 1.2, crystals of unchanged HgI_2 are obtained, together with some of the compound $\text{HgI}_2, \text{Hg}(\text{NO}_3)_2$ (Kraut, *Z* 18, 3461)—7 *Chlorine*, passed into water containing HgI_2 in suspension, produces a yellow solution containing HgCl_2 and ICl_2 (Fehlbol).

Combinations—1 With *hydriodic acid* (v *Reactions*, No 1)—2 With various metallic iodides to form double salts, regarded by Von Bonsdorff as *iodomercurates* (*P* 17, 265).

These salts have been examined chiefly by Boullay (*A Ch* [2] 34, 345). They are generally obtained by dissolving HgI_2 in a solution of the other iodide and evaporating. The following are the most important -

(i) With *alkali iodides* (a) With *potassium iodide* $2HgI_2 \cdot 2KI \cdot 3H_2O$, $HgI_2 \cdot 2KI$. The former is produced by saturating boiling $KIAq$ with HgI_2 , filtering, separating from HgI_2 , which crystallises out, and evaporating. Sol. alcohol and ether. Decomposed by water, with formation of HgI_2 , and $HgI_2 \cdot 2KI$ which separates on evaporation. These soluble salts may be prepared by boiling $KIAq$ with HgO , the solution contains the double salts and also KOH (*cf Jahn, Ar Ph* [3] 1, 97). (b) With *sodium iodide* $NaIAq$ reacts with HgI_2 similarly to $KIAq$, the compositions of the double salts of NaI with HgI_2 have not been satisfactorily determined. (c) With *ammonium iodide* $2(NH_4I \cdot HgI_2) \cdot 3H_2O$, obtained, as yellow needles, by dissolving HgI_2 in hot NH_4IAq , separating from HgI_2 , which forms on cooling, and evaporating.

(ii) With *other metallic iodides* BaI_2 , SrI_2 , CaI_2 , and MgI_2 solutions react with HgI_2 similarly to $KIAq$, the double salts are probably $M_2 \cdot HgI_2$. HgI_2 appears to form double compounds with CdI_2 and Kel . The compound $2HgI_2 \cdot CuI_2$ was obtained by Hess (*D P J* 218, 183) - 3 HgI_2 combines with *mercuric chloride*, also with *mercuric bromide* (*v Mercury, iodochlorides and iodobromides of, infra*) - 4 Combines with *mercuric sulphide* (*v Mercuric sulphide, Combinations*, No 5, p 225).

MERCURY, HEXA IODIDE OF HgI_2 . This periodide is said by Jørgensen (*J pr* [2] 2, 357) to be obtained by adding cold $HgCl_2$ Aq , followed by addition of water, to an alcoholic solution of KI , heated to 50° . If the solutions are mixed hot, large rhombic crystals of HgI_2 are formed, but they are always mixed with HgI_2 . HgI_2 is decomposed quickly by alcohol, slowly by water, with formation of red HgI_2 . HgI_2 possesses most of the optical properties of *tourmalin*.

Mercury, iodobromide of $HgIBr$. Sulphur yellow, translucent, rhombic prisms, $a b c = 0.443 \ 1 \ 0.9194$ (Groth, *B* 2, 574). Melts at $c \ 229^\circ$, and boils at $c \ 360^\circ$, crystallisable from ether, may be sublimed unchanged. Produced by crystallising a mixture of HgI_2 and $HgBr_2$ from ether or acetone, also by the reaction of an alkyl iodide, *e g* EtI , on $HgBr_2$, dissolved in acetone (Oppenheim, *B* 2, 571).

Mercury, iodochlorides of Two iodochlorides of Hg have been isolated. (i) $HgI_2 \cdot 2HgCl_2$ ($=Hg_3I_2Cl_4$). Obtained by dissolving HgI_2 in hot $HgCl_2$, and cooling (Liebig, *S* 49, 252), also by boiling $HgCl_2$ with excess of I and much water until I vapour is no longer given off (Selmi, *J* 1855 417). (ii) $HgICl$. Prepared by heating, in a closed tube at 140° - 160° , HgI_2 , $HgCl_2$, and a little water, until no HgI_2 remains unchanged (Köhler, *B* 12, 1187). Boullay (*P* 48, 175) said that $HgICl$ is produced by saturating hot $HgCl_2$ Aq with HgI_2 , but Köhler found that very little was formed in this way. $HgICl$ forms a citron-yellow crystalline mass, which becomes red after about 12 hours. The red modification forms tetragonal, the yellow forms rhombic, crystals. $HgICl$ is citron-yellow at

$c \ 125^\circ$, it melts at $c \ 158^\circ$ to a golden coloured liquid which solidifies at $c \ 146^\circ$, it is slightly soluble, with partial decomposition, in hot water, more soluble in alcohol. $HgICl$ may be sublimed, with only slight decomposition, in a stream of HCl , SO_2 , or H_2S . From a solution of $HgICl$ in dilute $HClAq$, H_2S ppt's a yellow solid, probably $HgICl \cdot HgS$ (Köhler, *lc*).

Mercury, iodosulphides of, *v Mercuric sulphide, Combinations*, No 5, p 225

Mercury, nitride of Hg_3N_2 (*Trimercuramine*). This compound was first isolated by Plantamour (*A* 40, 115). $Ppd \ HgO$ is dried at 40° - 50° , and then heated at 100° in a stream of dry NH_3 , so long as water is evolved (Hirzel, *J* 1852 419). Plantamour directed to pass NH_3 over cold HgO , then to heat to 150° in NH_3 , and finally to remove unchanged HgO by HNO_3 Aq , Hirzel found that HNO_3 Aq reacts with Hg_3N_2 . Mercury nitride is a brown powder, very explosive, but requires a higher temperature, or a stronger blow, than nitrogen iodide to explode it. Hg_3N_2 is not acted on by cold dilute H_2SO_4 , the hot acid decomposes it. Conc H_2SO_4 reacts energetically and explosively. Conc HNO_3 Aq forms $Hg(NO_3)_2$ and NH_4NO_3 , dilute HNO_3 Aq forms a white powder, $HClAq$ forms $HgCl_2$ and NH_4Cl . Mixed with KOH and heated, NH_3 and Hg are formed. Moist Hg_3N_2 is slowly decomposed by light, in water it is changed, after 24 hours, to a white powder.

Mercury, oxides of. Two oxides of Hg are known, Hg_2O and HgO , it is doubtful whether Hg_2O has been obtained free from Hg and HgO . Neither oxide has been gasified, and the mol w of neither is known with certainty.

MERCUROUS OXIDE Hg_2O (*Suboxide, or black oxide, of mercury*). This oxide is prepared by adding $NaOHAq$ or $KOHAq$ to solution of a mercurous salt, or to $HgCl$ suspended in water. The best salt to use seems to be $HgNO_2$, it should be dissolved in water with a very little HNO_3 , the solution being made dilute, ppn with $KOHAq$, and washing with cold water, should be performed in the dark, and the black pp should be dried in the dark without heating. Guibourt (*A Ch* [2] 1, 422) says that Hg_2O cannot be obtained quite free from Hg and HgO , this is confirmed by Bruns a O v d Flordten (*B* 21, 2010), who assert that Hg_2O oxidises to HgO at the ordinary temperature. Barfoed (*J pr* [2] 38, 441) says that the product of the action of $NaOHAq$ on mercurous salts is a mixture of Hg_2O , HgO , and Hg in varying proportions. The descriptions of Hg_2O probably apply to Hg_2O mixed with more or less Hg and some HgO .

Mercurous oxide is a black powder, *S G* 10 69 (Herapath, *P M* 64, 321), 8 95 (Karsten, *S* 65, 894). $H.F [Hg_2O] = 24,860$ (Thomsen, *Z P C* 2, 21). Hg_2O is decomposed very easily by light or heat, giving HgO and Hg , strongly heated gives Hg and O . Dilute $HClAq$ produces $HgCl$. Hg_2O is soluble in conc acetic acid. With dilute acids Hg_2O generally yields mercurous salts, with H_3PO_4 Aq it gives H_3PO_4 and Hg . Boiled with $KIAq$, Hg and $HgI_2 \cdot 2KIAq$ are formed (Berthelot, *J Ph* 14, 189). With conc. NH_4OHAq , NH_3 is evolved and Hg and $HgCl_2$ are produced (Pagenstecher, *R. P.* 27, 27, Thompson, *P. M* [3] 10, 179).

MERCURIC OXIDE HgO . (*Red oxide of mercury* *Red precipitate*) Mol w unknown, as compound has not been gasified SG 11 074 at 17.5° (Herapath, *P M* 64, 321), 11 136 to 11 344 at 4° (Playfair a Joule, *C S Mem* 3, 84, and *C J* 1, 137), 11 29 at 4° *in vacuo* (Le Royer a Dumas, in Bottger's *Tabellarische Uebersicht der Spec Gewichte der Körper*, Frankfurt, 1837) $S H$ $19^\circ-52^\circ = 053$ (Kopp, *T* 155, 71), $5^\circ-98^\circ = 0518$ (Regnault, *A Ch* [3] 1, 129) $H F$ (Hg_2O) = 22,000 (Thomsen, *Z P C* 2, 21) Crystallises in rhombic forms $a b c = 6523$ 1 9456 (Nordenskjöld, *P* 114, 621), in monoclinic forms, according to Des Cloiseaux (*A Ch* [4] 20, 201) $C E$ ($0^\circ-100^\circ$) 0058 (Playfair a Joule, *C J* 1, 137)

HgO was known to the Arabians in the eighth century, Geber prepared it by calcining the nitrate, and towards the end of the seventeenth century Boyle obtained it by strongly heating Hg in air

Formation—By heating Hg to near its B P in a loosely covered vessel for a long time

Preparation—1 Hg is dissolved in HNO_3Aq , the solution is evaporated to dryness, the residue is powdered and then heated on a sand tray so long as N oxides are evolved, the temperature being slowly raised, and the heating continued till a porcelain plate held over the vessel shows a slight deposit of Hg HgO thus prepared forms red lustrous scales—2 One of the oxychlorides 2HgO HgCl_2 , and the oxychloride 4HgO HgCl_2 , when treated with KOH Aq , give red HgO , another oxychloride 2HgO HgCl_2 , and also the oxychloride 3HgO HgCl_2 , give yellow HgO by treatment with KOH Aq The compound 2HgO HgCl_2 which yields red HgO is prepared by mixing 1 vol KHCO_3Aq , free from K_2CO_3 , saturated at 15° , with 3 vols HgCl_2Aq , also saturated at 15° , stirring with a glass rod till black streaks appear where the rod rubs the glass, pouring off, adding a fresh quantity of the mixed solution, and again stirring The compound 2HgO HgCl_2 which gives the yellow HgO is prepared by stirring a mixture of 1 vol KHCO_3Aq , saturated at 15° , with 6 to 10 vols HgCl_2Aq , also saturated at 15° The compound 4HgO HgCl_2 is prepared by mixing 1 vol HgCl_2Aq (saturated at 15°) with a large excess (4 to 6 vols) KHCO_3Aq (saturated at 15°), and allowing to stand for some time (*cf Mercury, oxychlorides of*, p. 223)—3 $\text{Hg}(\text{NO}_3)_2\text{Aq}$, or HgCl_2Aq , is poured into KOH Aq , the pp is thoroughly washed and dried at $100^\circ-120^\circ$ HgO thus prepared forms a yellow compact solid

Properties.— HgO exhibits allotropy, it forms heavy, red, crystalline scales, or a somewhat more bulky, amorphous, yellow powder (*vsupra*) HgO is a violent irritant poison HgO is very slightly soluble in water, 1 pt dissolves in 200,000 cold water, or in 125,000 pts if the water is boiled with HgO and then allowed to cool (Wallace, *Chem Gazette*, 1858, 845) The solution has a metallic taste Heated to redness, Hg is separated into Hg and O Heated below the temperature of decomposition, HgO turns black, but recovers its original colour on cooling. HgO is slowly blackened and decomposed to Hg and O by the action of light There are differences in the reactions of Cl , HgCl_2 , in

alcoholic solution, and oxalic acid, with the two varieties of HgO (*v Reactions*, Nos 8, 9, 10)

Reactions—1 Decomposed by heating to redness, giving Hg and O , Pelouze (*C R* 16, 50) said that yellow HgO is decomposed at a lower temperature than the red variety, but this was not confirmed by Gay Lussac (*C R* 16, 309), nor by Millon (*B J* 27, 112) Carnelleys Walker (*C J* 53, 80) found that HgO , ppd by NaOH Aq from HgCl_2Aq , began to give off O at a little above 175° , and that evolution of O became rapid at $c 415^\circ$, at which temperature red HgO was formed and decomposed Myers's results (obtained by measuring vapour pressures of O from HgO , *B* 6, 11) agree with those of *C* a W Debray (*C R* 77, 123) found that when a tube containing HgO was sealed, and the whole tube was heated to 440° , the O evolved was almost wholly again absorbed—2 A mixture of HgO with such easily oxidised bodies as *P*, *S*, *Sb*, detonates when heated more or less violently Some organic compounds are oxidised by boiling in solution with HgO SO_2Aq boiled with HgO produces SO_3Aq and Hg SnCl_2Aq forms SnCl_2Aq and Hg —3 HgO is decomposed, rapidly and somewhat violently, by heating with *magnesium*, in the ratio HgO Mg , with formation of MgO and Hg (Winkler, *B* 23, 128)—4 Heated with *sodium*, *Na* amalgam and the compound $\text{Na}_2\text{O HgO}$ are formed (Bickeloff, *B* 13, 2392), the compound $\text{Na}_2\text{O HgO}$ is not decomposed by heat, but very quickly by water—5 HgO dissolves in *molten potash* If the KOH is nearly saturated with HgO , allowed to cool slowly, and lixiviated with a little cold water, a heavy violet crystalline powder and a lighter greyish green powder are obtained The violet powder is said to consist of $\text{K}_2\text{O HgO}$, and the greyish powder to contain from 2 to 5 p.c. K_2O , the SG of $\text{K}_2\text{O HgO}$ is 10.31, it is decomposed at high temperatures, also by continued washing with water, but less completely by alcohol (St Meunier, *C R* 60, 557)—6 According to Foubert (*A Ch* [4] 1, 300), saturated solutions of the *alkali and alkaline earth haloid compounds* are decomposed, with separation of alkali or alkaline oxides, by boiling with HgO (*cf* Melsens, *A Ch* [3] 26, 220, and *H Rose*, *P* 107, 298) André, however, says that boiling saturated solutions of alkaline earth chlorides react with HgO to form either oxychlorides $x\text{HgCl}_2$, $y\text{HgO}$, or compounds of HgO with the alkaline chloride (*C R* 104, 431, *v Mercury oxychlorides of*, p. 223, and also *infra*, *Combinations*, No 1) According to Jehn (*Ar Ph* [3] 1, 97), HgO boiled with KI Aq produces KOH Aq and the double iodide HgI_2KI —7 HgO reacts with most acids to form mercuric salts HgX_2 ($\text{X} = \text{NO}_3$, $\frac{1}{2}\text{SO}_4$, $\frac{1}{2}\text{PO}_4$, &c)—8 The reaction of *chlorine* with HgO differs according to the conditions, and the variety of HgO used Cl scarcely reacts with red crystalline HgO , with yellow HgO , prepared by ppn and dried at $c 100^\circ$, Cl reacts energetically forming HgCl_2 and O , with the yellow oxide, dried at 300° and cooled, Cl reacts more slowly, forming Cl_2O and HgCl_2 , or, if water be present, forming HgClO Aq and HgO HgCl_2 (*cf vol* i pp 12, 16)—9 An alcoholic solution of *mercurous chloride* heated with yellow HgO at once produces black oxychloride HgO HgCl_2 , the same oxychloride is formed from the red variety of

HgO only after prolonged action of boiling alcoholic HgCl_2 solution—10 *Oxalic acid solution* reacts with yellow HgO, to form oxalate, in the cold, but the red variety is unchanged when boiled with $\text{H}_2\text{C}_2\text{O}_4\text{Aq}$

Combinations — 1 With *alkaline earth chlorides*, obtained by the reaction of HgO with solutions of CaCl_2 , BaCl_2 , and SrCl_2 . The compounds $2\text{HgO} \cdot \text{CaCl}_2 \cdot 4\text{H}_2\text{O}$, $\text{HgO} \cdot \text{BaCl}_2 \cdot 6\text{H}_2\text{O}$, and $\text{HgO} \cdot \text{SrCl}_2 \cdot 6\text{H}_2\text{O}$, are described by André (*C R* 104, 431)—2 With *ammonia*, to form $2\text{HgO} \cdot \text{NH}_3 \cdot \text{H}_2\text{O} = \text{NH}_4\text{OH} \cdot 2\text{HgO}$, this compound reacts with acids as a base forming *ammonium salts* (v *MERCURAM MONIUM COMPOUNDS*, p 206)

Mercury, oxybromide of $8\text{HgO} \cdot \text{HgBr}_2 = \text{Hg}_9\text{O}_8\text{Br}_2$ (*Basic mercuric bromide*) A yellow, crystalline, powder, obtained by boiling HgBr_2Aq with HgO, filtering, and allowing to crystallise, or by partial ppn of HgBr_2Aq by KOHAq, and boiling the liquid in contact with the pp (Lowig, *P* 14, 485, cf Rammelsberg, *P* 55, 248)

For H F of oxybromides of Hg v André, *Bl* [2] 41, 274

Mercury, oxychlorides of $\text{Hg}_2\text{O}_2\text{Cl}_2$ (*Basic mercury chlorides*) Several oxychlorides of Hg are known, they are produced by the reaction of HgO with HgCl_2Aq , by partial ppn of HgCl_2Aq by KOHAq or NaOHAq, by mixing KHCO_3Aq and HgCl_2Aq in different proportions, and by the reaction of Cl with yellow HgO

Millon (*A Ch* [3] 27, 253) described oxychlorides obtained by mixing solutions of KHCO_3 (free from K_2CO_3) and HgCl_2 , both saturated at 15° (i) $2\text{HgO} \cdot \text{HgCl}_2$, by adding 1 vol KHCO_3Aq to 6–10 vols HgCl_2Aq , also by adding 1 vol KHCO_3Aq to 3 vols HgCl_2Aq , stirring till ppn begins on the sides of the vessel, pouring off the liquid, adding a fresh quantity of the same mixture, and stirring again. The compound obtained by the first process forms a red non crystalline powder, and yields *yellow* HgO when acted on by KOHAq, as obtained by the second process, the compound is a heavy, black, lustrous solid, which yields *red* HgO with KOHAq (ii) $3\text{HgO} \cdot \text{HgCl}_2$, by mixing equal volumes of the solutions of KHCO_3 and HgCl_2 , and allowing to stand, a golden yellow crystalline pp, yielding *yellow* HgO with KOHAq (iii) $4\text{HgO} \cdot \text{HgCl}_2$, by mixing 1 vol HgCl_2Aq with 4–6 vols KHCO_3Aq , and allowing to stand, a brown crystalline solid, which yields *red* HgO with KOHAq

Roucher (*A Ch* [3] 18, 372) described seven oxychlorides, containing HgCl_2 and HgO in the ratios 1, 2, 3, 4, 5, and 6, each oxychloride exists, according to Roucher, in two varieties, one derived from and giving *red* HgO, the other derived from and giving *yellow* HgO, some of the oxychlorides also exhibit other differences. Roucher described 15 different oxychlorides. A tabular statement showing the chief differences between the oxychlorides is given by Roucher

André (*C R* 104, 431) described two oxychlorides, $\text{HgCl}_2 \cdot 3\text{HgO}$ and $2\text{HgCl}_2 \cdot 8\text{HgO}$, obtained by dissolving HgO in boiling CaCl_2Aq and MgCl_2Aq , respectively, and pouring the solutions into large quantities of cold water.

The oxychlorides of Hg have been examined more recently by Thümmel (*Ar. Ph.* 37, 589, v.

abstract in *C J* 56, 1050) f. says that only 5 definite compounds exist, viz, $\text{HgO} \cdot 2\text{HgCl}_2$, $\text{HgO} \cdot \text{HgCl}_2$, $2\text{HgO} \cdot \text{HgCl}_2$, $3\text{HgO} \cdot \text{HgCl}_2$, and $4\text{HgO} \cdot \text{HgCl}_2$. (i) $\text{HgO} \cdot 2\text{HgCl}_2$ is formed in all solutions prepared by dissolving HgO in HgCl_2Aq , but it is readily decomposed, it is best prepared by heating at 100° 1 pt HgO (red or yellow), with 10 pts HgCl_2 , and 60 pts water, stirring till no further separation of a yellow powder occurs, washing, drying, and then washing with ether (free from alcohol) to remove HgCl_2 . Warm water separates $2\text{HgO} \cdot \text{HgCl}_2$, NaOHAq ppts *red* HgO. (ii) $\text{HgO} \cdot \text{HgCl}_2$ has not yet been obtained pure. (iii) $2\text{HgO} \cdot \text{HgCl}_2$ exists in two varieties, (a) *red* variety, obtained by mixing solutions of NaHCO_3 and HgCl_2 in the ratio 2NaHCO_3 , HgCl_2 , (b) *black* variety, obtained by heating the red variety when dry, or by adding HgO to hot HgCl_2Aq , or by mixing equivalent quantities of *red* HgO and HgCl_2 in cold water, or by treating $3\text{HgO} \cdot \text{HgCl}_2$ with cold HgCl_2Aq . (iv) $3\text{HgO} \cdot \text{HgCl}_2$ is a yellow pp obtained by adding alkaline carbonate (normal or acid) to HgCl_2Aq , avoiding excess of the carbonate, also by treating freshly ppd *yellow* HgO with HgCl_2Aq (this preparation is yellow and yields *yellow* HgO with KOHAq), also by treating washed *red* HgO with HgCl_2Aq (this preparation is reddish yellow and yields *red* HgO by KOHAq). (v) $4\text{HgO} \cdot \text{HgCl}_2$ is obtained as a brown amorphous powder, by adding KHCO_3Aq to HgCl_2Aq , in the ratio 30 to 35 KHCO_3 , HgCl_2 , it is also obtained, as reddish-brown crystalline plates, by shaking HgCl_2Aq with *red* HgO in the ratio HgCl_2 , 6HgO , both varieties give *yellow* HgO by KOHAq. All these oxychlorides yield sublimates containing HgCl and HgCl when heated

For H F of oxychlorides of Hg v André, *Bl* [2] 41, 274

Mercury, oxycyanides of, v. CYANIDES, vol. ii p 342

Mercury, oxyfluoride of $\text{HgO} \cdot \text{HgF}_2 \cdot \text{H}_2\text{O}$ (*Basic mercuric fluoride*) A yellow crystalline solid, obtained by dissolving freshly ppd HgO in HFAq , and digesting the solution with HgO, or evaporating the solution (Finkener, *P* 110, 628). The oxyfluoride is also obtained by heating $\text{HgF}_2 \cdot 2\text{H}_2\text{O}$ (*q v* p 210) to 30° (*F*). Heated above 100° , the oxyfluoride gives off H_2O and HF, it is decomposed by water with separation of HgO

Mercury, oxyiodide of $3\text{HgO} \cdot \text{HgI}_2$ (*Basic mercuric iodide*) A yellowish brown solid, obtained by melting together HgO and HgI_2 in the ratio $3\text{HgO} \cdot \text{HgI}_2$, also by digesting HgI_2 with dilute KOHAq (Rammelsberg, *P* 48, 182)

Mercury, oxysulphides of Oxysulphides of Hg have been described, but according to Pollock (*B* 22, 2859) none has been isolated, and the existence of any is very improbable

Mercury, phosphide of No phosphide of Hg has been isolated with certainty. By passing P hydride, prepared by boiling amorphous P with KOHAq, into HgCl_2Aq , several compounds have been obtained. Aschan (*Chem. Zeitung*, 10, 82, 102) describes a yellow compound $3\text{Hg}_2\text{P}_2 \cdot 7\text{HgCl}_2$, a red compound $4\text{Hg}_2\text{P}_2 \cdot 5\text{HgCl}_2$, and a black compound $\text{Hg}_2\text{P}_2 \cdot \text{HgCl}_2$. A compound probably $2\text{PH}_3 \cdot 6\text{HgO}$ is said by Aschan (*lc*) to be formed by passing P hydride over Hg_2CO_3 . The ppn.

formed when P hydride is passed into $\text{Hg}(\text{NO}_3)_2\text{Aq}$ are probably mixtures of different compounds of Hg, P, and HNO_3 . (A) Most of the foregoing compounds are readily decomposed, some explosively, by warming.

Mercury, salts of Hg forms two series of salts, *mercurous salts* corresponding with Hg_2O , and *mercuric salts* corresponding with HgO . The former are obtained sometimes by dissolving Hg_2O in acids, sometimes by dissolving Hg in acids and digesting with excess of Hg, and in some cases by double decomposition from the nitrate, the mercuric salts are obtained by dissolving HgO or HgCO_3 in acids, or by double decomposition from the nitrate. The mercurous salts have the composition Hg_2X , where $\text{X} = \text{NO}_3$, $\frac{1}{2}\text{SO}_4$, &c., the mercuric salts have the composition HgX_2 . The mercuric salts are more stable, as a class, than the mercurous salts. The normal mercurous salts are generally decomposed by water into a basic salt, which separates, and an acid salt, which goes into solution. Many basic salts and a large number of double salts are known. The salts of Hg, as a class, act as violent and irritating poisons. The principal salts of Hg are the following (*v* CARBONATES, NITRATES, SULPHATES, &c.) *antimonates, arsenates and -ites, bromates, carbonates, chlorates and perchlorates, chromates, iodates and periodates, molybdates, nitrates and -ites, phosphates, selenates and -ites, sulphates and -ites, tantalates, thiosulphates, tungstates, vanadates*.

Mercury, selenide of HgSe Small quantities of more or less pure HgSe occur native in the Harz, accompanying PbSe . HgSe is produced by strongly heating together Hg and Se, and subliming at a higher temperature, forms grey, metal like, lustrous laminae, dissolved by *aqua regia* with formation of Hg selenite, slowly changed to the same compound by boiling conc nitric acid (Uelsmann, *A* 116, 126, cf Little, *A* 112, 211). HgSe is also formed by passing H_2Se for a long time into HgCl_2Aq , the white pp of $2\text{HgSe} \cdot \text{HgCl}_2$ at first produced is changed to black HgSe . S G of natural HgSe is 7.1 to 7.4, S G of artificially prepared HgSe is 8.77.

Combinations—1 With *mercuric chloride*, to form $2\text{HgSe} \cdot \text{HgCl}_2$, a white pp formed by adding H_2Se , or K_2Se , to excess of HgCl_2Aq .—2 With *mercuric oxide*, to form $2\text{HgSe} \cdot \text{HgO}$, a black powder formed by treating $2\text{HgSe} \cdot \text{HgCl}_2$ with NaOH Aq . Decomposed by heat, yielding sublimates of HgSe and Hg (Uelsmann, *A* 116, 126).

Mercury, selenochloride of $\text{Hg}_2\text{Se}_2\text{Cl}_4$, *v* *Mercury, selenide of, Combinations*, No 1.

Mercury, selenosulphide of A mineral approaching the composition $\text{HgSe} \cdot 4\text{HgS}$ is found near San Onofre in Mexico, it is known as *onofrite*.

Mercury, selenocyanides of HgSeCy and $\text{Hg}(\text{SeCy})_2$, *v* vol u p 348.

Mercury, silicofluorides of Two are known, $\text{Hg}_2\text{SiF}_6 \cdot 2\text{H}_2\text{O}$ and $\text{HgSiF}_6 \cdot 6\text{H}_2\text{O}$.

MERCUROUS SILICOFLOURIDE, $\text{Hg}_2\text{SiF}_6 \cdot 2\text{H}_2\text{O}$ Clear prismatic crystals, obtained by dissolving Hg_2CO_3 in $\text{H}_2\text{SiF}_6\text{Aq}$, evaporating, washing with a little water, and pressing between paper (Finkener, *P* 111, 246, cf Berzelius, *P* 1, 200).

MERCURIC SILICOFLOURIDE, $\text{HgSiF}_6 \cdot 6\text{H}_2\text{O}$. Obtained by dissolving HgO in $\text{H}_2\text{SiF}_6\text{Aq}$, evaporating till yellow needles ($\text{HgSiF}_6 \cdot \text{HgO} \cdot 3\text{H}_2\text{O}$) begin to separate, and then allowing to stand at a temperature not above 15° . Forms clear, colourless, rhombohedral crystals, very unstable, deliquescent in air, and efflorescent over H_2SO_4 , composition doubtful (Finkener, *P* 111, 246). The compound $\text{HgSiF}_6 \cdot \text{HgO} \cdot 3\text{H}_2\text{O}$ (Finkener) was described by Berzelius (*P* 1, 200) as the normal salt.

Mercury, sulphides of Two are known, Hg_2S and HgS , the latter reacts towards the alkali sulphides as an acidic sulphide, it also combines with many Hg compounds to form double compounds.

MERCUROUS SULPHIDE, Hg_2S This sulphide is best prepared, according to Berzelius, by dropping mercurous acetate solution into Na or NH_4 sulphide solution (cf Brande, *Q J S* 18, 292). HgNO_3Aq should not be used, as the HNO_3 produced oxidises the Hg_2S . Hg_2S is a black powder, easily decomposed by heat to HgS and Hg. According to Barfoed (*Bl* [2] 3, 183) the substance described as Hg_2S always contains some Hg, however it may be prepared.

MERCURIC SULPHIDE, HgS Mol w uncertain $\text{H F} [\text{Hg}_2\text{S}] = 8.220$ (Thomsen, *Z P C* 2, 21). This compound exists in two forms a black amorphous pp, and a red crystalline body known as *vermilion* or *cinnabar*.

Occurrence—In Illyria, Spain, Bohemia, Ural mountains, China, Japan, Mexico, California, Chili, and Peru.

Formation—1 By rubbing or heating to gether 1 pt S and 6 $\frac{1}{4}$ pts Hg.—2 By adding H_2S , or an alkaline sulphide, to solution of a mercuric salt.—3 By digesting Hg with an alkaline polysulphide.—4 By subliming a mixture of S and HgO or HgSO_4 .

Preparation—1 Amorphous black mercuric sulphide is prepared by passing excess of H_2S into slightly acidified HgCl_2Aq or $\text{Hg}(\text{NO}_3)_2\text{Aq}$, washing the pp thoroughly with dilute HNO_3Aq and then with water, and drying at a low temperature.—2 Red crystalline mercuric sulphide (vermilion) is prepared in different ways (a) A mixture of 6 pts Hg and 1 pt S is heated till combination occurs (attended generally with production of heat and light and partial projection of the mass), the product is powdered, mixed with a small quantity of S, and heated for some hours in a hard glass flask, sunk in sand in a wind-furnace, the flask is loosely closed by a charcoal stopper and is arranged so that the upper part is kept comparatively cold, red HgS sublimes on to the neck of the flask. The HgS is purified by grinding, boiling with KOH Aq , and washing with water. (b) Black amorphous HgS is heated with an alkaline persulphide. This may be done by triturating 100 pts Hg with 38 pts S for some hours, till black HgS is produced, heating with 25 pts KOH in 130–150 pts water to 45° – 50° for several hours (the water being replaced as it evaporates, and the mass being stirred from time to time) until reddening begins, after which the temperature is maintained at 45° till the whole is bright red. The red HgS is washed, and any Hg present is removed by levigation (Brunner, *P* 15, 598, cf Döbereiner, *S* 61, 860;

Firmenich, *D P J* 172, 370, Liebig, *A* 5, 289, 7, 49, Raab, *N R P* 24, 39) (HgCl_2Aq is poured into excess of dilute NH_4Aq , to the pp of NH_4HgCl thus produced is added conc $\text{Na}_2\text{S}_2\text{O}_3\text{Aq}$ in quantity rather more than sufficient to dissolve the pp, and the whole is heated for some time to $70^\circ\text{--}80^\circ$ (Hausmann, *B* 7, 1746) For details of the manufacture of *Vermilon v* DICTIONARY OF APPLIED CHEMISTRY—8 Colloidal mercuric sulphide, soluble in water, is obtained by ppg a dilute solution of a mercuric salt by H_2S , and washing with water or dilute H_2SAq for a long time (Winssinger, *Bl* [2] 49, 452)

Properties—The black sulphide is an amorphous, heavy powder, heated in a closed vessel, it yields a sublimate of red HgS , heated in air, Hg sublimes and SO_2 is formed, it is not acted on by dilute acids The red sulphide crystallises in hexagonal forms, $a=c=1.145$, S_G 8.1 to 8.99, polarises light, blackens by exposure to light (*v* Heumann, *B* 7, 750), heated to the sublimation temperature, red HgS is changed to the black variety (Fuchs, *P* 31, 581) Mitscherlich (*A* 12, 168) found V D of HgS at 670° to be 85.3, V and C Meyer found V D at 1660° to be 78, the formula HgS requires V D 116, a mixture of $\text{Hg} + \text{Hg} + 2\text{S}$ requires V D 77.3 The red sulphide reacts with acids more slowly than the black variety The colloidal sulphide is soluble in water, forming a solution which is black and opaque when conc, but brown, with a greenish tint by reflected light, when dilute, a very dilute solution may be boiled till all H_2S is expelled, or kept for some time, without change

Reactions—1 Heated in a closed vessel, HgS sublimes—2 Heated in air, SO_2 is formed, and Hg sublimes—3 Heated with solid alkalis or alkaline carbonates, Hg sublimes and alkaline sulphide remains—4 Heated with iron, tin, antimony, copper, zinc (and some other metals), a metallic sulphide and Hg are produced, HgS is decomposed by heating with finely divided Cu and water (*v* Heumann, *B* 7, 1388, 1486)—5 Digested for some time with iodine in KIAq , HgS is decomposed with formation of HgI_2 2KI and separation of S (Wagner, *J pr* 98, 23)—6 HgS is decomposed to Hg and SO_2 by heating with lead monoxide, Pb being separated.—7 HgS is scarcely attacked by dilute acids, conc nitric acid produces $\text{Hg}(\text{NO}_3)_2$ mixed with HgSO_4 , HNO_3Aq S G c 1.2 produces a white compound $2\text{HgS} \cdot \text{Hg}(\text{NO}_3)_2$.—8 According to Bolley (*A* 75, 239) HgS is at once decomposed by an ammoniacal solution of silver nitrate, with formation of a mercurammonium salt and Ag_2S —9 HgS is insoluble in caustic soda or sodium monosulphide solution, but it dissolves in a mixture of the two, i.e. in sodium hydrosulphide solution This solution contains a sulpho-salt (sulphohydrargyrate of sodium) The solution is readily obtained by adding KOHAq to ppd HgS , and passing in H_2S (excess of H_2S causes reppn of HgS) On evaporation, white crystals $\text{HgS} \cdot \text{K}_2\text{S} \cdot 5\text{H}_2\text{O}$ separate, mixed with KOH , these crystals are decomposed by water (Weber, *P* 97, 76, *cf* Brunner, *P* 15, 596) By ppg HgCl_2Aq with NH_4S sulphide, and adding KOHAq , a solution is obtained, which, on evaporation, yields KCl , and then a sulpho-salt mixed with KOH .

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According to Schneider (*P* 127, 488) the crystals of the sulpho-salt, when kept for some years in KOHAq , form six sided, lustrous, olive green tablets having the composition $2\text{HgS} \cdot \text{K}_2\text{S}$

Combinations—1 With alkali sulphides, *v supra*, Reactions No 9—2 Forms a compound with hydrogen sulphide, $31\text{HgS} \cdot \text{H}_2\text{S}$ Produced in solution by passing H_2S into HgS suspended in boiled water (Linder a Picton, *C J Proc* 1890 49)—3 With cuprous chloride to form $2\text{HgS} \cdot \text{Cu}_2\text{Cl}_2$, a yellow solid, obtained by boiling HgS with CuCl_2Aq and HClAq , and removing S by CS_2 ($3\text{HgS} + 2\text{CuCl}_2 = 2\text{HgS} \cdot \text{Cu}_2\text{Cl}_2 + \text{HgCl}_2 + \text{S}$) This compound is decomposed by boiling conc HClAq with evolution of a little H_2S and formation of Cu_2Cl_2 and HgCl_2 in solution, dilute hot $\text{H}_2\text{SO}_4\text{Aq}$ has no action, but with boiling conc H_2SO_4 , HCl and SO_2 are evolved and a compound of HgSO_4 and HgS is formed NaOHAq produces NaCl , HgS , and Cu_2O (*v* Heumann, *B* 7, 1390) 4 With mercuric chloride, and also with mercuric bromide, to form compounds $2\text{HgS} \cdot \text{HgX}_2$, ($\text{X} = \text{Cl}$ or Br) These sulphohaloid compounds are formed (a) by passing H_2S into excess of HgCl_2Aq or HgBr_2Aq , (b) by digesting freshly ppd HgS with boiling HgCl_2Aq or HgBr_2Aq (*H Rose*, *P* 13, 59), (c) by adding HCl (or HBr), or solution of a metallic chloride (or bromide), to HgS dissolved in mercuric acetate solution (*Palm*, *C C* 1863 120), (d) by heating HgS with 8-10 parts HgCl_2 or HgBr_2 , till the whole melts, and washing the cold mass with boiling water (Schneider, *P* 115, 167) The compounds are yellow white crystalline powders, decomposed by slow heating to HgS and HgX_2 , decomposed by alkali solutions, but not by HNO_3Aq or $\text{H}_2\text{SO}_4\text{Aq}$ (*cf* Barfoed, *J pr* 93, 230)—5 With mercuric iodide to form the sulpho iodide $\text{HgS} \cdot \text{HgI}_2$, a yellow solid obtained by digesting HgI_2 with less H_2SAq than suffices for complete decomposition (*H Rose*, *P* 13, 59), also by ppg HgO and HgI_2 in HClAq by a small quantity of H_2S (Rammelsberg, *P* 48, 175), also by saturating HgI_2 in HIAq with H_2S , and diluting (Kekulé) *Palm* (*C C* 1863 121) says that the yellow red pp obtained by adding HIAq , or a solution of an alkali iodide, to HgS dissolved in mercuric acetate solution, has the composition $2\text{HgS} \cdot \text{HgI}_2$.—6 With mercuric fluoride, to form the sulphofluoride $2\text{HgS} \cdot \text{HgF}_2$, obtained by passing a little H_2S into HgF_2 dissolved in HFAq (*H Rose*, *P* 13, 59)—7 With mercuric sulphate, to form several compounds, the chief of which are (1) $\text{HgSO}_4 \cdot \text{HgS}$, (2) $\text{HgSO}_4 \cdot 2\text{HgS}$, and (3) $\text{HgSO}_4 \cdot 3\text{HgS}$ These compounds are obtained, (1) by treating red HgS with warm $\text{H}_2\text{SO}_4\text{Aq}$ (*Palm*, *J* 1862 220), (2) by treating $2\text{HgS} \cdot \text{HgNO}_3$ (obtained by passing a little H_2S into $\text{Hg}(\text{NO}_3)_2\text{Aq}$ with $\text{H}_2\text{SO}_4\text{Aq}$ (Barfoed, *J* 1864 282, Kessler, *A Ch* [3] 6, 615), (3) by adding $\text{H}_2\text{S}_2\text{O}_8\text{Aq}$ to $\text{Hg}(\text{NO}_3)_2\text{Aq}$ and removing S by washing with CS_2 (*Spring*, *A* 199, 116, Wackenroder, *A* 60, 190)—8 With mercuric nitrate, to form $2\text{HgS} \cdot \text{Hg}(\text{NO}_3)_2$, a white solid, obtained by passing a little H_2S into $\text{Hg}(\text{NO}_3)_2\text{Aq}$, or by digesting freshly ppd HgS with $\text{Hg}(\text{NO}_3)_2\text{Aq}$, or by ppg a solution of HgS in mercuric acetate by HNO_3Aq or alkali nitrate solution (*H Rose*, Barfoed, *Palm*, *lc*) By heating HgS with HNO_3Aq , S.G. 1.2, to 120° in a closed tube, Gramp

Q

(*J pr* [2] 14, 299) obtained a white crystalline mass of $6\text{HgS} \cdot \text{HgO} \cdot 2\text{Hg}(\text{NO}_3)_2 \cdot 12\text{H}_2\text{O}$

Mercury, sulphobismide of, v *Mercuric sulphide, Combinations*, No 4

Mercury, sulphochloride of, v *Mercuric sulphide, Combinations*, No 4

Mercury, sulphocyanides of, v vol II p 850

Mercury, sulphofluoride of, v. *Mercuric sulphide, Combinations*, No 6

Mercury, sulpho-iodides of, v *Mercuric sulphide, Combinations*, No 5

Mercury, selenoselenide of, v *Mercury, selenosulphide of*, p 224

M M P M

MERCURY COMPOUNDS, ORGANIC Mercury forms compounds with two identical alcohol radicles, or with one alcohol radicle and one acid residue. Compounds of mercury with two different alcohol radicles appear to split up at the moment of their formation, e.g. $2\text{HgEtMe} = \text{HgMe}_2 + \text{HgEt}_2$, (Frankland, *A* 111, 57)

Mercury dimethide $\text{Hg}(\text{CH}_3)_2$, *Mercury dimethyl*. Mol w 230 V D 8.29 (calc 7.97) (93° – 96°)

Formation—1 When mercury is exposed with MeI to sunlight for a week, crystalline CH_3HgI is formed (Frankland, *A* 85, 361). When this body is distilled with KOH, lime, or KOCy, mercury dimethide passes over as a heavy liquid which may be purified by washing with water and rectifying over CaCl_2 (Buckton, *Pr* 9, 91, *A* 108, 103)—2 By the action of sodium-amalgam on MeI (10 pts) in presence of acetic ether (1 pt) (Frankland & Duppa, *C J* 17, 415, *A* 130, 105)

Properties—Colourless oil, with faint but mawkish taste. Insol water, v sol alcohol and ether. Dissolves phosphorus, resins, and caoutchouc. Very inflammable, burning with a bright flame.

Reactions—1 Iodine forms CH_3HgI and methane. Bromine acts in like manner.—2 SnCl_4 forms a crystalline compound which is decomposed by water with formation of CH_3HgCl .—3 Conc HClAq gives methane and CH_3HgCl .—4 Conc H_2SO_4 gives methane and $(\text{CH}_3\text{Hg})_2\text{SO}_4$.—5 PCl_5 gives CH_3HgCl .—6 KMnO_4 oxidises it to CH_3HgOH (Seidel).

Mercuric chloro-methide CH_3HgCl [170°] (Seidel, *J pr* [2] 29, 135) S G 4.063 (Schröder, *B* 12, 563). Formed from HgMe_2 by the action of HCl . Lammæ.

Mercuric iodomethide CH_3HgI [143°] From MeI and Hg in sunlight (Frankland). Formed also by heating HgMe_2 with HgI_2 . Small nacreous lammæ (from ether), insol water, m sol alcohol, v sol ether and MeI. Slightly volatile, emitting an unpleasant odour, and leaving a persistent nauseous taste. May be sublimed.

Mercuric nitromethide CH_3HgNO_2 [100°] From an alcoholic solution of CH_3HgI and AgNO_3 . Nacreous lammæ, v sol water, sl sol alcohol. Its solution is not pptd by KOH or baryta water, but HCl and soluble chlorides ppt CH_3HgCl (Strecker, *A* 92, 79).

Mercuric acetomethide $\text{CH}_3\text{Hg.OAc}$ [148°] Obtained by heating HgMe_2 with acetic acid at 180° (Otto, *Z.* [3] 6, 26). Thin tables,

with very offensive odour, nearly insol boiling water and cold HOAc .

Mercuric chloro-methylo iodide CH_2ClHgI [129°] Obtained by boiling CH_3I with alcoholic HgCl_2 (Sakurai, *C J* 41, 360). Silky plates (from alcohol). Decomposed by iodine into methylene chloro iodide and HgCl_2 .

Mercuric iodo-methylo-iodide CH_2IHgI [109°] Formed by leaving mercury and a little mercurous iodide in contact with methylene iodide in a sealed tube for several days (Wanklyn & Von Than, *C J* 12, 258, Sakurai, *C J* 37, 658). White crystals (from CH_2I_2), insol water, cold alcohol, ether, chloroform, EtI, and benzene, sl sol boiling alcohol, v sol methylene iodide. When heated with iodine dissolved in aqueous KI it is decomposed into CH_2I_2 and HgI_2 . Chlorine and bromine act in like manner. When heated with HClAq it is reduced to CH_3HgI .

Di-mercuric methyleno-di-iodide $\text{CH}_2(\text{HgI})_2$ [$c 230^\circ$] Obtained by exposing a mixture of an ethereal solution of methylene iodide with a little HgI_2 and an excess of mercury to sunlight (Sakurai, *C J* 39, 487). Formed also by the action of mercury on CH_2IHgI . Yellowish crystalline powder, insol all ordinary solvents, sl sol hot methylene iodide. When heated with dilute HClAq or with KIaq it yields CH_3HgI . Iodine forms HgI_2 and methylene iodide.

Mercuric methyleno-tri-iodide $\text{CH}(\text{HgI})_3$. From iodoform, mercury, and alcohol, by exposure to sunlight (Sakurai, *C J* 39, 488). Yellow mass, insol ordinary solvents and in CH_2I_2 . Gives iodoform when treated with iodine.

Mercuric diethide HgEt_2 . Mol w 258 (159°) S G 2.46 V D 9.97 (calc 9.4). Strecker (*C R* 39, 57, *A* 92, 97) by acting on EtI with mercury obtained the compound EtHgI . Dünhaupt (*Chem Gas* 1854, 263, *A* 92, 879) obtained the corresponding EtHgCl and EtHgBr by decomposing bismuth triethide with HgCl_2 or HgBr_2 . Buckton (*A* 109, 218, *C J* 18, 17) obtained HgEt_2 by the action of ZnEt_2 on HgCl_2 , and, lastly, Frankland and Duppa showed that mercuric diethide can be more easily obtained by the action of sodium amalgam on EtI in presence of acetic ether. The acetic ether remains in undiminished quantity at the end of the process, but nevertheless its presence is essential.

Preparation—Ethyl iodide (10 pts) is mixed with EtOAc (1 pt) and poured upon $\frac{1}{2}$ p.c. sodium amalgam, the flask being shaken and cooled in water. When separation of NaI has rendered the liquid pasty it is distilled off from a water bath and shaken with fresh amalgam. Finally it is mixed with water, and the oil dried over CaCl_2 and rectified.

Properties—Transparent colourless liquid, with faint ethereal odour, insol water, sl sol. alcohol and ether. Very poisonous. Burns with smoky flame. Takes fire in chlorine.

Reactions.—1 Acts violently on bromine and iodine, forming EtHgBr and EtHgI (Buckton, *A* 112, 220).—2 Alcoholic HgCl_2 forms EtHgCl .—3 Conc HClAq forms ethane and EtHgCl .—4 H_2SO_4 gives ethane and $(\text{EtHg})_2\text{SO}_4$.—5. *Sodium* forms a bulky grey spongy mass, which

takes fire in air and violently decomposes when heated—6 Heated with finely granulated zinc it yields ZnEt₂ and mercury (Franklanda Duppa, *C J* 17, 29) Cadmium forms CdEt₂. Bismuth gives BiEt₃.—7 Mercuric ethide is decomposed when heated at 150° with copper, iron, silver, or gold, but ethides of these metals are not formed 8 KMnO₄ forms EtHgOH—9 When heated with PCl₅ it forms PEtCl₄.—10 Iodoform at 90° gives acetylene and ethylene (Suida, *M* 1, 716)

Mercuric chloro-ethide EtHgCl [190°] (Seidel) SG 3 48 Formed by the action of HgCl₂ on BiEt₃, by treating HgEt₂ with alcoholic HgCl₂, and by the action of HgCl₂ on ZnEt₂. Ppd by adding NaCl to an aqueous solution of EtHgNO₃. Obtained also by treating EtHgOH with HCl. Light iridescent laminae (from alcohol). Nearly insol water, sl sol ether, v sol boiling alcohol. Sublimes at 40° With Ag₂CO₃ it gives (EtHg)₂CO₃, which crystallises with difficulty and is easily decomposed by heat and by acids. Ag₂PO₄ gives a very soluble phosphate (Dunhaupt)

Mercuric iodomethide EtHgI Formed by the action of mercury on EI in diffused daylight. Decomposed by direct sunlight (Strecker, *A* 92, 75) Shining laminae, insol water, sol alcohol and ether. Sublimes at 100° Dissolves without change in aqueous NH₃ or KOH With ZnEt₂ it yields ZnI₂ and HgEt₂

Mercuric ethylo hydroxide EtHgOH Obtained by adding moist Ag₂O to a boiling alcoholic solution of EtHgCl, filtering, and evaporating *in vacuo*. Strongly alkaline liquid, which blisters the skin. V sol water. Decomposes ammonium salts with expulsion of NH₃. It does not liberate KOH or MgO from their salts, but it ppt salts of Al, Zn, Sn, Cu, Au, and Pt. With a large excess of H₂S it forms a white ppt turning through orange to black. With metallic zinc it forms ZnEt₂. With acids it forms crystalline salts, eg HCl yields EtHgCl

Mercuric bromo ethide EtHgBr **Mercuric ethylo bromide** Formed from BiEt₃ and HgBr₂, or by treating EtHgOH with HBr (Dunhaupt, *A* 92, 379) Resembles EtHgCl

Mercuric ethylo cyanide EtHgCy Formed by saturating an alcoholic solution of EtHgOH with HCN. Crystallises readily, v sol alcohol and ether. Very volatile. Gives off a repulsive odour when heated

Mercuric ethylo nitrate EtHgNO₃, **Mercuric nitro ethide** From EtHgI and AgNO₃. Colourless prisms, v e sol water, m sol alcohol (Strecker)

Mercuric ethylo sulphate (EtHg)₂SO₄ Formed from EtHgCl and Ag₂SO₄. Shining laminae (from alcohol)

Mercuric ethylo sulphide (EtHg)₂S Obtained by adding ammonium sulphide to an alcoholic solution of EtHgCl. Yellowish white powder, v sol alcohol, ether, and CS₂. The alcoholic solution is decomposed by evaporation, leaving HgS

Mercuric ethylo-acetate EtHgOAc [178°] Obtained by dissolving HgEt₂ in HOAc (Otto, *Z* 1870, 25)

Mercuric dipropyl HgPr₂, (190°) (O); (180°) (Sohlscherbakoff, *J R* 18, 353) SG 12 214 Formed by the action of propyl iodide on sodium-amalgam in presence of acetic ether (Cahours,

C. R 76, 133, 1383) Oil, sl sol alcohol, v sol ether. With iodine and bromine it forms products which may be crystallised from alcohol. The effect of small additions of HgPr₂ of Hg(C₂H₅)₂, of Hg(C₃H₇)₂, and of HgPh₂ on the freezing point of ethylene bromide has been studied by Louise and Roux (*C R* 107, 600)

Mercuric diisobutyl Hg(CH₂Pr)₂ (206°) SG 1-1835 Obtained by treating a mixture of isobutyl iodide and acetic ether with 2 p.c. sodium amalgam (Cahours). Colourless liquid. Iodine forms CH₂Pr HgI, whence Ag₂O forms CH₂Pr HgOH. Bromine acts in like manner

Mercuric diisocamyl Hg(C₈H₁₇)₂ SG 2 1666 From isocamyl iodide (5 pts), acetic ether (1 pt), and sodium amalgam (Franklanda Duppa, *A* 130, 110). Colourless oil, decomposed on distillation even *in vacuo*. May be distilled with steam. Insol water, v sl sol alcohol, v sol ether. With chlorine it forms dense fumes of C₈H₁₇HgCl. It reacts with great violence with bromine and iodine. With an alcoholic solution of HgCl₂ it readily forms C₈H₁₇HgCl [86°], which crystallises from alcohol in hair like needles. An ethereal solution of Hg(C₈H₁₇)₂ treated first with an alcoholic solution of iodine and then with solid iodine forms isocamyl iodide and C₈H₁₇HgI [122°], which crystallises from alcohol in pearly scales. With zinc at 130° C₈H₁₇HgI forms zinc isocamyl and zinc amalgam

Mercuric dioctyl Hg(C₈H₁₇)₂ SG 17 1342 Prepared by the action of sodium amalgam on octyl iodide in presence of a little acetic ether (Eichler, *B* 12, 1880). Liquid. Insol water, sol alcohol, ether, and C₆H₆. Decomposes at 200° into Hg and hexadecane (dioctyl)

Mercuric octylo iodide C₈H₁₇HgI white silky pp. Prepared by the action of an alcoholic solution of iodine on mercury dioctyl

Mercuric octylo chloride C₈H₁₇HgCl White pp. Prepared by the action of mercuric chloride on mercury dioctyl

Mercuric octylo-hydroxide C₈H₁₇HgOH [75°] Prepared by the action of silver oxide on an alcoholic solution of mercury dioctyl (Eichler, *B* 12, 1881). Yellow leaflets. Insol cold, sl sol hot, water, v sol alcohol. Alkaline

Mercuric allylo-iodide C₃H₅HgI [135°] S (CS₂) 187 at 49° Formed by shaking allyl iodide with mercury, and extracting the resulting yellow mass with hot alcohol or ether (Zinin, *A* 96, 363, cf Linnemann, *A* 140, 180). It is best to mix the allyl iodide with an equal volume of alcohol (Oppenheim, *B* 4 670). Silvery scales, which turn yellow on exposure to light, especially on drying. It even turns yellow in the dark. Sublimes at 100° Sl sol cold alcohol, nearly insol water. Decomposes above its melting point. AgNO₃ added to its alcoholic solution ppt all the iodine as AgI. Moist Ag₂O ppt AgI, and the filtrate on evaporation leaves a strongly alkaline syrupy mass which forms salts with acids. HI easily decomposes it, forming HgI₂ and propylene. Iodine forms HgI₂ and allyl iodide (Linnemann, *A* 133, 133, *Suppl* 3, 262). PBr₃, acetyl chloride, and BzCl have no action on mercuric allyl iodide. On shaking it with a solution of ZnEt₂ in ether an energetic reaction sets in, the products being mercury, ZnI₂, diallyl, and HgEt₂. A solution

of K₂O acts quickly in the cold, forming diallyl, mercury, HgCy₂, and KI, if in distilling off the diallyl the distillation is not stopped in time an explosion occurs. Aqueous KI at 100° forms mercury, diallyl, and KHgI. The mercury allyl compounds attack the skin, producing blisters which appear after the lapse of 7 hours. The compound C₃H₅HgCl obtained by the action of HCl on the hydroxide is sparingly soluble (Krasowsky, *Z* 6, 527).

Mercuric propargyl iodide C₃H₃HgI. From propargyl iodide and mercury (Henry, *B* 17, 1152). Small yellowish crystalline masses.

Mercuric diphenyl Hg(C₆H₅)₂ [120°] (above 800°) SG 2318. Formed by heating a mixture of bromo benzene, benzene, HgCl₂, and sodium (Michaelis a. Reese, *B* 15, 2876). Prepared by boiling for some time a solution of bromo-benzene (10 pts) in an equal volume of xylene with 27 p.c. sodium amalgam and a little acetic ether (1 pt). The product is recrystallised from benzene (Dreher a. Otto, *Z* [2] 4, 685, 6, 9, *A* 154, 93). Small white needles or prisms, turning yellow in daylight. Insol water, v sol chloroform, CS₂, and benzene, m sol ether and boiling alcohol. May be sublimed. Partially decomposed on distillation into mercury, diphenyl, benzene, and charcoal.

Reactions—1 Dry HCl gives benzene and HgCl₂. HI, nitric acid, HBr, and H₂SO₄ act in like manner.—2 When melted with sulphur (2 at) the products are HgS and phenyl mercaptan.—3 Iodine (2 at) in alcoholic or CS₂ solution forms PhHgI and iodobenzene. Excess of iodine forms HgI₂ and iodo benzene.—4 Chlorine forms PhHgCl and chloro benzene. Br acts in like manner.—5 Glacial acetic acid yields benzene and PhHgOAc.—6 Aluminium foil at 130° forms AlEt₃ [230°] (Friedel a. Crafts, *A Ch* [6] 14, 460).—7 Alcoholic HgCl₂ at 100° gives PhHgCl.—8 When boiled with sodium, benzene and sodium amalgam are formed.—9 SO₂ gives mercuric benzene sulphonate (Otto, *J pr* [2] 1, 179).—10 Allyl iodide forms diallyl, diphenyl, and PhHgI (Suida, *M I* 715).—11 C₆H₅SO₂Cl at 160° forms C₆H₅SO₂C₆H₅ and PhHgCl (Otto, *B* 18, 246).—12 COCl₂ CO₂Et forms phenylglyoxylic ether (Claisen a. Morley, *B* 11, 1596). 13 When oxidised by KMnO₄ the hydroxide PhHgOH is probably first formed, but on adding HCl this is changed to PhHgCl (Seidel, *J pr* [2] 29, 134, Otto, *J pr* [2] 29, 136).

Mercuric phenylo chloride C₆H₅HgCl. **Mercuric chlorophenylide** [250°]. Obtained by slowly passing chlorine over HgPh₂ or into its solution in CS₂. Prepared by heating HgPh₂ with HgCl₂ in alcoholic solution at 110°. Trimetric tables (from benzene). May be sublimed. Sl sol alcohol and benzene, insol water.

Mercuric phenylo-bromide C₆H₅HgBr [275°]. Resembles the preceding in preparation and properties. Br forms HgBr₂ and bromobenzene. When heated with an alcoholic solution of K₂S the products are KBr, HgS₂, and HgPh₂.

Mercuric phenylo-iodide C₆H₅HgI [266°]. From HgPh₂ and I in alcohol. Satiny tables (from alcohol benzene). Insol water, nearly insol. cold alcohol, ether, and benzene, m. sol. hot benzene and chloroform, v. sol. CS₂.

May be partially sublimed. Sodium amalgam reduces it in alcoholic solution to HgPh₂.

Mercuric phenylo hydroxide C₆H₅HgOH. From PhHgCl by boiling with alcohol and moist Ag₂O. White trimetric prisms (from alcohol), sol boiling water, alcohol, and benzene, sl sol cold water. Softens at 160°, but is not melted at 200°. The aqueous solution is alkaline in reaction. Expels NH₃ from its salts, ppts alumina from alum, and absorbs CO₂ from the air.

Mercuric phenylo cyanide PhHgCy. [204°]. From HgPh₂ and HgCy₂ in alcoholic solution at 128°. Long slender trimetric prisms, v sl sol boiling water, m sol boiling alcohol and benzene. Conc HClAq at 120° forms HgCl₂, formic acid, benzene, and NH₃. Alcoholic KOH forms mercury, benzene, and potassium cyanate. Iodine forms PhHgI and ICy. H₂S forms HgS, benzene, and HCy.

Mercuric phenylo-sulphocyanide PhHgSCy [227°]. From HgPh₂ and Hg(SCy)₂. Silky plates, sol boiling alcohol.

Mercuric phenylo-nitrate PhHgNO₃ [c 167°]. From PhHgCl and AgNO₃ in alcohol. Trimetric silky plates, v sl sol boiling water. Conc HClAq gives benzene and mercuric nitrate.

Mercuric phenylo carbonate (PhHg)₂CO₃. From PhHgCl and Ag₂CO₃. Small white needles, sl sol boiling water, m sol boiling alcohol and benzene. Not decomposed on fusion.

Mercuric phenylo-formate PhHgOCHO [171°]. From HgPh₂ and concentrated formic acid. Small tables.

Mercuric phenylo acetate PhHgOAc. [149°]. Obtained by boiling HgPh₂ with excess of HOAc, ppg with water, and crystallising from hot water. Radiate groups of prisms, sl sol cold, m sol hot, water, m sol alcohol and benzene. Decomposed by distillation yielding diphenyl, benzene, AcO, HOAc, carbon, and mercury. Boiling aqueous HCl forms benzene, HgCl₂, and HOAc. Other mineral acids act in like manner. Sodium amalgam reduces it in alcoholic solution to benzene. Iodine, acting on its aqueous solution, forms HgI₂, iodo benzene and HOAc. H₂S forms HgS, benzene, and HOAc.

Mercuric phenylo-propionate PhHgOC₂H₃O [c 166°]. From HgPh₂ and boiling propionic acid. Crystals, sol hot water, alcohol, and benzene.

Mercuric phenylo-myristate PhHgOC₄H₇O. From HgPh₂, myristic acid, and alcohol at 120°. Small scales, sol boiling alcohol and benzene. Boiling HClAq splits it up into HgCl₂, benzene, and myristic acid (Otto, *J pr* [2] 1, 179).

Mercuric tetra-methyl-di-amido-di-phenyl (NMe₂C₆H₄)₂Hg [169°]. Formed by the action of sodium amalgam in presence of EtOAc on *p*-bromo-di-methyl-aniline dissolved in xylene (Sohenk a. Michaelis, *B* 21, 1501). Colourless crystals, turning green in air. Crystallises from benzene with C₆H₆ (1 mol.) v sol chloroform, and benzene, v sl sol alcohol and ether.

Mercuric di-*o*-tolyl Hg(C₆H₄Me)₂ [107°]. From *o* bromo toluene sodium-amalgam, and

acetic ether (Ladenburg, *A* 178, 162). Large trichinic tables (from benzene)

Mercuric o-tolylo-chloride

[2] $C_6H_4MeHgCl$ *o*-Tolyl-mercuric chloride [146°] Formed by treating mercuric *o* tolyl with $HgCl_2$ in ethereal solution (Michaëlis a Genzken, *A* 242, 180) Needles, v sol chloroform, m sol alcohol and benzene, sl sol ether and petroleum ether Sublimes with partial decomposition

Mercuric m-tolylo-chloride

[3] $C_6H_4MeHgCl$ [160°] Obtained by boiling $(C_6H_4Me)_2SbHgCl_2$ with alcohol (M a G) Needles (from alcohol), v sol chloroform and benzene, m sol alcohol H_2S passed into its alcoholic solution ppts HgS

Mercuric di-p tolyl $Hg(C_6H_4Me)_2$ [238°] Formed from *p* bromo toluene, sodium amalgam, and $EtOAc$ (Dreher a Otto, *A* 154, 171, Ladenburg, *A* 173, 163) Needles or tables, insol water, sl sol cold alcohol, m sol hot benzene, CS_2 , and $CHCl_3$ May be distilled Boiling conc $HClAq$ gives toluene and $HgCl_2$

Mercuric p tolylo chloride

$C_6H_4MeHgCl$ [187°] Small tables (Otto, *J pr* [2] 1, 185)

Mercuric p tolylo iodide C_6H_4MeHgI [220°] From mercuric di *p* tolyl and iodine Trimetric satiny tables, insol water, sl sol boiling alcohol, m sol hot benzene May be sublimed

Mercuric p tolylo-acetate

$C_6H_4MeHgOAc$ [153°] Small trimetric prisms, nearly insol cold water, sl sol boiling water, m sol alcohol Behaves like the corresponding phenyl compound

Mercuric di-p xylyl $(C_6H_4Me)_2Hg$ [123°]

Prepared by the action of sodium amalgam on bromo *p* xylene (Jacobsen, *B* 14, 2112) Thick prisms, sol CS_2 , chloroform, and benzene, sl sol alcohol and ether

Mercuric di-m-xylyl [1 3 4] $(C_6H_4Me)_2Hg$ [170°] From bromo *m* xylene and 2 *p* c sodium amalgam by heating at 145° for 12 hours (Weller, *B* 20, 1718) Slender felted needles, sl sol ether, alcohol, and cold benzene, v sol hot benzene Boiling $HClAq$ gives $HgCl_2$ and *m* xylene PCl_5 forms $C_6H_4Me_2PCl_2$ (257°) and $C_6H_4Me_2HgCl$

Mercuric di n-propyl-di phenyl $Hg(C_6H_5Pr)_2$ [110°] From BrC_6H_5Pr [1 4] and sodium-amalgam (R Meyer, *J pr* [2] 34, 103) Long needles

Mercuric di cymyl $Hg(C_{10}H_{11})_2$ [134°]

Formed by heating a mixture of bromo cymene with xylene and sodium amalgam with addition of a little acetic ether (Paternò a Colombo, *B* 10, 1749) Long slender needles, m sol boiling alcohol, benzene, and xylene

Mercuric di pentamethyl-phenyl $Hg(C_6Me_5)_2$

[266°] From bromo-pentamethyl-benzene, chloroformic ether, and sodium-amalgam (Jacobsen, *B* 22, 1220) Prisms (from xylene), v sl sol alcohol and ether, v sol hot xylene

Mercuric di-naphthyl $Hg(C_{10}H_7)_2$ Mol w

464 [243°] SG 1.93 (Schröder, *B* 12, 564) Formed by boiling (a) bromo naphthalene with several times its volume of crude xylene, pasty sodium-amalgam, and a little acetic ether for 19 hours, and filtering while hot (Otto a Moraes, *A* 147, 164, *Z* [2] 3, 877, 4, 162). Small minute, trimetric prisms (from benzene

or CS_2) Not affected by air or light Insol water, sl sol boiling alcohol, cold benzene or ether, m sol hot CS_2 , chloroform, and benzene Decomposed by heat Conc $HIAq$ forms HgI_2 and naphthalene HCl and HBr act in a similar way Iodine forms HgI_2 , $C_{10}H_7HgI$, and ultimately $C_{10}H_7I$ It does not react with $HgCl_2$

Mercuric naphthyllo-bromide

$C_{10}H_7HgBr$ [196°] From $Hg(C_{10}H_7)_2$ and bromine Formed also by heating $Hg(C_{10}H_7)_2$ with $HgBr_2$ Needles, insol water, sol hot alcohol

Mercuric naphthyllo iodide $C_{10}H_7HgI$ [185°] From mercuric di naphthyl and iodine (1 mol) in CS_2 Soft satiny needles or dendritic groups Not altered by light, insol water, sl sol hot alcohol and benzene Sodium-amalgam converts it into mercuric di naphthyl

Mercuric naphthyllo acetate

$C_{10}H_7HgOAc$ [154°] Formed, together with naphthalene, by heating mercuric di naphthyl with excess of $HOAc$ Small needles (from alcohol), insol water, v sol hot $HOAc$, alcohol, CS_2 , benzene, and chloroform, m sol ether Hot $HClAq$ gives naphthalene and $HOAc$ Sodium amalgam acting on its alcoholic solution forms naphthalene and $HOAc$ Water at 140° has no action

Mercuric naphthyllo butyrate

$C_{10}H_7HgOC_4H_9O$ [200°] Very slender needles, sol hot water

MESACONIC ACID $C_6H_4O_4$ *aa*

$CH_2C(CO_2H)CHCO_2H$ Mol w 130 [202°] S 2.7 at 18°, 118 at 100° (Pebal, *A* 78, 130), 2.6 at 14°, 3.4 at 22° (Baup, *A* 81, 97) S (90 p c alcohol) 30.6 at 17°, 95.7 at 78° (P), (88 p c alcohol) 38 at 22° (B) R_{20} 46.9 in a 3.69 p c aqueous solution (Kanonnikoff, *J pr* [2] 31, 349) Heat of solution - 5493 Heat of combustion 27,334 (Gal a Werner, *Bl* [2] 47, 159) H C 479.063 (Lougounne, *C R* 106, 1291)

Formation—1 By boiling a dilute solution of citraconic acid for half an hour with one-sixth of its volume of nitric acid Mesaconic acid is deposited on cooling (Gottheb, *A* 77, 268, Pebal, *A* 78, 129, Baup, *A* 81, 96)—2 By boiling citric acid with conc $HIAq$ or $HBrAq$ (Kekulé, *A Suppl* 2, 94, Fitig, *A* 188, 77, 80). 3 By heating a conc aqueous solution of itaconic or citraconic acid at 180° to 200°, CO_2 and an empyreumatic oil being also formed (Swarts, *Bull Acad. Royale Belgique*, [2] 36, No 7) 4 From *citra-* and *mesa-* di bromo pyrotartaric acids by heating with KI and copper at 150° (Swarts, *Z* 1868, 259)—5 The mono-anilide is heated with (3 mol of) KOH and a little water for 1½ hours in the water bath The aniline separates out and is removed by means of ether, the acid being ppd by means of SO_2H_2 (Anschütz, *A* 254, 136)—6 'Oxy-tetric acid,' which is formed from methyl aceto acetic acid by successive treatment with bromine and alcoholic potash (Demarçay, *A Ch* (5) 20, 473), is identical with mesaconic acid (Gorboff, *J R* 1887, 605, Cloëz, *Bl* [3] 3, 698, 602)

Preparation—By evaporating a mixture of citraconic anhydride (10 pts), water (22 pts), and conc HNO_3 (3 pts), until red fumes begin to appear The product is crystallised from water (Fitig, *A* 188, 73)

Properties—Slender needles (from water) or

prisms (from alcohol), sl sol cold, v sol hot, water V sol alcohol and ether. May be sublimed without decomposition. Its molecular weight, determined by Raoult's method, is identical with that of citraconic and of itaconic acid (Paternò, *B* 21, 2157). Mesaconic acid is slightly coloured by FeCl_3 , and the solution when boiled yields a brown gelatinous pp which re-dissolves on cooling. The addition of more FeCl_3 prevents its re-dissolving. Neutral solutions of mesaconates give a brown pp with FeCl_3 , insoluble in excess of the reagent or when heated.

Reactions—1 Conc HIAq at 160° forms pyrotartaric acid—2 Sodium amalgam also reduces it to pyrotartaric acid. The same reduction may be effected by zinc dust (Bottinger, *B* 9, 1821)—3 Bromine has no action in the cold (difference from citraconic acid) but above 60° it combines, forming meso di bromo pyrotartaric acid [170°] (Kekulé, *A Suppl* 2, 85)—4 Chlorine passed into water in which mesaconic acid is suspended forms a solution which when evaporated leaves chloro citramalic acid $\text{C}_4\text{H}_5\text{ClO}_4$ (chloro oxy pyrotartaric acid). The same acid is formed, together with tri chloro acetone, by passing chlorine into an aqueous solution of sodium mesaconate—5 Does not combine with HBr in the cold, but by prolonged heating with HBrAq at 100° or 140° it is converted into citra bromo pyrotartaric acid, an acid that is formed also by the union of HBr with citraconic acid in the cold—6 When repeatedly heated with fuming HClAq at 160° it is converted into citra chloro pyrotartaric acid [130°], which is decomposed by boiling water into HCl and mesaconic acid (Swarts)—7 The electrolysis of potassium mesaconate yields, at the positive pole, CO , and allylene (Aarland, *J pr* [2] 6, 256, 7, 142)—8 AsCl_3 forms citraconic anhydride. Mesaconic anhydride has not been obtained (Petrì, *B* 14, 1636)—9 On evaporating a solution of the aniline salt water is not eliminated from the molecule (difference from citraconic acid).

Salts— $\text{NH}_4\text{A}''$ S 12.5 at 15° . Very small prisms— BaA'' 4aq. monoclinic crystals, more sol water than barium citraconate and itaconate (Petrì, *B* 14, 1634)— BaHA'' 2aq. pearly hexagonal plates— CaA'' 5aq. S 6 at 20° . Very small needles, insol alcohol— PbA'' 1½aq. deposited in the cold as a crystalline pp, v sl sol water— PbA'' 4aq. deposited from a hot solution as an amorphous pp— $\text{PbH}_2\text{A}''$ (dried at 100°). Small needles— $(\text{HO Pb})_2\text{A}''$ 2 ppd by adding lead subacetate to a solution of sodium mesaconate (Otto, *A* 127, 182)— CuA'' 2aq— AgA'' crystalline pp, sl sol water— AgHA'' needles, m sol hot water.

Methyl ether MeA'' (205° 1 V) S 8 at 15° SG $\frac{15}{15}$ 1.1254, $\frac{30}{15}$ 1.1138 MM 1.154 at 24° μ_D 1.4564 at 16° (Gladstone), 1.4570 at 17° (Knops, *A* 248, 195), μ_H 1.4813 at 16° (Gladstone). From the acid, MeOH , and HCl (Perkin, *C J* 39, 556).

Ethyl ether (229° 1 V) SG $\frac{15}{15}$ 1.051, $\frac{30}{15}$ 1.039 (Perkin), $\frac{32}{15}$ 1.0468 (Knops, *A* 248, 196) MM 1.168 at 25° μ_D 1.4433, μ_H 1.4727 at 16° (Gladstone, cf Knops). Formed by the action of alcohol and HCl on mesaconic and on citraconic acids.

Chloride $\text{C}_4\text{H}_4(\text{COCl})_2$. (80° at 17 mm).

Prepared by the action of PCl_5 on mesaconic acid or citraconic anhydride (Petrì, *B* 14, 1634). Colourless liquid.

Amide $\text{C}_4\text{H}_5\text{O}_2(\text{NH}_2)_2$: [177°], colourless flat crystals, sol water.

Anilide $\text{C}_4\text{H}_5\text{O}_2(\text{NHPh})_2$ [186°], flat white silky needles, sol alcohol and ether, sl sol water. Heated to 268° it decomposes into aniline and citraconenil (the phenylimide of citraconic acid) (O Strecker, *B* 15, 1639).

Mono anilide $\text{C}_4\text{H}_5(\text{CO}_2\text{H})(\text{CONHPh})$. [158°] Formed by heating citraconic anhydride with aniline at 170° , rectifying *in vacuo* the resulting phenylimide (172° at 12 mm), dissolving it in baryta water, ppg excess of barium by CO_2 , and then adding HCl (Anschutz, *A* 254, 133). Identical with the product which separates from an aqueous solution of acid aniline citraconate on standing. White powder. Converted by heating in a sealed tube with KOH into mesaconic acid.

Constitution—Mesaconic acid stands to citraconic acid in the same relation that fumaric acid stands to maleic acid. This is shown by their behaviour towards Br, HBr, and aniline, and by the physical constants of their ethers. Hence mesaconic acid is methyl fumaric acid, while citraconic acid is methyl malic acid. The constitution of mesaconic acid will therefore be known as soon as that of fumaric acid has been satisfactorily determined (v MALEIC ACID).

MESA-DI BROMO PYROTARTARIC ACID α . DI BROMO PYROTARTARIC ACID

TRIMESIC ACID $\text{C}_6\text{H}_3\text{O}_6$, α $\text{C}_6\text{H}_3(\text{CO}_2\text{H})_3$ [135] Benzene α tri carboxylic acid Mol w 210 [α 325°] H C v 768,500 H C p 767,600 H F 285,400 (Stohmann, Kleber & Langbein, *J pr* [2] 40, 140).

Formation—1 By the oxidation of mesitylene or mesitylenic acid by chromic acid mixture (Fittig, *A* 141, 153)—2 By the oxidation of uvitic acid (Baeyer, *Z* 1868, 119, Fittig, *A* 147, 301), and of α tri ethyl benzene (Jacobsen, *B* 7, 1435, Friedel & Baisohn, *Bl* [2] 34, 636) by chromic acid mixture—3 One of the products obtained by heating hydromellitic or isohydromellitic acid with conc H_2SO_4 (Baeyer, *A Suppl* 7, 40, 48)—4 By heating mellitic acid with glycerin (Baeyer, *A* 166, 340)—5 In very small quantity by fusing the corresponding bromo benzene sulphonic acid with sodium formate (Bottinger, *B* 7, 1781)—6 By fusing benzene α trisulphonic acid with KCy , and saponifying the resulting nitrile with KOH (Jackson & Wing, *Am* 9, 347)—7 From di sodium saheylate $\text{C}_6\text{H}_4(\text{ONa})_2\text{CO}_2\text{Na}$ by heating in a current of carbonic acid, converting the resulting $\text{C}_6\text{H}_4(\text{OH})(\text{CO}_2\text{H})_2$ into $\text{C}_6\text{H}_4\text{Cl}(\text{CO}_2\text{H})_2$, and reducing with zinc and dilute acids—8 By the polymerisation of propionic acid, which takes place to some extent when it is exposed for some weeks to sunlight, air being excluded (Baeyer, *B* 19, 2185)—9 Trimesic ether is formed by the action of sodium upon a mixture of ethyl formate and ethyl acetate, formyl acetic ether $\text{HCO CH}_2\text{CO}_2\text{Et}$ is probably first formed and subsequently condensed (Putti, *B* 20, 537). When mixtures of methyl formate and ethyl acetate, or of ethyl formate with methyl acetate

are used a mixture of ethyl and methyl trimesates is obtained

Properties—Colourless prisms (from water), partially sublimes before melting M sol cold water and ether, v sol hot water, v e sol alcohol When distilled with lime it yields benzene

Salts— $\text{NaH}_2\text{A}'''$ plates, sl sol cold water— $\text{Na}_2\text{A}'''$ — $\text{KH}_2\text{A}'''$ needles, sl sol cold water— $\text{Ca}_2\text{A}'''$, aq nodules— $\text{Ba}_2\text{A}'''$, aq (dried at 150°) needles, almost insol cold, v sl sol boiling water (difference from mesitylenic acid)— $\text{BaH}_2\text{A}'''$, 2aq slender hair like needles, sl sol cold water— $\text{Zn}_2\text{A}'''$, 2aq glittering prisms, almost insol cold water— $\text{Cu}_2\text{A}'''$, 2aq (dried over H_2SO_4) Pale blue pp— $\text{Ag}_2\text{A}'''$

Methyl ether $\text{Me}_2\text{A}'''$ [143'] Small silky needles H F 249,500 (Stohmann, *J pr* [2] 40, 353)

Ethyl ether $\text{Et}_2\text{A}'''$ [133'] Prepared by adding a mixture of formic and acetic ethers by drops to twice the amount of ether, in which sodium is placed On adding dilute H_2SO_4 , an oil is obtained, which gives a blue violet colour with FeCl_3 , and on standing in a desiccator deposits crystals of trimestic ether (Wiskenus, *B* 20, 29 10) Long glistening prisms A mixture of this ether and the preceding in equal proportions melts at 105° to 110°

Sulpho trimestic acid *Amide*

$\text{C}_6\text{H}_2(\text{SO}_2\text{NH}_2)(\text{CO}_2\text{H})_2$ Formed by oxidising the amide of either of the sulpho mesitylenic acids with KMnO_4 (Jacobsen, *A* 206, 203)— $\text{KH}_2\text{A}'''$, 2aq crystalline mass, m sol water Conc HCl aq at 210° gives trimestic acid, NH_3 , and H_2SO_4 Potash fusion gives oxy trimestic acid

MESICERIN v TRI OXY MESITYLENE

MESIDIC ACID v UVITIC ACID

MESIDINE $\text{C}_6\text{H}_3\text{N}$ v e

$\text{C}_6\text{H}_2(\text{CH}_3)_2\text{NH}_2$ [1 3 5 6] *Amido mesitylene* (229') (Ladenburg, *A* 179, 172) SG 963

Formation—1 By boiling nitro mesitylene with tin and HCl aq (Fitting a Storer, *A* 147, 1)—2 From di methyl aniline methyl iodide, by heating in a sealed tube at 335° (Hofmann, *B* 5, 715, 8, 61)—3 By heating *um* xyldine hydrochloride or *cm* xyldine hydrochloride with methyl alcohol at 300° (Eisenberg, *B* 15, 1012, Nolting a Forel, *B* 18, 2631)—4 By heating aniline hydrochloride with MeOH at 300° (Lumpach, *B* 21, 640)

Properties—Liquid Gives *m* xyloquinone on oxidation When heated with MeOH and HCl for forty eight hours at 230° it yields dimethyl mesidine (c 215°) ClCO Et yields $\text{O}_2\text{H Me}$, $\text{NH CO}_2\text{Et}$ [629'] (Eisenberg, *B* 15, 1016)

Salts— B^+HCl feathery crystals or prisms, v sol water and alcohol— $\text{B}^+\text{H}_2\text{SnCl}_4$ Sparingly soluble needles— $\text{B}^+\text{H}_2\text{PtCl}_6$ — $\text{B}^+\text{H}_2\text{H}_2\text{O}_4$ plates, sl sol cold water

Acetyl derivative $\text{C}_6\text{H}_2(\text{CH}_3)_2\text{NHAc}$ [217'] Prisms (from alcohol) May be sublimed

Benzoyl derivative $\text{C}_6\text{H}_2(\text{CH}_3)_2\text{NHBz}$ [204'] Needles (Schaek, *B* 10, 1711)

MESIDINE SULPHONIC ACID $\text{C}_6\text{H}_2\text{NSO}_3$

v e $\text{C}_6\text{H}_2(\text{NH}_2)(\text{CH}_3)_2\text{SO}_3\text{H}$ [6 1 3 5 4] *Amido-mesitylene sulphonic acid* Formed from nitro mesitylene sulphonic acid by reduction with NH_3 and H_2S (Rose, *A* 164, 70) Slender prisms or needles (containing aq) (from water), v sol. hot

alcohol, sl sol cold water Does not combine with HCl or H_2SO_4 — BaA' , nodules, m sol. cold water— MgA' , 8aq— ZnA' , 5aq— PbA' , aq. crystalline, v sol cold water— AgA'

MESITENE LACTONE v vol 1 p 21 and

BROMO MESITENE LACTONE

TRI-MESITIC ACID v PYRIDINE TRI CARB OXYLIC ACID

MESITOL $\text{C}_6\text{H}_2\text{O}$ v e

$\text{C}_6\text{H}_2(\text{CH}_3)_2(\text{OH})$ [1 3 5 6] *Oxy mesitylene* [699] (219.5° 1 V) Obtained from mesidine by the diazo reaction, and by fusing mesitylene sulphonic acid with potash (Biedermann a Ledoux, *B* 8, 57, 250, Jacobsen, *A* 195, 268) Crystals, v e sol alcohol and ether Volatile with steam Insol NH_4aq and aqueous Na_2CO_3 , v sol NaOH aq Not coloured by FeCl_3 Forms a sulphonic acid, which has an easily soluble barium salt, coloured deeply by FeCl_3 This sulphonic acid yields oxy mesitylenic acid by potash fusion

Methyl derivative $\text{C}_6\text{H}_2(\text{CH}_3)_2(\text{OCH}_3)$ (c 202°) Liquid Bromine gives a bromo derivative [809] V AMIDO- and Bromo mesitol

MESITONIC ACID $\text{C}_6\text{H}_2\text{O}_2$ or $(\text{CH}_3)_2\text{C}(\text{CO}_2\text{H})\text{CH}_2\text{COCH}_3$ (?) *Di a methyl B acetyl propionic acid* [749'] (Anschutz) [909'] (P) (138° at 15 mm) (230° – 240°) at 760 mm Prepared, together with an acid ($\text{C}_6\text{H}_2\text{NO}_2$), and phoronic nitrile ($\text{C}_6\text{H}_2\text{N}_2\text{O}_2$), by boiling the product of the action of gaseous HCl on acetone, with alcoholic KCN In this reaction the substance first formed is probably the chloride $(\text{CH}_3)_2\text{CClCH}_2\text{COCH}_3$, which by KCN would give the nitrile of mesitonic acid (Pinner, *B* 14, 1071) Plates or prisms Sol water, alcohol, ether, and benzene, sl sol petroleum ether Its salts are v e sol water On distillation it splits off water, forming mesito lactone $\text{C}_6\text{H}_2\text{O}_2$ Yields di methyl malonic acid on oxidation with HNO_3 (Anschutz, *A* 247, 103) On reduction it yields the lactone of γ oxy di a methyl-valeric acid $(\text{CH}_3)_2\text{CCH}_2\text{CHMeOCO}$ [529']

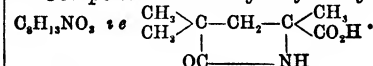
Ethyl ether $\text{C}_6\text{H}_2(\text{OH})(\text{CO}_2\text{Et})$ (210') Colourless liquid

Acetyl derivative of the ethyl ether $\text{C}_6\text{H}_2(\text{OAc})\text{COEt}$ (206') Liquid (Pinner, *B* 15, 578)

Phenyl hydrazide

$\text{PhHN}, \text{CMeCHCMe}, \text{CO}_2\text{H}$ [121 59] Small prisms (from benzene or dilute alcohol) On boiling with a few drops of HCl aq it yields crystalline $\text{C}_6\text{H}_2\text{N}_2\text{O}$ [849']

Compound with Hydrogen Cyanide



Mesitylic acid [1749'] Prepared, together with mesitonic acid and phoronic nitrile, by boiling the product of the action of HCl on acetone with alcoholic KCN In this reaction it is probably formed by the addition of HCN to mesitonic acid or its nitrile (Simpson, *A* 148, 351, Pinner, *B* 14, 1071, 15, 580) Distils undecomposed at a high temperature Large flat prisms (containing aq) Very stable body Forms easily soluble salts By heating with HCl to 140° CO_2 and NH_3 are split off Heated with concentrated H_2SO_4 to 150° it is converted

into mesitonic acid On oxidation with KMnO_4 in acid solution it gives di methyl succinimide and di-methyl malonic acid [107°]

Ethyl ether A^tEt [90°] Colourless prisms Sol alcohol and in acid, sl sol water

Amide $\text{C}_6\text{H}_7\text{N}_2\text{O}_2$ [222°] Colourless crystals Sol water and alcohol

Mesito lactone $\text{C}_8\text{H}_{10}\text{O}_2$ ψ e

$(\text{CH}_3)_2\text{C}(\text{CH}_3)\text{CH}_2\text{CO}$

Lactone of Oxy heptenone

acid [24°] (167°) Prepared by distillation of mesitonic acid (Pinner, *B* 15, 579) Large colourless prisms, sl sol water Neutral body By boiling with KOH it is reconverted into mesitonic acid It combines with bromine

MESITYL This name was given by Kane (*P* 44, 476) to a radicle C_6H_3 , of which he supposed acetone to be the hydroxide and MESITYL oxide (q v) the oxide More recently it has been employed to denote both s di methyl-benzyl and s tri methyl phenyl

MESITYL ALCOHOL ψ ω OXY MESITYLENE

MESITYL BROMIDE ψ BROMO MESITYLENE

MESITYLENE C_6H_6 ψ e $\text{C}_6\text{H}_5\text{Me}$, [135]

s-Tri-methyl benzene Mol w 120 (164.5°)

(Schiff, *A* 220, 94) S G $\frac{8}{8}$ 8694 (S), $\frac{2}{2}$ 8558 (Bruhl, *A* 200, 190) CE (9.8° to 161.5°) 001159 (S) VD 4.4 (obs and calc) SV 162.4 (Schiff), 162.2 (Rumery) $\mu_p = 1.5015$ $R_\infty = 65.75$ (B) $\text{H C} = 1,251,660$ [$\text{C}_6\text{O}_2 = 94,000$, $\text{H}_2\text{O} = 69,000$] (Stohmann, *J pr* [2] 85, 41), 1,282,310 (Thomsen, *Th* 4, 63) $\text{H F p} = 490$ $\text{H F v} = -2410$ (*Th*)

Occurs in coal tar oil (Fittig a Wackenroder, *A* 151, 292, Jacobsen, *A* 184, 179, *B* 9, 256, 10, 855) It is also one of the products of the manufacture of oil gas (Armstrong, *C J* 49, 74) Occurs in all kinds of petroleum (American, Russian, &c) (Engler, *B* 18, 2234)

Formation—1 By distilling acetone with H_2SO_4 (Kane, *P* 44, 474, Hofmann, *C J* 2, 101, Cahours, *C R* 24, 255, *C J* 3, 17)—2 By the action of H_2SO_4 on allylene (Fittig a Schroebe, *B* 8, 17)—3 By treating toluene with MgCl in presence of AlCl_3 , ψ cumene being also formed (Friedel a Crafts, *A Ch* [6] 1, 461, Ador a Rilliet, *B* 12, 329)

Preparation—1 By mixing 180 g acetone with 800 g sulphuric acid, allowing to stand for an hour, then distilling gently, using a current of steam at the end of the operation The yield is 40 g (Varenne, *Bl* [2] 40, 266)—2 H_2SO_4 (2 vols) diluted with water (1 vol) is slowly added to a mixture of crude acetone (2 vols) and sand After 24 hours the mixture is distilled The oil thus obtained is washed with aqueous NaOH and rectified (Fittig a Bruckner, *A* 147, 42)—3 The fraction of coal tar oil boiling between 160° and 168° is agitated with H_2SO_4 and the resulting sulphonic acids converted first into Ba salts and then into Na salts The Na salts are then treated with PCl_5 and the resulting chlorides converted into amides by NH_3Aq . On crystallising from alcohol, the ψ cumene sulphonamide separates first, and then mesitylene sulphonamide The mesitylene sulphonamide is then heated with conc HClAq at 160° (Jacobsen)—4 The sulphonic acids of ψ cumene and of mesitylene may also be separated by heating

with HClAq at 100° for 1 hour, when mesitylene sulphonic acid is decomposed into mesitylene and H_2SO_4 , while ψ cumene sulphonic acid is not affected (Armstrong, *B* 11, 1697)—5 When steam is passed through a heated solution of mesitylene sulphonic acid in dilute H_2SO_4 , hydrolysis into mesitylene and H_2SO_4 begins as soon as the temperature reaches 100° (Armstrong a Miller, *C J* 45, 148)

Properties—Light oil

Reactions—1 Br and Cl form crystalline substitution products—2 Fuming HNO_3 mixed with fuming H_2SO_4 forms tri nitro mesitylene, 3 Boiling dilute HNO_3 oxidises it to mesitylenic acid and uvitic acid (Fittig, *A* 141, 142)—4 $\text{K}_2\text{Cr}_2\text{O}_7$ and H_2SO_4 yield acetic acid (Fittig) 5 KMnO_4 oxidises it to uvitic and trimelic acids (Jacobsen, *A* 184, 191)—6 When heated with AlCl_3 in a current of HCl it yields MgCl and *m* xylene, toluene, and benzene (Jacobsen, *B* 18, 312)—7 MeI in presence of AlCl_3 yields (1, 2, 4, 5) tetra-methyl benzene (durene) (Barbier a Roux, *Bl* [3] 2, 4)—8 Benzyl chloride and AlCl_3 at 100° form benzyl mesitylene (q v) 9 Benzoyl chloride in presence of aluminium chloride forms phenyl tri methyl phenyl ketone $\text{C}_6\text{H}_5\text{COCH}_2\text{Me}$, [36°] and the compounds $(\text{C}_6\text{H}_5\text{CO})_2\text{CMe}$, [117°] and $(\text{C}_6\text{H}_5\text{CO})_3\text{CMe}$, [215°] (Louise, *A Ch* [6] 6, 200)—10 PCl_5 at 180° gives a small quantity of $\text{C}_6\text{H}_5(\text{CH}_2\text{Cl})$, (Colson a Gautier, *Bl* [2] 45, 6)—11 Mesitylene is very readily attacked by halogens In the dark, mono- di- or tri- (*eso*) bromo mesitylene ($\text{C}_6\text{H}_4\text{Me}_2\text{Br}$, $\text{C}_6\text{H}_3\text{Me}_2\text{Br}_2$, and $\text{C}_6\text{Me}_3\text{Br}_3$) are formed according as 1, 2, or 3 mols of bromine are employed Chlorine acts similarly The presence of iodine, though not required, does not effect the reaction At the boiling point the bromine, as usual, enters the side chains, 1 mol Br forming ω bromo mesitylene (mesityl bromide) $\text{C}_6(\text{CH}_3)_2\text{CH}_2\text{Br}$ melting at [38°] Direct sunshine, which usually acts like heat, in the case of mesitylene, owing to the great tendency to displacement of the nuclear hydrogen, produces a different result, (*eso*) mono bromo mesitylene is first produced but is partly attacked by the rest of the bromine forming the liquid p ω di-bromo mesitylene (p bromo mesityl bromide) $\text{C}_6\text{H}_4(\text{CH}_3)_2\text{Br}(\text{CH}_2\text{Br})$ [53.4.1] If *eso* bromo mesitylene be exposed to further bromination in sunshine, the reaction proceeds normally, producing p ω di bromo mesitylene (as above), and, on further action, p ω - α_2 tri bromo mesitylene $\text{C}_6\text{H}_3(\text{CH}_3)_2\text{Br}(\text{CH}_2\text{Br})_2$ [54.3.1] melting at [122°] (Schramm, *B* 19, 212)—12 When a 10 p solution of mesitylene in CS_2 is mixed with a similar solution of chromyl chloride CrO_2Cl_2 a pp ($\text{C}_6\text{H}_4\text{Me}_2$)(CrO_2Cl_2) is obtained which when decomposed by water yields di methyl benzoic aldehyde (221°), which is oxidised by air to mesitylenic acid (Etard, *C R* 97, 909)—13 The diacetyl derivative of *ortho formic aldehyde* (1 pt) mixed with mesitylene (1 pt) and HOAc (10 pts) condenses on adding a mixture of HOAc (10 pts) with an equal volume of H_2SO_4 , and after 24 hours crystals of $(\text{C}_6\text{H}_4\text{Me}_2)_2\text{CH}_2$ [130°] separate The formation of this body may be used as a test for mesitylene, a mixture of methyl alcohol and CrO_3 in HOAc being used instead of $\text{CH}_3(\text{OAc})_2$ (Baeyer, *B* 5, 1094, 6, 220)

Constitution.—The explanation of the for.

mation of mesitylene by condensation of acetone was first given by Baeyer (A 140, 306). That the three methyls are symmetrically situated in the benzene nucleus would appear from the existence of only one mono-derivative derived by displacing an atom of hydrogen in the nucleus, *eg* there is only one nitro-mesitylene. A systematic proof of the symmetry of mesitylene has been given by Ladenburg (A 179, 163). He prepares in succession $C_6Me_3(NO_2)(NO_2)H$, $C_6Me_3(NO_2)(NH_2)H$, $C_6Me_3(NO_2)(NH_2)(NO_2)$, and $C_6Me_3(NO_2)H(NO_2)$. Since the last di-nitro-mesitylene is identical with the first, the second and third hydrogen atoms are similarly situated in the nucleus. Again, from the above nitro-mesidine $C_6Me_3(NO_2)(NH_2)H$ Ladenburg obtained $C_6Me_3(NO_2)HH$ and, by reducing this, mesidine $C_6Me_3(NH_2)HH$. Nitro-mesidine, obtained by nitrating $C_6Me_3(NH_2)HH$, must be either $C_6Me_3(NH_2)(NO_2)H$ or $C_6Me_3(NH_2)H(NO_2)$. But since it has been shown that the second and third atoms of hydrogen are similarly situated, these two formulae are identical. And since the nitro-mesidine is found to be identical with the $C_6Me_3(NO_2)(NH_2)H$ prepared from the di-nitro-mesitylene $C_6Me_3(NO_2)(NO_2)H$, it follows that the first and second atoms of hydrogen are similarly situated. Hence all three atoms of hydrogen are similarly situated, and the formula will be $C_6Me_3H_3$ [1 3 5].

Mesitylene hexahydride C_6H_8 , *ie* $C_6Me_3H_3$, (*c* 137°). Obtained by heating mesitylene with PH_3 at 280° (Baeyer, Z [2] 5, 320, A 155, 273). Oil, smelling like petroleum. Converted by prolonged heating with fuming HNO_3 into tri-nitro-mesitylene.

References -- DI AMIDO, BENZYL, BROMO, BROMO NITRO, BROMO OXY, CHLORO, DI CHLORO BROMO, CHLORO NITRO, NITRO, NITRO BENZYL, DI OXY- and TRI OXY MESITYLENE. V also MESIDINE, MESITOL, and AZO COMPOUNDS.

MESITYLENE-DIAMINE *v* DIAMIDO MESITYLENE, vol 1 p 166.

MESITYLENE CARBOXYLIC ACID *v* (B)-ISO CUMINIC ACID.

MESITYLENE GLYCOL *v* DI OXY MESITYLENE.

MESITYLENE PHTHALOYLIC ACID *v* PHENYL TRI METHYL PHENYL KETONE CARBOXYLIC ACID.

MESITYLENE SULPHINIC ACID
 $C_6H_3Me_3SO_2H$ [1 3 5 6] [99°]. Formed by adding a mixture of sodium amalgam and $C_6H_3Me_3SO_2Cl$ to a dry mixture of benzene and toluene, and decomposing the resulting pulpy mass with HCl (Holtmeyer, Z 1867, 686). Fan-shaped groups of needles -- BaA' , aq -- AgA' .

MESITYLENE SULPHONIC ACID $C_6H_3Me_3SO_3H$ [*c* 77°]. Formed by dissolving mesitylene in warm H_2SO_4 (Hofmann, C J 2, 113, Jacobsen, A 146, 85, 184, 185, Beilstein a Kögler, A 137, 317, Fittig, J 1866, 610, Z [2] 4, 583). Trimetric six-sided tables (containing 2aq) (Rose, Z [2] 6, 341, A 164, 53, Bodewig, J 1879, 737). Decomposed by distillation with dilute H_2SO_4 into mesitylene and H_2SO_4 , the hydrolysis beginning at 100° (Armstrong a Miller, C J 45, 149). Gives mesitol on potash-fusion (Jacobsen, A 195, 265).

Salts -- KA' , aq . Roundish groups of silky

laminae S 14 at 13° -- NH_4A' , aq [250°]. Plates, *v* sol water and alcohol -- CaA' , $5aq$ efflorescent crystals -- BaA' , $9aq$ S 6 6 at 18°, 5 5 at 11 5° -- SrA' , $7aq$ -- MgA' , $6aq$ -- CoA' , $6aq$ flesh coloured laminae, *v* sol water and alcohol. -- CuA' , $4aq$ Pale green laminae S 6 at 10° -- PbA' , $9aq$ Pearly plates S 15 at 20°.

Chloride $C_6H_3Me_3SO_2Cl$ [57°]. Wedge-shaped tables (from ether), insol water, *v* sol alcohol and ether (Holtmeyer, Z 1867, 686).

Amide $C_6H_3Me_3SO_2NH_2$ [142°] S 033 at 0°, 54 at 100° S (83 p.c. alcohol) 5 5 at 0°, 114 at 78°. Fibrous mass (from alcohol) or hair like needles (from ether). May be oxidised to $C_6H_3Me_3(CO_2H)_2SO_2NH_2$ and an acid $C_6H_3Me_3(CO_2H)_2SO_2NH_2$, and the anhydride of

an isomeric acid $C_6H_3Me_3\langle\frac{SO_2}{CO}\rangle NH$ (Hall a.

Remsen, Am 2, 130, Emerson, Am 8, 268).

Imide $(C_6H_3Me_3SO_2)_2NH$ [124°]. Formed by heating the amide with a little HCl in a sealed tube. Long needles, *m* sol hot water, sol $NaOH$ aq , and reppd by HCl .

Mesitylene disulphonic acid $C_6HMe_3(SO_3H)_2$. Formed by dissolving mesitylene (1 pt) in fuming H_2SO_4 (10 pts) and adding P_2O_5 (3 pts), the temperature being kept low during the operation (Barth a Herzog, M 1, 807). Deliquescent needles. Gives oxy-mesitylenic acid on potash fusion -- K_2A'' , $2aq$ -- NaA'' , $1\frac{1}{2}aq$ needles -- BaA'' , $3aq$ needles -- CuA'' (dried over H_2SO_4) greenish white needles.

Reference -- BROMO-MESITYLENE SULPHONIC ACID.

MESITYLENIC ACID $C_6H_3O_2$, *ie*

$C_6H_3Me_3CO_2H$. Di-methyl benzoic acid. Mol w 150 [166°] $H C v$ 1,084,300 $H C p$ 1,085,200 $H F$ 105,800 (Stohmann, Kleber, a Langbein, J pr [2] 40, 135). Colourless plates.

Formation -- 1 By boiling mesitylene with dilute HNO_3 (1 vol of SG 14 and 2 vols water) for 18 hours. The product is distilled with steam, boiled with a little tin and HCl aq to remove a nitro compound, and crystallised from alcohol (Fittig, A 141, 144, Fittig a Brückner, Z [2] 4, 493, A 147, 45) -- 2 By boiling *s*-dimethyl ethyl-benzene with nitric acid (SG 1.1) (Jacobsen, B 7, 1430, Wroblewsky, B 9, 495). 3 By passing CO over a mixture of $NaOEt$ and $NaOAc$ at 205°, or by heating such a mixture with zinc dust (Geuthner a Fröhlich, A 202, 310).

Properties -- Monoclinic crystals (from alcohol), *v* sol water, *v* sol alcohol. Sublimes below its melting point. Distillation with lime yields *m*-xylene. Chromic acid mixture oxidises it to trimelic acid.

Salts -- NaA' (dried at 130°). *V* sol water and alcohol -- CaA' , $\frac{1}{2}aq$. Not more soluble in hot than cold water -- BaA' , silky prisms -- MgA' , $5aq$ groups of monoclinic prisms -- ZnA' , (dried at 130°) laminae or needles, *sl* sol water -- MnA' , $-NaA'$, (dried at 130°) -- AgA' , Minute needles, *sol* hot water -- AgA' , aq (Hall a Remsen, Am 2, 130).

Ethyl ether EtA' (241°). Heavy oil; solidifies below 0°.

Amide $C_6H_3Me_3CONH_2$ [133°]. Needles; *v* *sl* sol cold water.

References -- AMIDO- BROMO-, and CHLORO-MESITYLENIC ACID.

MESITYLENIC GLYCOL *v* Di oxy mesityl

ENE

MESITYLENIC GLYCERIN *v* Tri-oxy me

SITYLENE

MESITYLIC ACID *v* compound of Mesitonic acid with HO₂**MESITYL MERCAPTAN** *v* Tri-methyl-

PHENYL MERCAPTAN

MESITYL OXIDE C₆H₃O₂(CH₃)₂C CH CO CH₃, *Methyl isobutenyl ketone*
Isopropylidene acetone (132°) at 751 mm
S G $\frac{4}{25}$ 8578 μ , 1.4440 V D 3.67 (calc 3.39)
R_∞ 49.7 (Kanonnikoff, *J pr* [2] 31, 352)

Formation—1 One of the products of the action of H₂SO₄ on acetone (Kane, *P* 44, 476)
2 Together with phorone by leaving acetone for several weeks in contact with quicklime (Fittig, *A* 110, 32)—3 One of the products of the action of ZnEt₂ or ZnMe₂ on acetone (Pawlow, *B* 9, 1311, *A* 188, 130)—4 In small quantity, together with other bases, by distilling diacetone amine (Heintz, *A* 174, 133, 181, 70, *B* 8, 89)—5 By boiling with lime the lactonic products of condensation of acetoacetic ether (Hantsch, *A* 222, 21)—6 By boiling acetyl chloride with acetone (Beilstein a Wiegand, *Bl* [2] 38, 167)—7 Among the products of the distillation of glycerin with BaO (Destrem, *A Ch* [5] 27, 5)

Preparation—Acetone is saturated with HCl and then left to itself for a fortnight. The product is mixed with water, and the heavy oil, containing hydrochlorides of mesityl oxide and phorone, decomposed by alcoholic or conc aqueous KOH, and distilled with steam. The product is fractionally distilled (Baeyer, *A* 140, 297)

Properties—Oil, smelling like peppermint, insol water, miscible with alcohol and ether

Reactions—1 Boiling dilute sulphuric acid forms acetone. Conc H₂SO₄ gives mesitylene (Claisen, *B* 7, 1168)—2 Boiling dilute nitric acid yields acetic and oxalic acids—3 KMnO₄ oxidises it to acetic and oxy isobutyric acid (Pinner, *B* 15, 591)—4 Slowly combines with NaHSO₃ forming NaC₆H₃SO₃aq, the sodium salt of methyl isobutyl ketone sulphonic acid CH₃COCH₂OMe, SO₃H, which is decomposed by conc NaOH aq regenerating mesityl oxide (Pinner, *B* 16, 1727)—5 Bromine acts violently. But when it is added to a solution of mesityl oxide in CS₂, only C₆H₃Br₂O is formed. This can be distilled with steam, but in the dry state soon gives off HBr—6 A solution in dilute alcohol is strongly attacked by sodium amalgam and water then ppts a heavy oil C₆H₃O 'mesitic ether' or 'mesitic camphor' (c 215°) smelling like camphor. A small quantity of a solid [110°-120°] is also formed (Claisen, *A* 180, 7)—7 PCl₅ forms 'mesityl chloride' C₆H₃Cl₂, which separates on adding water as a heavy oil, smelling like oil of turpentine. It resinifies on exposure to air, and is decomposed by distillation over KOH, baryta, or lime into HCl and C₆H₃Cl, a liquid (130°) smelling like turpentine—8 Anmonia forms diacetoneamine C₆H₃NO—9 Hydriodic acid forms a heavy oil C₆H₃IO decomposed by alcoholic KOH into HI and mesitylene (Pawlow, *A* 188, 138)—10 HCl passed into a mixture of mesitylene and benzoic aldehyde forms C₆H₃COOHCHPh (Claisen a

Claparède, *B* 14, 849)—11 With acetamide and ZnCl₂ it forms oxy-tri methyl pyridine dihydride (Canzoneri a Spica, *G* 14, 349, *B* 19, 818, cf Engler a Riehm, *B* 19, 40)—12 With amyl nitrite and alcoholic NaOEt it forms the nitroso-derivative CMe, CH CO CH NOH, crystallising in colourless prisms [102°] (Claisen a Manasse, *B* 22, 529)

Oxim C₆H₃NO (c 185°) Formed by the action of hydroxylamine on mesityl oxide (Nageli, *B* 16, 495) Oil. Sol alcohol, ether, benzene, CS₂, ligroin, alkalis and acids. On warming with acids it is resolved into its constituents

MESOCALPHORIC ACID *v* CAMPHORIC ACID**MESORCIN** *v* TRI-METHYL RESORCIN**MESOTARTARIC ACID** *v* TARTARIC ACID**MESOXALIC ACID** C₆H₃O₄aq

CO₂H CO CO₂H + aq or CO₂H C(OH), CO₂H (Petrieff, *B* 11, 414) Mol w 136 [115°] (Deichsel), [108°] (Böttger, *A* 203, 140), [120°] (Fischer)

Formation—1 Together with urea, by boiling alloxan or alloxanic acid with baryta. The resulting Ba salt is recrystallised and decomposed by H₂SO₄ (Liebig a Wohler, *A* 26, 298, Svanberg, *B J* 27, 165, Deichsel, *B* 1864, 587)—2 By treating amido malonic acid with iodine and water (Baeyer, *A* 131, 298)—3 By boiling di bromo malonamide CBr₂(CO NH₂), with precipitated Ag₂O (Freund, *B* 17, 782)—4 By boiling di bromo malonic acid with baryta water (Petrieff, *J R* 10, 72)—5 By heating caustic acid with concentrated aqueous lead subacetate (E Fischer, *A* 215, 283)

Properties—Very deliquescent prisms, melting without loss of aq. M sol alcohol and ether, v e sol water. Its concentrated aqueous solution decomposes above 80° into CO₂, glycolic, oxalic, and glyoxylic acid (Böttger, *A* 203, 138). With acetates of Ba and Pb it gives flocculent pps gradually becoming crystalline. In neutral solutions CaCl₂ and BaCl₂ give pps. It reduces hot ammoniacal AgNO₃.

Reactions—1 H₂S passed into an aqueous solution to which silver oxide has been added forms thio glycolic acid CH₂(SH)CO₂H and thio di glycolic acid (sulphido diacetic acid) S(CH₂CO₂H)₂ (Böttger)—2 Sodium amalgam acting on its dilute aqueous solution forms tartaric acid CO₂HCH(OH)CO₂H—3 Silver mesoxalate boiled with water yields CO₂, oxalic acid, and silver—4 Urea at 100° gives allantoin C₄H₆N₂O₃—5 Hydroxylamine yields the oxim which is identical with nitroso malonic acid

Salts—[A⁻ = C₆O₄]—(NH₄)₂A⁻ granular crystals turning red in air (Deichsel, cf Engel, *G R* 98, 628)—(NH₄)₂A⁻ aq. Obtained by evaporation *in vacuo* over H₂SO₄.—Needles (Petrieff)—Na₂A⁻ aq thin laminae, v sol water.—CaA⁻ aq white pp insol dilute acetic acid.—CaA⁻ 4aq (dried at 100°) (Petrieff)—BaA⁻ aq (dried at 110°). White crystalline powder, v sl sol water.—BaA⁻ 1/3 aq.—BaA⁻ (dried at 180°)—(HO Pb)₂A⁻ nearly insol water.—Ag₂A⁻ aq amorphous pp changing to minute needles. Explodes when heated

Ethyl ether Et₂A⁻ aq. From the silversalt and EtL. Oil.

Di acetyl derivative of the ethyl ether $(\text{CO}_2\text{Et})_2\text{C}(\text{OAc})_2$ [145°] From the ether (1 mol) and AcCl (2 mols) at 100° (P) Long needles Partially decomposed by fusion and by solution in water

Di acetyl derivative $\text{C}(\text{OAc})_2(\text{CO}_2\text{H})_2$ [180°] From $(\text{CO}_2\text{Et})_2\text{C}(\text{OAc})_2$ and dilute alcoholic KOH Needles Its alkaline salts are v sol water, its silver salt $\text{Ag}_2\text{C}_4\text{H}_4\text{O}_8$ is an insoluble powder

Amide $\text{C}(\text{OH})_2(\text{CONH}_2)_2$ Deliquescent tables, which turn red in the air

Phenyl-hydrazide

$\text{PhHN}_2\text{C}(\text{CO}_2\text{H})_2$ [158°-164°] From mesoxalic acid and phenyl hydrazine hydrochloride in dilute acid solution (Elbers, A 227, 355) Crystals

Phenyl hydrazide of the Nitrile

$\text{PhHN}_2\text{C}(\text{CN})_2$ [185°] May be formed from the oxim phenyl hydrazide of glyoxylyl cyanide $\text{HO N CH C}(\text{N}_2\text{HPh})\text{CN}$ by dissolving in POCl_3 , heating with PCl_5 , pouring into ice, and treating the crystals with ether and alkali (Von Pechmann & Wehsarg, B 21, 3001) Yellow needles (from ether ligroin) Turns brown at 126° Sol hot water, alkalis, and the usual menstrua Conc H_2SO_4 forms a blood red solution not changed by FeCl_3 On warming with conc HClAq it forms yellow needles of $\text{C}_8\text{H}_5\text{N}_3\text{O}$ [215°]

MESOXALIC ALDEHYDE CHO CO CHO

Oxim HON CH CO CH NOH or

$\text{ON CH CO CH}_2\text{NO v Di NITROSO ACETONE}$

Mesoxalic semi aldehyde $\text{*CHO CO CO}_2\text{H}$ *Glyoxylyl carboxylic acid*

Diphenyl hydrazide

$\text{CH}(\text{N}_2\text{HPh})\text{C}(\text{N}_2\text{HPh})\text{CO}_2\text{H}$ [203°] From di bromo pyruvic acid in aqueous solution and phenyl hydrazine hydrochloride (Nastvogel, A 248, 87) Reddish yellow needles (from hot alcohol), almost insol water, sl ether and chloroform, v sol hot alcohol, acetone, benzene and HOAc Dyes wool and silk yellow Cold conc H_2SO_4 forms a dark red solution from which it is ppt by water unaltered

Di p tolyl hydrazide

$\text{CH}(\text{N}_2\text{HC}_6\text{H}_4\text{Me})\text{C}(\text{N}_2\text{HC}_6\text{H}_4\text{Me})\text{CO}_2\text{H}$ [188°] Formed in like manner Golden needles (from benzene) Its alkaline salts are v sol hot, sl sol cold, water

Nitrile $\text{*CHO CO CN v Glyoxylyl cyanide}$

Mesoxalic bromo semi aldehyde

$\text{*CBrO CO CO}_2\text{H}$

Methyl ether of the oxim

$\text{CBrO C}(\text{NOH})\text{CO}_2\text{Me}$ [c 170°] From di methyl di bromo pyrrole di carboxylate and HNO_3 at -18° (Ciamocian & Silber, B 20, 2601) Crystalline, sol alcohol, m sol cold water, insol petroleum ether Decomposed by fusion and by boiling with water

MESOXALYL-UREA is ALLOXAN See also its compounds with METHYL ANILINE, NAPHTHYL AMINE, and PYRROLES

META Use of this prefix applied to inorganic compounds, for *Meta-acids* and *Meta salts* v the acids or salts to the name of which *Meta-* is prefixed Thus *Meta-phosphoric acid* will be found under **PHOSPHORIC ACIDS**, and *Meta stannates* under **stannates**, a subdivision of the article **TIN**.

Names of organic bodies beginning with 'meta' will be found under the word to which 'meta' is prefixed

METACETONE The mixture obtained by distilling sugar with quicklime, called metacetone by Fremy (A Ch [2] 59, 6), is composed according to E Fischer & Laycock (B 22, 101) of propionic aldehyde, di methyl-furfurane, and hydrocarbons

METACETONIC ACID An old name for **PROPIONIC ACID**

METALBUMIN v PROTEIDS, Appendix C

METALDEHYDE v ALDEHYDE

METALLIC ACIDS In the article **ACIDS** (vol 1 p 47, cf **CLASSIFICATION**, vol 11 pp 201, 202), it is shown that compounds of H with certain negative elements or groups of elements react with metallic oxides, hydroxides, and carbonates, in presence of water, to produce substances composed of metal, and the elements of the hydrogen compound excepting the H or a part of the H Such hydrogen compounds are called acids The negative elements which are found intimately combined with H in acids are fluorine, chlorine, bromine, iodine, oxygen, sulphur, selenium, tellurium, carbon always enters into the composition of those negative groups of elements which combine with H or with H and other elements to form acids Only a few acids are binary compounds, the greater number are compounds of H with two or three other elements, among which are always found at least one of the eight negative elements enumerated above, or at least one of the negative carbon containing groups of elements By far the greater number of the more stable and definite acids are composed of H combined with non metallic elements, but some acids have been isolated which are composed of H combined with metals and one or more of the eight strongly negative elements already enumerated, or one or more of the carbon containing negative groups of elements The following table gives the composition of most of those acids which contain metallic elements —

Metallic acids

H_2AsO_4	(H_2TiO_4)	$\text{H}_2\text{PtCy}_4\text{Cl}_4$
H_2AsO_3	(H_2TiO_3)	H_2PtBr_4
$\text{H}_2\text{As}_2\text{O}_7$	H_2SnO_3	$\text{H}_2\text{Pt}_2\text{S}_4$
	H_2SnO_2	H_2OsCy_4
(H_2AsS_2)	H_2SnS_3	H_2RuCy_4
H_2SbO_4	(H_2PbO_4)	H_2IrCy_4
HSbO_3	H PbI_3	$\text{H}_2\text{N}_2\text{Os}_2\text{O}_6$
$\text{H}_2\text{Sb}_2\text{O}_7$	H_2CrO_3	
H_2SbO_3	$(\text{H}_2\text{Cr}(\text{SCy})_3)$	
HVO_3	H_2MoO_4	
$(\text{H}_2\text{V}_2\text{O}_7)$	H_2WO_4	
$\text{H}_2\text{V}_2\text{O}_5$	H_2UO_4	
$\text{H}_2\text{Ta}_2\text{O}_7$	$(\text{H}_2\text{M}_2\text{O}_4, \text{M} = \text{Mo, W, U})$	
$(\text{H}_2\text{Nb}_2\text{O}_7)$	$(\text{H}_2\text{M}_2\text{O}_{10}, \text{M} = \text{Mo, W, U})$	
(H_2MnO_4)	HAuCy_4	(H_2ZnO_4)
H_2MnCl_4	HAuCl_4	HHgCl_4
H_2MnCy_4	HAuBr_4	HHgBr_4
H_2FeCy_4	H_2PtCl_4	HHgI_4
H_2FeCy_3	H_2PtCl_3	H_2HgCl_4
$\text{H}_2\text{FeCy}_2\text{NO}$	H_2PtI_4	H_2HgBr_4
H_2CoCy_4	$\text{H}_2\text{Pt}(\text{NO})_4$	H_2HgI_4
H_2CoCy_3	$\text{H}_2\text{Pt}(\text{NO})_3\text{Cl}_2$	$(\text{H}_2\text{ZnCl}_4)$
(H_2AlO_4)	$\text{H}_2\text{Pt}(\text{SCy})_4$	$(\text{H}_2\text{Zn}_2\text{Cl}_4)$
	H_2PtCy_4	

The isolation of some of the acids in the fore-

going list is doubtful, these acids are placed in brackets. The compounds H_2AlO_3 and H_2ZnO_3 are also bracketed, because the reactions of these bodies show that they may be classed as feeble acids and at the same time as basic hydroxides.

An examination of the composition of the metallic acids shows that the reacting atomic aggregates of these compounds all contain a number of atoms of a negative element, or group of elements, which is larger relatively to the number of atoms of metal present. The accumulation of negative atoms so modifies the functions of the H atoms that these are replaceable by metals, and this notwithstanding the presence of the positive metallic atoms. The H of the metallic hydrogen-containing compounds H_2CrO_4 ($= Cr_2O_3, H_2O$) and H_2CrO_5 ($= CrO, H_2O$) is not replaced by metals when these compounds react with metallic hydroxides or carbonates, but when 4 atoms of the negative O are associated with one atom of Cr and 2 atoms of H, the H of this compound (H_2CrO_4) is distinctly acidic. The negative character, or acid forming character, of the groups CN and SCN is rendered evident by the number of acids which are formed by the union of these groups with H and metals.

If attention is paid to the general chemical characters of the metals which form acids it is seen that most of these metals occur in groups (using this term as it is used in the nomenclature of the periodic law) which also contain several distinctly non-metallic elements: thus Ti, Sn, and Pb belong to Group IV, which group contains C and Si, Cr, Mo, W, and U belong to Group VI, in which group also occur the negative elements O, S, Se, and Te, As, Sb, V, Nb, and Ta form part of Group V, which is distinctly negative in its general chemical character, and includes the markedly non-metallic elements N and P, I, e, Co, Ni, Au, and the Pt metals occupy a peculiar position in Group VIII (*cf* CLASSIFICATION, vol. i pp 203-210, also IRON ELEMENTS, this vol p 65).

Several salts exist which, on account of their methods of formation and general stability, are probably best regarded as derivatives of metallic acids that have not yet been isolated, among such salts may be mentioned the stannous and zirconic fluorides $M_2Sn(Zr)F_6$. Some chemists would class most, if not all, the double metallic haloid compounds as salts of metallic acids, *e.g.* $BiF_3, 3KF$ as the K salt of the acid H_3BiF_6 , $ZnCl_2, BaCl_2$ as the Ba salt of the acid H_2ZnCl_4 , and MgI_2, KI as the K salt of the acid $HMgI_2$ (*v.* especially Remsen, *Am* 11, No 5).

The consideration of those metal-containing compounds which are acids brings out the inadequacy of that classification which would divide the elements into two classes only, metals and non-metals, it also well illustrates the difficulties of chemical classification, as shown by the way in which the chemical properties of an element are modified according to both the nature and the number of other elementary atoms with which that element is combined (*cf* the article METALS in this vol and CLASSIFICATION in vol. i).

M M P M

METALLOIDS This name was at one time applied, most mistakenly, to the non-metallic elements. It is sometimes used to denote those elements which on the whole are non-metallic,

but yet closely approach the metals in some of their properties, As, Sb, Ti, V, Nb, Ta, for instance, are sometimes called metalloids. The term cannot be defined. There are certain elements which one chemist would class among metals, another would place with the non-metals, and a third would prefer to put into neither class, but call them metalloids.

M M P M

METALLURGICAL CHEMISTRY The chemical reactions utilised in metallurgy are divisible into two distinct classes, viz 'wet,' those which take place in aqueous solutions, and 'dry,' those which take place in furnaces, or their equivalent, at a relatively high temperature. The present tendency is more and more in the direction of combining these two methods, metals being now frequently extracted from their ores partly by wet processes and partly by dry processes.

The ore of any metal may be defined as a collection of mineral substances containing that particular metal in sufficient quantity to pay for its extraction on a commercial scale. Although each metallic element exists in nature in a great number of combinations, yet very few of these compounds occur in sufficient quantity to be of direct importance to the metallurgist, except in so far as they may tend to introduce impurities into the metal to be extracted.

Chemically, ores may be broadly divided into three classes, containing respectively—I NATIVE METALS, *i.e.* metals uncombined with any non-metallic element II SULPHIDES and ARSENIDES III OXIDES, *including carbonates and silicates*.

Class I NATIVE METALS A native metal may be separated from its ores in one of four ways (a) By LIQUATION. In order that this may be done it is essential that the metal be fusible at a temperature insufficient to cause the earthy portions of the ore to frit or agglomerate, *e.g.* Bi. (b) By fusing the ore, when the metal will sink to the bottom by reason of its high SG., *e.g.* Bi, Cu. (c) By dissolving the metal out by means of another metal, *e.g.* Au, Ag, Pt. These metals may be dissolved out of their ores by Pb, which is then removed by cupellation, or, in the case of Ag, the separation of the Ag from the Pb may be effected by Zn, which does not alloy with Pb, and being specifically lighter rises to the surface, carrying the Ag and some lead with it. The Zn is subsequently distilled off, and the residual Ag and Pb cupelled. Similarly, Au and Ag may be dissolved out by Hg, which may then be distilled off. (d) By dissolving the metal by means of an acid or a gas in solution, *e.g.* Au extracted by a solution of Cl, and Pt, Pd, Ir, Rh, and Ru by *aqueous regia*. The Au and Pt may be refined by wet processes, taking advantage of the fact that Au and Pt are insoluble in sulphuric, hydrochloric, and nitric acid, when these acids are used separately, but are soluble in *aqueous regia*, while the Ag, Cu, Pb, and Fe are freely soluble in one or other of the three acids named. In 'parting' Au, sufficient Ag must be present to allow the acid free access to the impurities, which would otherwise be protected by the insoluble gold. Pt, when present in Au only in small quantity, may be separated by 'parting' with Ag, as under these conditions the Pt is

acted on by the 'parting acid' If present in larger quantity, it must be ppd from a solution. Ir may be separated from Au by difference of SG, and from Pt it may be removed to a certain extent by using *aqua regia* of medium concentration, in which it is not freely soluble. Pt is ppd from its solutions as a double salt of Pt and NH₃, by adding NH₄Cl. Pd, if present, may be ppd before the Pt by neutralising with Na₂CO₃Aq and adding HgCy₂. Ir, if it has gone into solution, will not be ppd with the Pt, but its double salt is difficult to wash out. Pd, Rh, Ru, and Os will also be found with the Ir in solution after the Pt has been ppd. Rh may be removed completely by fusing the platinum double chloride with KHSO₄, and a small quantity of NH₄H₂SO₄. Ir may be ppd at the same time as platinum by KCl, and the pp fused with K₂CO₃, which will oxidise the Ir and not the Pt. Remove the potassium salt by boiling water, and then dissolve out the platinum with *aqua regia*, in which the oxide of Ir is insoluble. Ir may also be separated from the pp by KCy, the Ir salt being soluble while the Pt salt is insoluble. Pt is obtained in the metallic state by carefully heating the double chloride, which then breaks up. Au is ppd from its solutions as metal by FeSO₄Aq, SO₂Aq, or H₂C₂O₄Aq. For dental purposes, Au is frequently deposited by electrical means. Ag is first thrown down as chloride, which is afterwards reduced by Cu, Zn, or Fe. Au, containing not more than 10 p.c. Ag, is also refined by Miller's process, at the Australian Mint, in the dry way, by passing Cl into the molten gold. The impurities As, Sb, Bi, Pb, and Zn are converted into chlorides, which volatilise, and the Ag becomes AgCl, which forms a fused layer on the surface of the gold.

Class II. SULPHIDES AND ARSENIDES. Dry methods. Sulphides and arsenides are either (a) *infusible*, at such temperatures as can be obtained in furnaces on the large scale, (b) *fusible*, or (c) *volatile without fusion*.

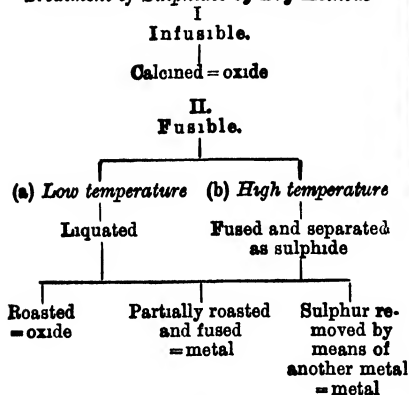
(a) *Infusible sulphides*. In these cases the S must be replaced by O, as an infusible sulphide cannot be properly reduced to the metallic state. This is done by calcining or roasting the ore, so that air has free access to it. The sulphide is oxidised to a sulphate at low temperatures, and at higher temperatures the sulphate breaks up into SO₂ and an oxide of the metal. Practically there is only one sulphide under this head, viz ZnS, *sinc blende*. For the subsequent treatment of the oxide see 'oxide class'.

(b) *Fusible sulphides*. Sulphides and arsenides of this class may be subdivided as follows—(i) *those which are fusible at a very low temperature, insufficient to produce fritting, & incipient fusion causing agglomeration of the constituents of the ore*, (ii) *those requiring a higher temperature, at which fritting would take place*. A sulphide in division (i) may be liquated out, e.g. Sb₂S₃. The sulphides and arsenides belonging to (ii) may be separated by fusing the ore, when the sulphide or arsenide would collect together beneath the slag, e.g. sulphide of copper (*copper pyrites*), arsenides of nickel and cobalt (if sufficient arsenic is not present in the ore more is added), the arsenide separates in a distinct layer from the sulphides of other metals during the

fusion, sulphide of nickel obtained by fusing nickel ores, or products, free from arsenic, with iron pyrites. The sulphides and arsenides thus separated from the gangue would next be treated in one of the following ways—1 *Converted into oxide by roasting*, e.g. Sb₂S₃, *copper matte*, arsenides of nickel and cobalt (the arsenious acid being condensed in coke towers), sulphides of nickel and cobalt free from arsenic, the oxides of nickel and cobalt are subsequently treated in the wet way. 2 *Partially roasted to form a certain amount of oxide and sulphate, and then fused*, the oxygen of the oxides combines with the sulphur of the sulphides and arsenides, forming SO₂, and liberating the metal, e.g. Sb₂S₃ and PbS. In the case of the double sulphide of Cu and Fe, the Fe is first removed by a series of calcinations and fusions, S passing to the Cu and O to the Fe, the oxide of iron thus formed uniting at the same time with silica to form slag. This process goes on so long as any iron remains. As soon as the iron is all removed, the reaction between Cu₂S, CuO, and Cu₂O takes place, liberating metallic copper. The principal impurities in copper ores likely to pass into the Cu are As, Sb, Zn, Pb, Bi, Sn, Ni, Co, Au, and Ag. The greater proportion of these present either volatilises or becomes oxidised and removed in the slags. Au and Ag, being neither appreciably volatile under the conditions nor oxidisable, become concentrated in the copper. It is particularly difficult to get rid of the last traces of As and Bi. The use of a basic lining to the furnace—say, dolomite—greatly facilitates the removal of As in the slag. The elimination of As is also assisted by the use of 'soda nitre' in refining. Bi can most readily be removed by what is known as the *best selecting* process, in which advantage is taken of the circumstance that copper has a greater affinity than bismuth for sulphur. This process comes in just before the copper is first reduced from the sulphide. A little copper is made to separate by the reaction between sulphide and oxide, this throws out and collects as 'bottoms' the bismuth, tin, lead, and antimony—3 *Fused in the presence of another metal which combines with, and so removes, the sulphur*, e.g. sulphides of Bi, Pb, and Sb treated with Fe, Ag separated by metallic Pb from sulphide of Pb containing sulphide of Ag. Copper can only be partially separated from sulphur in this way, a double sulphide forming which cannot be reduced by iron.

The operations described under (i) and (ii) are in some cases applied directly to the ore without first separating the sulphide by liquation or fusion.

(c) *Sulphides volatile without fusion*. Amongst the metallic sulphides there is only one which sublimes without fusion, viz HgS. When heated in presence of air HgS yields SO₂ and Hg. As the Hg has no tendency to combine with oxygen under these conditions, and is volatile at a very low temperature, it distils over, and may readily be condensed. For these reasons advantage is not taken of the fact that the sulphide is itself volatile, it being simpler to distil the metallic mercury direct from the ore. The sulphur is sometimes removed by roasting the mercury ore with lime or oxide of iron.

Treatment of Sulphides by Dry Methods**III.**

Sublimed without fusion

- 1 Sulphur separated by calcination alone = metal
- 2 Sulphur separated by lime, oxide of iron, or iron = metal

Wet processes for sulphides and arsenides. The sulphide and arsenide ores of Ni, Cu, and Co are partially, and those of Cu and Ag partially or wholly, treated by wet methods. In these processes the metal is first made to form a soluble compound. This may be done by roasting the ore with, or in some cases only mixing it with, some compound, which by double decomposition will convert the metal to be extracted into a soluble compound. In other cases it is done by roasting the ore alone at a low temperature, or even by simply exposing it to the action of the atmosphere without using artificially produced heat, in either case the sulphide will be converted into a sulphate, *e.g.* copper pyrites and silver ores.

Thus, soluble sulphate of copper may be formed by roasting copper pyrites or more slowly at the ordinary temperature of the atmosphere. About one third of the copper extracted in this country is obtained from Spanish pyrites, averaging only 3-4 p.c. of copper, by roasting the burnt ore—received from the sulphuric acid works—with common salt, which reacts with the CuSO_4 produced during the burning, to form CuCl_2 , which is then dissolved out, together with the chlorides of Ag, Pb, and Au, these metals being present to a small extent. The Ag is thrown down, along with Au and Pb, by a soluble iodide, KI or ZnI_2 . The iodine is removed from the AgI by Zn . Finally the Cu is pptd by addition of iron.

The reactions which take place in the wet treatment of silver ores are of special interest and importance. They are divisible into three classes.

I *The insoluble silver compound is decomposed and the Ag amalgamated while still in the ore ('free milling ore'), without the aid of any furnace-operation whatever.* If there is much S or As present the ore, where practicable, is first roasted with salt. Under this head are included all the amalgamation-processes.

II. *The insoluble silver compound is converted*

into sulphate by roasting alone and washed out with water, or the Ag is converted into chloride by roasting with common salt, and washed out by a solvent for chloride of silver. Under this head are included the Augustin, Ziervogel, Von Patera, and Russell processes.

III *The insoluble silver compound is converted into a soluble compound by the action of certain salts in solution without roasting the ore or employing any furnace-operation whatever.* This class includes the Von Patera process when applied to 'amalgamation tailings,' and the Russell process applied to both ores and tailings.

The amalgamation methods without roasting have been of great service where fuel is scarce, as in Mexico. These methods depend on the power of certain salts, such as the chlorides of Cu, to decompose Ag_2S . If mercury is present at the same time, the AgCl will be reduced and the Ag taken up by the excess of Hg, from which it can be separated by squeezing and distillation. Iron or Cu may be employed to decompose the AgCl , Hg being used only to collect the Ag. This reduces the loss of Hg, but to bring about sufficient contact, more power, and consequently more fuel, is required. Where fuel is available there is often great advantage in roasting the ore with salt before amalgamation.

Where possible the amalgamation methods have been superseded by the other wet processes included under heads II and III, viz the Augustin, Ziervogel, Von Patera, and Russell methods.

The Augustin process. Ore or regulus is first partially roasted, and then undergoes further roasting with common salt, the AgCl thus formed being washed out with NaClAq and pptd by Cu. The Ag is usually first concentrated in copper mattes.

The Ziervogel process. This method depends on the difference between the temperatures required to break up the sulphates of different metals. Thus by suitably regulating the temperature, the sulphates of Fe, Cu, &c., formed by roasting mattes at a low temperature, may be decomposed into oxides and SO_2 , while the sulphate of Ag will be unaltered, and can therefore be extracted by warm water, and then pptd as in the previous method. The regulation of the temperature throughout a large furnace is obviously a point requiring considerable skill. For this reason, in practice, mattes are usually first worked by the Ziervogel process, and the residue extracted by the Augustin method.

Von Patera process. The soluble salts produced by roasting the matte are first dissolved out with water, after which the matte is further roasted with NaCl , and the AgCl washed out with $\text{Na}_2\text{S}_2\text{O}_3\text{Aq}$ or $\text{CaS}_2\text{O}_3\text{Aq}$, from which solution the Ag is pptd by a soluble sulphide or SH_2 . The silver is reduced from the Ag_2S , either by roasting or by boiling with freshly slaked lime, forming calcium polysulphide. When there is much of the base metals present, more particularly lead, the Von Patera process is not so suitable. The lead is mostly present as sulphate after roasting, and this is soluble in $\text{Na}_2\text{S}_2\text{O}_3\text{Aq}$. Some of the most important objections to the Von Patera process, as applied to poor ores containing base metal, are obviated in the process next described.

The Russell process In this process what is known as the 'extra solution' is used, in addition to or in substitution for that ordinarily employed in the Von Patara process. The 'extra solution' is prepared by making solutions of $\text{Na}_2\text{S}_2\text{O}_3$ and CuSO_4 , containing respectively 18 parts of the former and 10 parts of the latter salt, and mixing them together. The pp after washing is dissolved in a solution containing 1-2½ p.c. $\text{Na}_2\text{S}_2\text{O}_3$. Although this 'extra solution' is not so good a solvent for AgCl as $\text{Na}_2\text{S}_2\text{O}_3\text{Aq}$, yet it acts energetically on native silver and the compounds of silver with S, As, and Sb. Owing to the want of permanence special precautions have to be taken in using this solution. To get the best effect the solutions should be used warm. In order to obtain finer bullion by this process, advantage is taken of the fact that PbCO_3 is not soluble in the solution, although other salts of lead are soluble. Na_2CO_3 , free from NaOH and Na_2S , is added to the solution containing the silver and lead which have been washed out. It is stated that the whole of the Pb may thus be removed. Carbonate and sulphate of Cu are the only compounds of Cu likely to occur which are soluble in the 'extra solution'.

It is probable that this process in the near future will supersede very largely the fusion and amalgamation processes, besides which it is applicable to ores which cannot profitably be treated by either of the other methods. The cost of the necessary chemicals is much less than that due to loss of mercury, in addition to which, lead and copper are lost in the older processes and saved in the Russell process.

In some cases the ores may be treated direct by the Russell process without previous roasting, and both the Von Patara and Russell methods are largely used in the treatment of 'tailings' from amalgamation.

Nickel and cobalt It has been seen that these metals may be separated from others as arsenide. In practice, however, the separation is not so complete, some of the nickel and cobalt passing into the regulus, and some of the other metals into the speise. The following are the reactions usually employed to separate the various metals from one another. The calcined speise is treated with HClAq . The resulting solution, which will contain Ni, Co, Fe, Cu, Pb, Bi, and As, is diluted, and oxidised by bleaching-powder, the proportion added being adjusted by a rough analysis. Milk of lime is added to the requisite extent to throw the iron down as sesquioxide, any arsenic present at the same time ppg as basic arsenate of iron. The solution is next treated with SH , to ppt Cu, Pb, and Bi, after which the Co can be thrown down as sesquioxide by a further addition of bleaching-powder, and subsequently the Ni ppt as hydrated oxide by the addition of milk of lime.

Class III. OXIDES, INCLUDING CARBONATES AND SILICATES Ores of this class are reduced to the metallic state by means of carbonaceous matter such as charcoal, coal, or co's, or by means of the gaseous product of the incomplete oxidation of carbonaceous matter, viz carbon monoxide.

The only exceptions among the oxides of the commoner metals are Al_2O_3 and MgO . These oxides cannot be reduced to the metallic state in this way; they have to be converted into

double chlorides, from which, while in a state of fusion, the metals are liberated by metallic sodium, or by electrolysis. The metals, the oxides of which are in practice reduced by carbon or CO, are Sn, Fe, Ni, Co, Mn, Cr, W, and Zn. Of these Sn and Zn present the simplest reactions. The reactions in the other cases are complicated by the fact that all these metals combine more or less freely with carbon. When it is desired to obtain the metals in the most malleable condition, + e free from carbon, the fact that the oxides and the carburised metals react with one another eliminating both the carbon and the oxygen, as CO and CO_2 , is utilised. Every iron- and steel making process is dependent on this reaction. According as the conditions are made more or less favourable to carburisation, or decarburisation, so will the resultant metal be either cast iron, hard steel, mild steel, or malleable iron. It is possible to reduce the oxides under consideration to the metallic state, and also to carburise the metals, by the action of CO, without the metals becoming fused. The oxides are first reduced, and then carburised by the dissociation of some of the CO. The liberated oxygen combines with CO and is thus removed. Nickel is thus reduced to the metallic state, more or less combined with carbon, and bars of metallic iron are carburised to produce 'cementation' steel. In a similar way a carburised metal may be decarburised by O or CO_2 , in this way 'malleable cast iron' is made. Whether the action consists of carburising or decarburising is entirely dependent on which agent preponderates at the time in the atmosphere surrounding the body.

Reactions for removal of 'impurities' from metals of oxide class The refining of Ni and Co, as we have seen, precedes the reduction to the metallic state. The refining of Sn is effected after reduction, partly by liqation and partly by oxidation, by which means the principal impurities, Fe, As, and W, are removed. Mn and Cr are reduced directly from their ores, in the form of alloys with iron, only pure ores being used. Tungsten is either reduced directly from the ore, alloyed with iron, or undergoes a preliminary purification and ppg as oxide in the wet way. The impurities which it is important to remove from iron are S and P. The latter can only be passed into the slag when the slag is basic, and the conditions tend to oxidation. Thus, P is removed in the primitive iron making processes, also when iron is made by the 'finery' and the 'puddling' processes, also in making 'ingot-iron' by the Siemens and Bessemer processes, when a basic lining to the furnace or converter is used. P is not removed in processes in which pig iron or high carbon steel is made, nor in the Siemens or the Bessemer processes when the furnace or converter is lined with siliceous material, and malleable metal is being produced. In the Bessemer process, with an 'acid lining,' the necessary heat for the process is obtained mainly by the oxidation of silicon in the pig-iron, when a phosphiferous pig iron is used, and a 'basic lining,' most of the heat is obtained by the oxidation of the phosphorus and less silicon is required in the pig-iron. In both processes a portion of the heat is obtained from the oxidation of S. Sulphur can be removed econo-

mically only in the blast furnace, because when treating pig-iron for the production of malleable iron the sulphur is one of the last substances to oxidise out, and its removal would result in a great waste of iron. To remove S in the blast-furnace the temperature must be high, and the conditions must be strongly reducing and carburising: i.e. the charge must contain plenty of charcoal, coal, or coke, and lime must be added to the charge in considerable quantity. Under these conditions the S combines with the Ca, and passes into the slag as sulphide.

SLAG REACTIONS Slags consist of the non-metallic constituents of an ore or furnace product, and of the useless or objectionable metallic elements. They are produced by simply fusing the ore alone, if it is in itself sufficiently fusible, or by fusing it with such materials—fluxes—as will bring about the requisite fusibility. Except in a few special cases, e.g. the reduction of aluminium or magnesium, slags consist of more or less complex silicates. In processes for making more or less malleable iron, the slags are composed mainly of silicates of Fe, Ca, Al, and Mg, with smaller proportions of silicates of Mn, alkalis, and alkaline earths. In pig iron slags, the iron is comparatively small in quantity, or is even entirely absent when much lime is used. In other slags, excepting those produced in treating the 'noble metals,' iron will generally be present in large proportion.

ELECTRO METALLURGY Electro metallurgical processes are divisible into two distinct classes: one class includes processes for the extraction of metals from their ores, the other includes processes for refining metals already extracted. Although a great many processes have been devised for the electrical extraction of metals, except for the extraction of aluminium and magnesium—in which cases the metals are first got into the condition of double chlorides, or, in the case of aluminium, sometimes of fluoride—there is not much prospect of such processes being successfully and economically worked, even when power can be obtained from waterfalls. By the Cowles process, according to Sterry Hunt, nearly pure Al is produced in small buttons, but up to the present it has been necessary to reduce some other metal with the Al to collect it together. In this process the Al_2O_3 and the oxide of the metal to be alloyed with the Al are mixed with carbon, and the mixture is placed round two carbon poles between which an electric discharge is made to pass.

The second class of processes has been advantageously applied to the refining of Cu, more particularly Cu containing small quantities of Au and Ag. In refining Cu electrically, the electrolyte is usually $CuSO_4$, kept at as uniform a temperature and concentration as possible, the anode being formed of the copper to be refined, pure copper depositing on the cathode. The silver and gold collect at the bottom of the tank as a muddy deposit.

METALS. A K H
An element is a definite and distinct kind of matter which has resisted all attempts to separate it into unlike portions. The classification of the elements, in accordance with their chemical properties, necessarily carries with it the classification of many compounds, inasmuch as the chemical properties of an ele-

ment cannot even be stated without considering the composition, general chemical behaviour, and conditions of formation, of compounds of that element. For instance, certain elements are placed in the same class because they all form hydroxides which are alkalis: this statement implies an acquaintance with the composition, methods of production, and chemical properties, of the alkalis, but one of the chemical properties of an alkali is that it neutralises acids, and in so doing forms salts, hence it is necessary to know something about acids and salts, in order to understand what is meant by an alkali, or by an alkali forming element.

The elements may be classified in accordance with their physical properties. If a binary compound is electrolysed, one of its elements separates at the positive electrode, and the other at the negative electrode (secondary reactions which may occur are supposed to be overlooked). That element which separates at the positive electrode is said to be electro positive to the other element. By studying the electrolysis of binary compounds, the elements may be arranged in an electrical series. This series may be divided into two parts: all the elements on one side of any chosen element are electro positive to all the elements on the other side of the chosen element. Taking hydrogen as the central element we are able to subdivide the elements into two classes, all the elements on one side of H are electro positive to the elements on the other side of H. Thus we arrive at a classification of the elements founded on one chemical-physical property. Now we find that the electro-positive elements, on the whole, more resemble one another in certain physical properties, and also in their general chemical character, than they resemble the electro negative elements. Those elements which are electro positive to H as a class are greyish white in colour, lustrous, fairly malleable and ductile, comparatively good conductors of heat and electricity, those elements which are electro negative to H vary much in colour and appearance, they are not usually lustrous, they are generally brittle, and they do not conduct heat or electricity well. Turning to the chemical characters of the two classes of elements, we find that those placed in the electro-positive class generally combine with O to form basic oxides: their compounds with O and H are also usually basic, they do not, as a rule, enter into the composition of acids, very few of them form hydrides, their haloid compounds, as a whole, are tolerably stable as regards the action of heat, and they are not readily decomposed by water, if they are thus decomposed they generally produce oxyhaloid compounds, speaking broadly, these elements do not exist in allotropic forms. On the other hand, we find that most of the elements which are placed in the electro negative class combine with O to form acidic oxides, their compounds with H and O are usually acids. All acids contain one or more of these elements, they generally form hydrides, many of their haloid compounds are decomposed by heat, and many of them are also decomposed by reacting with water, thereby producing haloid acids and either oxides or oxyacids of the electro negative elements; speak-

ing broadly, these elements exhibit the phenomena of allotropy

Hence, it would appear that the division of elements into two classes, those which are electro positive to H, and those which are electro negative to H, is a good classification, because with this one class mark many other properties, both physical and chemical, are associated

The electro positive elements are called metals, the electro negative elements are called non metals

A further examination of metals and non-metals shows that the classification implied in these terms is very far from being sufficient for chemical purposes. We cannot define the term metal, just as we cannot define any of the chemical names which are given to classes of bodies. We can sketch the ideal metal. Considered physically, the ideal metal is a hard, fairly heavy, greyish-white, lustrous, malleable, ductile, and tenacious solid, which melts only at a high temperature, it is a good conductor of heat and electricity, it crystallises in forms belonging to the regular system, its emission spectrum consists of lines, and is comparatively simple in character. Considered chemically, the ideal metal is hardly, if at all, changed in the air, it combines with O at fairly high temperatures, and forms one or more oxides which are basic, &c which react with acids to form salts, it reacts with water or steam to produce an oxide or hydroxide and evolve H, no hydride of the ideal metal is known, it does not enter into the composition of acids, but it reacts with acids to form salts, its sulphides react with acids to produce salts and evolve H₂S, they combine with the sulphides of decidedly electro-negative elements, the salts of the ideal metal are numerous and stable, it forms but few acid salts, but the number of double salts into which it enters is large, it combines directly with the halogens, producing compounds which are volatilised without decomposition at rather high temperatures, and which dissolve in water without change, the ideal metal forms alloys with elements of its own class, which alloys are long rather to the group of physical, than to that of chemical, compounds, lastly, the ideal metal exists in only one modification, &c it does not show allotropy. The ideal non-metal is the opposite, chemically and physically, of the metal

No element exhibits all the properties which we have placed in the category 'metal', nor is there any element which possesses even some of these properties without at the same time also possessing some of the properties which belong to the typical non metal

The elements sodium and potassium possess most of the chemical properties enumerated as characteristic of metals, but these elements are instantly oxidised by exposure to air, they probably form unstable hydrides, they are very soft, lighter than water, and melt at moderately low temperatures.

The element gold possesses most of the physical properties characteristic of metals, but its hydroxide reacts with alkalis to form salts, e.g. KAuO₂; Au also forms the acids H₂AuBr₂ and H₂AuCl₂; Au₂S combines with the sulphides of

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the very metallic elements K and Na to form salts. The element chromium exhibits many of the characteristic physical properties of metals, it also decomposes steam with evolution of H, it combines with the halogens to form stable compounds, some of which have been gasified at high temperatures, chromium does not form a hydride, the oxides CrO and Cr₂O₃ are basic, the element reacts with many acids to form salts, which are well marked, stable compounds, it does not exist in allotropic forms. On the other hand, CrO₃ is a distinctly acidic oxide, reacting with water to form the acid H₂CrO₄, from which is obtained a large number of salts. In other words, chromium belongs to the class metals, and also to the class non-metals. In an even more marked way than Cr, manganese combines in itself both metallic and non metallic properties

The chemical properties of an element depend on the properties of the other elements with which it combines, and on the relative quantities of these other elements entering into combination. It is this fact which makes it impossible to apply the definition of metal or non metal, in its entirety, to any element. The classification of elements into metals and non metals is nevertheless a useful one, provided it is employed with judgment and knowledge. If we find that a certain element is hard, lustrous, unchanged or only slowly changed in the air, and is a good conductor of electricity, or if we find that the oxide of a certain element is basic, and that the chloride is not decomposed by water, or if decomposed produces an oxychloride, we have at once a guide to lead us in our further examination of the element. We shall probably find that the element in question possesses several of the other physical characteristics of metals, and we shall also probably find that it reacts with acids to form salts, decomposes steam with evolution of H, produces at least one sulphide which combines with sulphides of some of the negative or non metallic elements, and so on

The following division of the elements usually placed in the class metals is that arising from the application of the periodic law

CLASS I division	1, Li Na K Rb Cs
"	2, Cu Ag Au
CLASS II division	1, Be Ca Sr Ba
"	2, Mg Zn Cd Hg
CLASS III division	1, Sc Yt La Yb
"	2, Al Ga In Tl
CLASS IV division	1, Ti Zr Ce Th
"	2, Ge Sn Pb
CLASS V division	1, V Nb D ₁ Ta
"	2, As Sb Er Bi
CLASS VI division	1, Cr Mo W U
"	2, none isolated.
CLASS VII division	1, Mn
"	2, none isolated.
CLASS VIII division	1, Fe Ni Co (Cu)
"	2, Rh Ru Pd (Ag)
"	3, Os Ir Pt (Au)

The metals in division 1 of Class I are generally known as the *alkali metals*. They possess in the most marked way the chemical characters of the ideal metal; none of their compounds exhibits any acidic functions, they are electro-positive to all the other elements. The metals

R

in division 2 of Class I are very distinctly metallic in their physical properties, chemically considered they show considerable differences among themselves, several compounds of gold are almost non-metallic in their reactions, the chemical and physical analogies of this element are best represented by placing it both with Cu and Ag, and also with Os, Ir, and Pt, in division 3 of Class VII.

The *alkaline earth metals* which form division 1 of Class II are more metallic in their physical properties than the metals placed in division 1 of Class I, chemically they are also distinctly metallic. Coming to Mg, Zn, Cd, and Hg, which form division 2 of Class II, we have four elements whose properties closely approach those of the ideal metal, Cd may be taken as on the whole the best actual representative of the class metal. As we pass to the higher classes we find many metals exhibiting properties characteristic of non-metals, until in Class VII we arrive at Mn, an element which is at once distinctly metallic and decidedly non-metallic in its chemical properties.

In connexion with the subject of this article, references should be made to the following articles wherein the different classes of metals are described — ALKALINE EARTHS, METALS OF THE, vol 1 p 112, ALKALIS, METALS OF THE, vol 1 p 114, CHROMIUM GROUP OF ELEMENTS, vol 11 p 168, COPPER GROUP, vol 11 p 250, EARTHS, METALS OF THE, vol 11 p 424, IRON GROUP, vol 11 p 65, MAGNESIUM GROUP, vol 11 p 163, NITROGEN GROUP (for Class V, V to Bi), vol 11 *infra*, NOBLE METALS, vol 11 *infra*, TITANIUM GROUP, in vol 14, TIN GROUP, in vol 14.

M M P M

METALS, RARE Under this name are included a number of presumed elementary bodies concerning which our knowledge is at present very imperfect. We see that in their general properties they approximate more or less closely to cerium, yttrium, and lanthanum, but we are not sure how far we have yet obtained them in a state of purity. Consequently we are in doubt not merely as to their at. w. and S.G., but even as to their number and their rank as elements, compounds, or mere mixtures. Our ignorance is due to the great rarity of these bodies, to the high complexity of the minerals in which they are found, but most of all to the fact that they differ among themselves merely by very minute shades.

The principal sources of the rare metals are *gadolinite*, *keilhauite*, *fergusonite*, *euxenite*, *cerite*, and *thorite*, which are Norwegian minerals, and further, *samarshkite*, which was first obtained from the Urals, but has since been found in relatively large quantities in North Carolina. It is to be noticed that specimens of any one of these minerals, if from different localities, are not identical in the earths they contain. Hence if it is desired to isolate any particular earth it is best to select as the first material that mineral in which nature has, so to speak, commenced the task of separation. (This method was proposed by the writer in his address to the chemical section of the British Association, Birmingham Meeting (*C N* 64, 128, *ibid* 54, 157, *Pr* 40, 505). Nilson and Krüss have since adopted and recommended the same method (*B* 20,

2134, *C N* 56, 74, 85, 135, 145, 154, 165, 172).

The recognition of the various rare earths is a matter of no little delicacy. Here spectroscopy in its several modifications renders the greatest service. In endeavouring to ascertain by this means what substances are present in a mineral containing rare earths, chemists may employ either the spark spectrum, the absorption spectrum, or the incandescence or the phosphorescence spectrum. They may further apply any of these tests either at once to the original matter, or to some of its portions after a partial separation has been effected by chemical treatment. The question of course arises, how are we to know when we have obtained any one earth separated from all other bodies, and absolutely pure? In the case of those earths and their solutions which present an absorption spectrum, *e.g.* didymia, samaria, holmia, erbia, &c., the writer has shown that as an element approaches simplicity the absorption spectrum of its solutions will become less and less complicated, hence it would appear that when absolutely free from its associates, each element would have an absorption spectrum of great simplicity, in many cases consisting of one band only (the 'one band, one element' hypothesis). But as certain earths, *e.g.* lanthana, mosandra, philippia, scandia, terbia, &c., give no absorption spectra, this test is not applicable in all cases.

Great caution is required in drawing conclusions from the examination of spectra. Concerning the influence of one body upon another little is yet known, but that little is of sufficient importance to make us very careful how we interpret absorption-spectra when not corroborated by chemical results. Lecoq de Boisbaudran and J. Lawrence Smith have pointed out some important modifications produced in absorption spectra by the presence of an excess of acid in the solution (*C R* 88, 1167). Soret subsequently verified these observations. Brauner and others have put on record experiments on mixing solutions of didymium and samarium. They find in the case of a didymium solution showing the group of three bands, 476, 469, 428 [$1/\lambda^2$ 430.4, 441.3, 454.6], that, by adding a dilute solution of samarium, these three bands vanish, without the appearance of any of the samarium bands, until a certain proportion is reached, when the samarium bands gradually come into their places (Brauner, *C J* 43, 286).

Many of the earths that do not yield solutions giving absorption spectra can be made to give characteristic spectra by phosphorescence. This is known as the 'radiant matter' test. When the spark from a good induction coil traverses a tube having a flat aluminium pole at each end, the appearance of the spark changes according to the degree of exhaustion. If atmospheric air is the gas under exhaustion, at a pressure of about 7 mm a narrow black space is seen to separate the luminous glow and the aluminium plate connected with the negative pole of the induction coil. As the exhaustion proceeds this dark space increases, until at a pressure of about 0.02 mm the dark space nearly fills the tube; the luminous cloud showing the presence of residual gas has almost disappeared, and the molecular discharge from the negative pole

begins to excite phosphorescence on the glass where it strikes the side. There is a very wide difference in the degree of exhaustion at which various substances begin to phosphoresce. Under the influence of this discharge of 'radiant matter,' a great many substances emit, more or less intensely, a phosphorescent light. On examining this light in the spectroscopic, most of these bodies give a faint, continuous, spectrum with a more or less decided concentration in some one part, the superficial colour of the phosphorescing substance depending on this preponderating emission in one or other part of the spectrum. Sometimes, but less commonly, the spectrum of the phosphorescent light is discontinuous.

If we examine the rare earths by this 'radiant matter' test we find they present phenomena of a striking character. Some of them remain unaffected, and are thus at once referred to a distinct group. Others, such as thoria, do not phosphoresce and offer great obstruction to the passage of the spark. Other earths become very phosphorescent and vary greatly in their power of retaining a residual phosphorescence. On examining phosphorescent earths glowing in a vacuum tube, the writer found remarkable differences in the duration of this residual glow. Some of the earths remain luminous for many minutes after the cessation of the current, while others cease to phosphoresce immediately on the stoppage of the current. Take the case of yttria. This earth, the writer finds, can be resolved by chemical treatment into a series of simpler bodies of unequal basicity, to which he has given the provisional designations of Ga, G β , G δ , G ζ , G η , and Sy. The after-glow of these bodies differs somewhat in colour from that which the earth exhibits while the current is still passing. The spectrum of the after glow also shows that some of the lines are missing. In the electrical phosphoroscope—an instrument similar to Becquerel's phosphoroscope, but having the substance acted on electrically instead of by direct light—the different bands of the new constituents of yttria (*in vitro*) do not all appear at the same speed of rotation. At the lowest speed the double greenish blue band of G β is first seen, followed next by the dark-blue band of Ga. As the velocity increases there follows the bright citron yellow band of G δ , and as the utmost speed approaches the red band of G ζ is seen, but not without difficulty. As another instance, if lanthanum sulphate, with traces of Sm as impurity, along with a little lime is examined in the phosphoroscope, the band of Ge is visible at the lowest speed, G δ follows at an interval of 0.035 second, and the Ga band immediately afterwards. All the earths of the yttrium and samarium groups yield discontinuous spectra when submitted to the induction discharge *in vacuo*.

A modification of phosphorescence spectroscopy is produced by the previous addition of other earths to the specially phosphorescent earths. Lime exerts a remarkable action. By itself, it phosphoresces with a continuous spectrum, while yttria phosphoresces with a discontinuous spectrum. But if these two bodies are mixed together, the phosphorescing energy of the lime does not extend over the whole spectrum,

but concentrates itself in strengthening the yttria bands. These bands become broader, but at the same time less sharply defined, in proportion as the lime is increased in quantity. Lime also brings out the phosphorescent bands of samaria. It also suppresses the sharp line S δ , the most striking feature in the phosphorescent spectrum shown by pure samarium sulphate. On the other hand an addition of 'old' yttria deadens the other lines of samaria, but brings out the line S δ more strongly. Lanthanum sulphate in the 'radiant matter' tube phosphoresces with a reddish colour. If lime is added to lanthanum sulphate the phosphorescence changes its colour from red to yellow. Lime also brings out the bands of yttrium and samarium if these are present as impurities. So sensitive is this test that it will show the presence of one part of yttria or samaria in more than a million parts of lime. When G δ , Ga, and G β are present in small proportions with lime the bands of G δ and Ga become intensified, but a dark space appears instead of the green band of G β . Hence if only a small trace of G β is present in lime the green band is not only obliterated, but the quenching action suppresses that part of the continuous lime spectrum which has the same refrangibility as the G β line, and thus gives a black space in the spectrum.

There are many instances of the modifications induced in the normal spectrum of one earth by the admixture of others when treated as anhydrous sulphates. One of the most striking instances is that of a mixture of samaria with yttria, since the presence of even 40 p.c. of yttria practically obliterates the spectrum of samaria. The most minute proportion of lime added to samaria causes the sharp line at $1/\lambda^2$ 269 to vanish, while at the same time it much intensifies the other bands (Tr, pt II, 1885, C R, June 15, 1885). The action of lime upon yttria is of great use in detecting very minute traces of this earth when in admixture with elements which would otherwise prevent its phosphorescence.

Alumina is also active in inducing new spectra when mixed with the rare earths. A moderate amount of fractionation has enabled the writer to penetrate beneath the veil of red phosphorescence observed in crude alumina and to see a complicated sharp line spectrum (C N, 56, 62, 72). The new body of which glimpses have been obtained is probably one of the unknown earths in decipia, since the new spectrum may be fairly reproduced by adding one of the fractionations of decipia to alumina. Hence, it will be seen that the performance of a long series of check and counter-check experiments often becomes necessary before the presence or the absence of any particular earth can be inferred.

The quantitative separation of the rare metals is much more difficult than their mere recognition. These substances are not linked to one another, or to other elements with which they are associated, by any strong affinities, but they are nearly identical in their behaviour and properties. Hence we have so far been unable to find any reagent or any mode of treatment which at once quantitatively separates one of these substances from all the others. We are therefore obliged to have recourse to tedious processes of fractionation.

In attempting to enumerate or describe the rare metals, we meet with the additional difficulty that the unitary character of many of them is still a matter of extreme doubt.

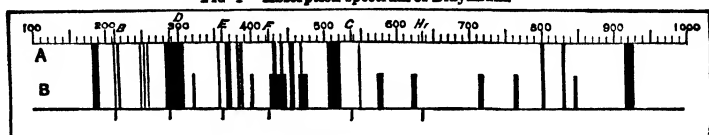
Several of the rare metals will be found described in their proper alphabetical order in this Dictionary, *e.g.* beryllium, cerium, didymium. Many of these, as will be seen below, are probably capable of being further split up, but as they are frequently mentioned in chemical treatises and memoirs their description as at present, or lately, known has been found necessary.

There are certain other metals which are still under discussion, *e.g.* decipium, philippium, holmium (Soret's X), and dysprosium. Roscoe has indeed proved that philippium is a mixture of terbium and yttrium, and the experiments of the writer have confirmed his results, but until we know more about the constitution of terbium and of yttrium, both of which are undoubtedly compound bodies, these experiments do not carry us much further. Samarium is also identical

separate a mixture of two bodies into two parts, just as the addition of a reagent only divides a mixture into two portions, a precipitate and a solution. These divisions will be effected on different lines according to the reagent employed. Thus, if we add ammonia to a mixture we may get a separation into two parts, but if we add oxalic acid to the same original solution we split up the mixture differently and obtain two other parts. Thus, if we crystallise a solution of old didymium, as was done by Auer von Welsbach, we divide its components into neodymium and praseodymium. But by fusing didymium nitrate we divide its components in a different way and obtain different products. Now, it is clear that so long as by different modes of attack we obtain different products, we have not yet reduced the original substance to its ultimate elements, we have not yet reached bed rock.

We find that a compound molecule may behave as an element, as has been shown in the case of old didymium. Chemists have a certain

Fig 1—Absorption-spectrum of Didymium.



The scale is $\frac{1}{\lambda}$

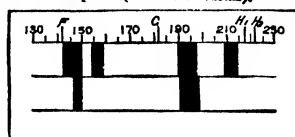
with a body which other chemists have named 'yttrium β '.

It has been mentioned elsewhere that the old didymium, after the elimination of Delafontaine's decipium, was found by Lecoq de Boisbaudran to contain another body, which he named samarium, characterised by the bands of Delafontaine's decipium, together with two additional bands (*cf.* figs 1 and 2). After the removal of these bodies the residual didymium was split up by Auer von Welsbach into the two bodies, neodymium and praseodymium, the absorption spectra of which are shown respectively in figs 3 and 4. It will be observed, however, that two of the bands of old didymium are not to be found in the neodymium and praseodymium spectra taken conjointly. Hence it becomes extremely probable that there exists a third body distinct from neodymium and praseodymium to which one of these extra bands, or possibly both, is due. This probable metal the writer has provisionally named D α . But we still encounter the question whether neodymium, praseodymium, and D α are ultimate elements, or are capable of still further scission. The researches of several investigators point very decidedly in the latter direction. Thus Nilson & Krüss in 1887 appear to have obtained from didymium no fewer than nine bodies, each of which may possibly prove to be an element. These bodies have been provisionally named by the discoverers D α , D β , D γ , D δ , D ϵ , D ζ , D η , D θ , D ι .

It seems to the writer that neodymium and praseodymium are simply the products into which the original didymium is split up by one particular mode of attack. Any single chemical operation, whether it be crystallisation, precipitation, fusion, partial solution, &c., can only

number of reagents, operations, or processes in regular use, and if a substance resists all these and otherwise behaves as a simple body, they call it an element. But for all this it may prove to be a compound. Hence, we may legitimately pause before conceding to neodymium and praseodymium the rank of elements. We need some criterion for an element which shall appeal to our reason more clearly than the old untrustworthy characteristic of having not as yet been decomposed, and to this point chemists would do well to turn their most serious attention.

Fig 2—Absorption spectrum of Samarium and Decipium (De Boisbaudran).



In samarium the writer, by means of the 'radiant matter' test, has recognised four bodies, named provisionally S β , G ϵ , G γ , and G θ . Very similar observations seem to prove that, like didymium and samarium, erbium, holmium, thulium, dysprosium, &c., are compounds or mixtures of a number of closely allied bodies.

In order to ascertain the existence of supposed new elements, chemists have proposed as a test that certain absorption-bands seen in different solutions follow the same variations of intensity. If this is the case we may infer that they are all characteristic of one and the same substance. But if one of the bands dies out while others remain unaltered we may judge that two or more distinct bodies are present.

In erbium the writer has detected two bodies, which he has characterised by their absorption bands $\lambda 550$ and $\lambda 493$. Krüss a Nilson apply apparently to the same bodies the names $\text{Er}\alpha$ and $\text{Er}\beta$. To obviate all confusion it must be remembered that the name 'erbia' has been given to two bodies which are not identical. The substance which ten years ago was called erbia, and which was then supposed to be the oxide of a simple metal, has been resolved by the investigations of Delafontaine, Marignac, Soret, Nilson, Clève, Brauner, and others into at least six distinct earths. Three of these—scandia, ytterbia, and terbia—give no absorption spectra, while three others—erbia (new), holmia, and thulia—give absorption spectra. The first to question the elementary character of old erbium was Delafontaine (*C R* 87, 559, *C N* 38, 202). He obtained from it and described philippia, a yellow oxide having a strong band in the violet ($\lambda 400$ to 405), a broad black absorption band in the indigo blue (about $\lambda 450$), two rather fine bands in the green, and one in the red.

The history of philippium is very instructive. Soon after Delafontaine's discovery, Soret (*C R* 89, 521, *C N* 40, 224) stated that he was unable to identify 'Soret's X' with Delafontaine's philippia, the latter being characterised by an absorption band in the blue occupying the same place as one of the erbia bands. In 1880 Delafontaine (*C R* 90, 221, *C N* 41, 72) described ten new earths as present in gadolinite and samarskite, viz., mosandra, philippia, ytterbia, decipia, scandia, holmia, thulia, samaria, and two others to which he did not assign names. He concluded that the properties of philippia were identical with those of Soret's X and of Clève's holmia, and proposed that the name holmia should be set aside in favour of philippia. In *C R* 91, 328 (also *C N* 42, 185), Clève repeated his earlier assertion that philippia was not identical with Soret's X or holmia. Delafontaine then withdrew all he had said about the absorption spectrum of philippium and decided that it had no absorption spectrum at all (*Archives de Genève* [3] 999, 15). Lastly Roscoe (*C J* 41, 277) gave an elaborate account of the earth metals in samarskite, proving philippia to be a mixture of ytterbia and terbia. The present writer, after prolonged chemical examination of these earths, has come to a similar conclusion, but a spectroscopic examination of the earth left on igniting some very carefully purified crystals of philippium formate, tested in the radiant matter tube, has shown that in the separation of Delafontaine's 'philippium' the ytterbia undergoes a partial fractionation.

Shortly after the announcement of philippium, Soret (*C R* 86, 1062) described an earth which he provisionally named X. It was subsequently found to be identical with Clève's holmia (*C R* 89, 479, *C N* 40, 125). The absorption spectrum of this earth is marked by a very strong band in the extreme red ($\lambda 804$), two characteristic bands in the orange and green ($\lambda 640$ and 536), with fainter lines in the more refrangible part of the spectrum, and a number of bands in the ultra violet (see fig 7). The claim of holmium to rank as an element has been disputed by Nilson and Krüss, who assert that it consists of, or at least contains, four distinct

bodies, provisionally named $\text{X}\alpha$, $\text{X}\beta$, $\text{X}\gamma$, $\text{X}\delta$. By submitting Soret's X to fractional precipitation, and examining the fractions spectroscopically, Lecoq de Boisbaudran found that this X, otherwise holmium, consisted of at least two elements one of these he has named dysprosium, reserving the name holmium for the residue left after the elimination of the dysprosium. The absorption spectrum of dysprosium shows four bands, $\lambda 451.5$, 475 , 756.5 , and 427.5 . The absorption spectrum of what may be called new holmium is shown in fig 8. What relation this new holmium bears to any of the components observed in the original spectrum of holmium by Krüss and Nilson is not as yet determined.

The writer (*Pr* 40, 502) obtained an earth by repeated fractionation, in which one of the bands ascribed to dysprosium, that namely at $\lambda 451.5$, was very strong, though the others were absent. As de Boisbaudran regards the bands $\lambda 475$ and 451.5 as both belonging to dysprosium, and as the earth obtained gives $\lambda 451.5$ strong but with scarcely a trace of $\lambda 475$, dysprosium consists of at least two simpler bodies. Krüss and Nilson in fact resolve it into three bodies to which they have given the provisional names $\text{X}\zeta$, $\text{X}\epsilon$, $\text{X}\eta$, and de Boisbaudran gives the absorption spectrum of dysprosium as shown in fig 9.

Simultaneously with the discovery of holmia, Clève announced the separation of a second earth from erbia, which he called thulia. Its absorption spectrum consists of a very strong band in the red $\lambda 680$ to 707 , and one in the blue $\lambda 464.5$ (fig 10). The ultimate character of thulium is by no means established. Krüss and Nilson resolve it into two bodies, $\text{Tm}\alpha$ and $\text{Tm}\beta$. Nevertheless the atomic weight of thulium has been determined as 170.7 , and the composition Tm_2O_3 has been assigned to its oxide, determinations which for the present must be regarded as premature.

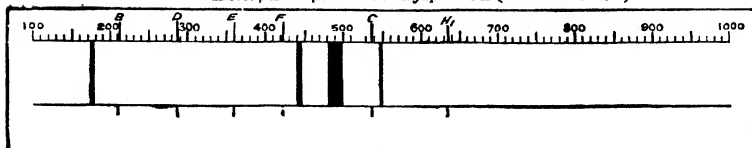
Further, it must be mentioned that the spectrum of old erbium has two faint bands, one at $\lambda 550$ and a second broader one at $\lambda 493$ (fig 11). These bands are not to be found in the spectrum of holmium, thulium, dysprosium, or the new erbium (fig 12). In a long continued fractionation of the erbia group of earths, conducted with an ample supply of the old erbia, the writer finds an earth giving these two bands concentrated at one end, the bands becoming stronger, while at the same time two other bands make their appearance. This phenomenon indicates the existence of another earth as yet unknown, belonging to the erbium group.

We next come to the yttrium group, comprising the metals yttrium, terbium, gadolinium, ytterbium, scandium, mosandrium, columbium, and rogerium. Of these yttrium, terbium, ytterbium, and scandium form the subject of distinct articles in this Dictionary.

Columbium and rogerium were discovered in the samarskite of North Carolina by J. Lawrence Smith in 1879, but nothing further has been published concerning them. This columbium is perfectly distinct from an element sometimes called columbium, but better known as tantalum.

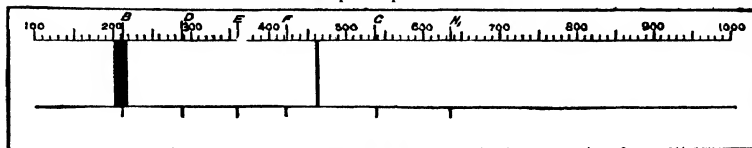
Mosandrium was also discovered by J. L. Smith, and has been the subject of a little controversy. Delafontaine pronounced it a mixture

FIG 9 —Absorption spectrum of Dysprosium (De Bolsaudran)



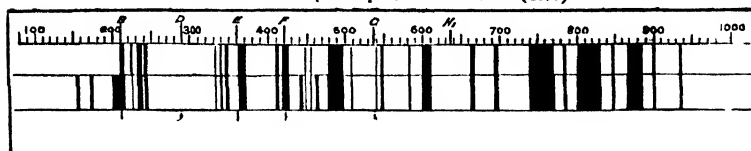
The scale is $\frac{1}{\lambda^2}$

FIG 10 —Absorption spectrum of Thulium



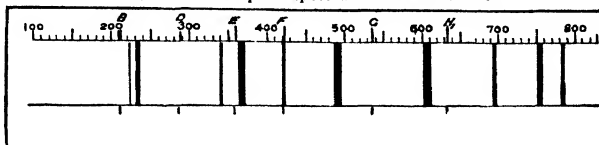
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FIG 11 —Absorption spectrum of Erbium (1878)



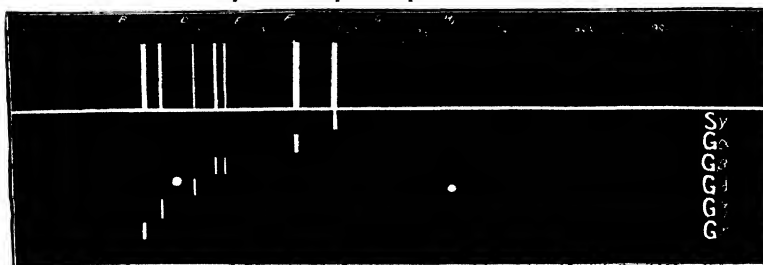
The scale is $\frac{1}{\lambda^2}$

FIG 12 —Absorption-spectrum of New Erbium.



The scale is $\frac{1}{\lambda^2}$

FIG 13 —Phosphorescence-spectra of products of fractionating Yttrium



The scale is $\frac{1}{\lambda^2}$

of terbium, yttrium, erbium, didymium, and phosphorus. In November 1878 and in September 1879 Smith reasserted the elementary character of mosandrum. He states that its compounds are of a deep orange colour, that its double potassium sulphate is not easily soluble, and that its equivalent weight is 51.2. No recent and conclusive investigations on this alleged metal have appeared, and its existence must in the meantime be regarded as exceedingly doubtful.

Decipium has been considered as holding an intermediate position between the didymium and the yttrium groups of rare metals, but its elementary character is very questionable. It has not formed the subject of any recent researches. It is said to show a characteristic absorption band about λ 416.

Gadolinium, otherwise known as the Y_a of Magnac, gives no absorption-spectrum. In the writer's investigations its phosphorescence spectrum was found to consist of those of two of the constituents of yttrium, which will be mentioned below.

Yttrium proves to be an exceedingly complex substance. The body to which all chemists would have applied the name yttria as recently as four years ago may be split up into possibly six, but certainly five, bodies, G_a, G _{β} , G _{δ} , G _{ϵ} , and G _{γ} , two of which, G _{β} and G _{ϵ} , are also met with as the components of gadolinium. Hence it is convenient to speak of the original substance as 'old yttria' and to remember that such is the substance to which everything written concerning yttria prior to 1885 will be found to refer. Fig. 13 shows the simple phosphorescence spectra of the several components into which yttria may be split up by fractionation. If these components are taken in the order of their apparent basicity—the chemical analogue of refrangibility—the lowest of these constituents gives the deep-blue band G_a, then follows a strong citron band, G _{δ} , which increases in sharpness as it becomes more separated from its associates until it may be called a line, then a red band, G _{ϵ} , then a crimson band, G _{γ} , and, lastly, very close together a pair of greenish-blue bands, G _{β} .

The diagram, fig. 14, shows a series of nineteen phosphorescence spectra obtained from a prolonged examination of 'old yttria'. The central spectrum, J, is approximately that given by crude 'old yttria,' though this differs slightly according to the mineral from which the old yttria is extracted. After a time fractionation splits up the earth J into two earths here marked κ and λ , giving slightly different spectra. Fractionating κ gives π and ν , while λ on fractionation yields J and λ . It must not be thought that there is so great a difference between any two adjacent spectra as is here shown. To make the diagram accurately represent what is actually seen in the laboratory it would be necessary to place between each of these nineteen spectra about 1,000 intermediate spectra. Beginning at the extreme red it will be seen that a strong band at λ 647 [$1/\lambda^2$ 289] is at its maximum intensity from ϵ to κ , when it rapidly disappears and is not seen beyond ϵ and κ . The component giving this band the writer names provisionally G _{γ} . The next band in the red λ 689 [$1/\lambda^2$ 245] reaches its maximum at λ or even higher, and fades out be-

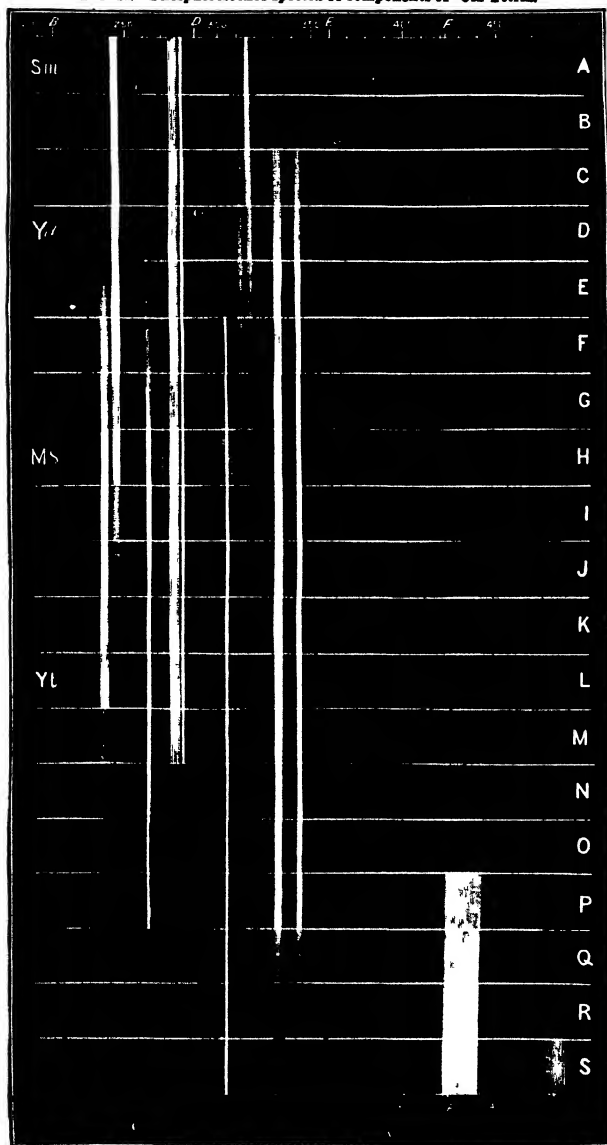
tween κ and λ . The band at λ 619 [$1/\lambda^2$ 261] has its maximum between κ and ϵ , dying out rapidly below but being more persistent above. It is called G _{ϵ} . Then comes an extremely sharp band λ 609 [$1/\lambda^2$ 269] which appears to belong to an earth absent in gadolinite but present in samarskite and a few other minerals. Its greatest brilliancy is between π and κ , and on either side it dies rapidly away. For this the writer proposes the name S _{δ} . Then follows a double orange band, and its two components, though very closely united, are probably capable of separation. The maximum brightness of the first component λ 603 [$1/\lambda^2$ 275] extends from ϵ to the top of the figure. The second component λ 597 [$1/\lambda^2$ 280] begins to fade about ϵ , and is at its greatest brilliancy at the highest spectrum shown on the figure. This band occurs almost isolated in a specimen of crude lanthana, and may be provisionally called G _{ϵ} . Next follows the citron, or G _{δ} band, λ 574 [$1/\lambda^2$ 305.5] which is the most prominent feature in the spectrum of old yttrium. This band extends with scarcely diminished sharpness from ϵ to π , above ϵ it fades rapidly and disappears above ν . Then follows a double green band separable into two components. The first of these, λ 568 [$1/\lambda^2$ 310] is nearly absent in λ , reaches a maximum at ν , and disappears at κ . The second member of this green pair, λ 563 [$1/\lambda^2$ 315] has its maximum at λ and extends only to π . The substance producing this pair of bands may be called for the present G _{γ} . Then follows a pair of bright green bands which so far show no signs of dividing. They begin at π , reach a maximum at ϵ , and continue with scarcely diminished brightness to ϵ . The body giving this double green band is remarkably persistent and may provisionally be called G _{β} . Next comes a dark interval followed by a broad, hazy, double blue band, with its centre at λ 482 [$1/\lambda^2$ 430.5], this band appears at ν and grows brighter to the last fraction at π . The substance to which it is due is called G_a. Lastly, at λ 456 [$1/\lambda^2$ 481] appears a deep violet band beginning at about ϵ and brightening as we proceed lower down. In some samples of ytterbia, supposed to be pure, this band is intensely brilliant, but it is absent in a specimen received from Nilson and considered by him to be perfectly pure. Hence it is probably due to another new body which may be provisionally named S _{γ} .

It must be remarked that the writer's fractionations have been carried far beyond the limits shown in the diagram. Fractions above λ and below π afford evidence that the process of differentiation has not yet reached its utmost limit.

On the left side of the diagram will be seen chemical symbols attached to some of the spectra. Thus the top spectrum, λ , is the one shown by samarium. At ν is the spectrum of Magnac's Y_a, or gadolinium. π shows the spectrum of mosandrum, and κ that which is generally pronounced to be pure yttrium. A careful study of this diagram will lead the observer to conclude that samarium, gadolinium, mosandrum, and yttrium are not true chemical elements but compounds, or perhaps very intimate mixtures, of certain simpler bodies. For these bodies the name 'meta-elements' is proposed (W. Crookes,

B A Birmingham Meeting, *C. N.* 54, 155, | be a complex body, and recognises in it three
 Krüss & Nilson, *B.* 20, 2134 and *C. N.* 54, 71, 85, | distinct substances which he names $Z\alpha$, $Z\beta$, and
 135, 142, 164, 165, 172, Marignac, *Archives des* | 'new yttria' (*C. R.* 103, 627) It is possible

FIG. 14.—Phosphorescence-spectra of components of 'Old Yttria.'



The scale is $\frac{1}{\lambda}$

Sciences Physiques, 16, No 5 and *C. N.* 57, W. | that $Z\alpha$ is $G\beta$, and that $G\beta$ and $Z\beta$ are identical,
 Crookes, *Pr.* Feb 10, 1887) | though de Boisbaudran now regards the latter
 De Boisbaudran likewise finds yttrium to | body as terbium (*C. R.* 102, 395, 902)

The right of the 'new yttria' to bear this name is in the highest degree questionable. De Boisbaudran assigns to it an atomic weight close upon 89, and states that it has a characteristic spark-spectrum but gives no phosphorescence spectrum either in the radiant matter tube or by his reversion-process. It evidently cannot be regarded as the old yttrium freed from impurities, since the purest specimens of such yttrium give a phosphorescence spectrum *in vacuo* of such remarkable intensity that it cannot be due to mere traces of impurities. Ga, Gβ, &c., can no more be spoken of as impurities in old yttrium than can neodymium and praseodymium (assuming them to be elementary) be viewed as impurities in old didymium.

Marignac's Y_a is probably a compound. The writer recognises in it two at least of the constituents of yttrium, Gβ and Gγ.

A. E. Nordenskjöld gives the name oxide of gadolmium (a totally different body from Marignac's Y_a or gadolinium) to the mixture of earths in gadolinite which are precipitated by ammonia and oxalic acid, but not by sulphate of potash. This mixture consists of yttria, erbia, and ytterbia. A very interesting point is that however different the minerals from which it has been obtained, and however the percentage of the rare earths varies, the equivalent weight of the mixture is always approximately the same, viz 261.9. This value is determined by transforming a weighed quantity of the oxide into sulphate by digestion with dilute sulphuric acid, and driving off excess of water and acid by heating to incipient redness (Nordenskjöld, *C. R.* 102, 795, W. Crookes, *C. N.* 54, 239).

On the other hand, commenting on this, De Marignac shows that there is sometimes a variation of 15 p.c. in the equivalent weights of these natural mixtures of rare earths (De Marignac, *Arch. des Sci. Phys.* 17, No 5).

We have therefore some thirty bodies of which the so called rare metals are composed, or, at least, which they contain, and a variety of facts points to the conclusion that we have by no means come to the end. Several even of the new bodies give signs of a capability of further splitting up, if they are examined with sufficient nicety and persistence. It is far from unlikely that when the various methods of research known as fractionation have been more generally applied we may have to deal, not with thirty, but with nearer sixty, unknown bodies.

But whatever may be the number of these bodies the question must be raised, What are they? Are they each and all independent elements? We have certainly no good *a priori* grounds for asserting that the number of elements now recognised is not capable of considerable extension. But before any body can be accepted as simple it should certainly undergo a very severe scrutiny, more severe than of these newly-discovered bodies has yet undergone. The case of didymium is here a warning. It had been closely examined by some of the ablest chemists in Europe, it had been freed from several foreign bodies, its atomic weight had been established, when a novel mode of examination proved its compound nature.

Pending therefore the completion of a series of investigations, chemical and optical, which

will probably occupy several generations of chemists, it may be safest to call these recently observed bodies not, as yet, elements, but quasi or meta elements. Our notions of a chemical element have been enlarged, hitherto the elemental molecule has been regarded as an aggregate of two or more atoms, and no account has been taken of the manner in which these atoms have been agglomerated. The structure of a chemical element is certainly more complicated than has hitherto been supposed. We may reasonably suspect that between the molecules which we are accustomed to deal with in chemical reactions, and the component or ultimate atoms, there may intervene sub molecules, sub aggregates of atoms, or meta elements, differing from each other according to the positions which they occupy in the very complex structures commonly known as didymium, yttrium, and the like. W. C.

METAMERISM This term is generally applied to those cases of *isomerism* wherein compounds show identity of elementary composition, but belong to different types or classes (*v.* *ISOMERISM*, pp. 79, 80, 81, 88).

METEORITES As regards composition, meteorites have been divided into two classes, those which consist for the most part of metals only, and those which are chiefly composed of silicates, sometimes accompanied by unoxidised iron and nickel. Iron and nickel are the principal constituents of those meteorites which are mainly metallic. The quantity of Fe varies from 80 to 95 p.c., and of Ni from 6 to 10 p.c. Co occurs in many meteorites, varying from a mere trace to 2 or 3 p.c., the other metals found in small quantities are Sn, Mn, Cr, and Cu, small quantities of Fe-Ni phosphide, and also carbide, phosphide, silicide, and sulphide of Fe are also frequently found in metallic meteorites. The silicates which form the chief constituents of meteorites of the second class are silicate of Al, Ca, and Na (*labradorite*), silicate of Ca and Mg (*augite*), and silicate of Fe and Mg (*olivine*). These silicates are often accompanied by nickel and iron, also by iron pyrites, and sometimes by chrome iron, and magnetic oxide of iron. M. M. P. M.

METHACRYLIC ACID C₄H₆O₂, *ε*
OH, CMe CO₂H *a Methyl acrylic acid* [16°].
(160.5° i. v.) S.G. ²⁰ 1.0153 (Bruhl, *B.* 14, 2800) $\mu_D = 1.4314$ $n_D^{20} = 35.07$

Occurrence—In small quantity in Roman oil of chamomile (Kopp, *A.* 195, 89).

Formation—1 Obtained as ethyl ether by the action of PCl₅ on oxy-isobutyric ('dimethoxalic') ether CMe₂(OH)CO₂Et (Frankland & Duppa, *C. J.* 18, 133, *A.* 136, 12, Paul, *A.* 188, 52)—2 By the action of fuming HBr on citraconic acid, mesaconic acid, or citraconic anhydride, the resulting bromo-pyrotartaric acid being boiled with conc. NaOH aq. (Fittig, *A.* 188, 95, *B.* 10, 517)—3 By boiling citra and meso-chloro-pyrotartaric acid with alkalis (Pehrn, *A.* 188, 42)—4 Together with oxy-isobutyric acid, by boiling a bromo-isobutyric acid (1 pt) with water (25 pts) (Thomson, *A.* 200, 86).

Preparation—Citraconic anhydride is mixed with a saturated solution of HBr (2 vols.) at 0°. In a few days a mass of crystals of citra-bromo-pyrotartaric acid is got. These are collected,

boiled with Na_2CO_3 , acidified with HCl , and distilled. The distillate is neutralised by CaCO_3 , filtered, evaporated to dryness, and mixed with HCl . Methacrylic acid separates as a light oil (Fittig & C Kolbe, *J pr* [2] 25, 372).

Properties—Long prisms (from water), with strong but not unpleasant odour. V sol water, v a sol alcohol and ether.

Reactions—1 *Bromine* forms di-bromo isobutyric acid—2 *Potash fusion* gives hydrogen, formic acid, and propionic acid (F & D)—3 *Sodium-amalgam* reduces it to isobutyric acid—4 Conc HClAq forms, in the cold, iodoisobutyric acid crystallising in tufts of prisms—5 Methacrylic acid dissolves easily in fuming HBr even at 0° , the solution, after some time, depositing a crystalline addition product and a thick oil (Paul, *B* 9, 122)—6 *Bromine* forms CH_3Br $\text{CBrMe CO}_2\text{H}$ —7 On contact with cold conc HClAq or by heating at 130° , it is converted into an amorphous (? polymeric) modification. The ammoniacal solution of this substance gives white pps with Ba and Ca salts. It is not altered by conc H_2SO_4 and HNO_3 (Fittig & Engelhorn, *A* 200, 70)—8 *Hypochlorous acid* forms chloro oxy isobutyric acid [107°] (α 235°) (Mehlkoff, *Bl* [2] 41, 311).

Salts— CaA' , tufts of long needles, v sol water. Appears to change on keeping to a salt of the polymeric acid (v Reaction 7)— AgA' needles (from boiling water), scarcely affected by light. Suddenly decomposes at 100° .

Constitution—Inasmuch as the di-bromo isobutyric acid, formed by the addition of bromine, is converted by boiling water into a bromo oxy butyric acid that can be reduced to a oxy isobutyric acid, $(\text{CH}_3)_2\text{C}(\text{OH})\text{CO}_2\text{H}$, it is clear that methacrylic acid is not $\text{CH}_3\text{CHMe CO}_2\text{H}$ but $\text{CH}_3\text{CMe CO}_2\text{H}$.

References—BROMO and CHLORO METHACRYLIC ACIDS

METHACYL BROMIDE v BROMO ACETONE

METHÆMOGLOBIN v HÆMOGLOBIN

METHAMIDO v METHYL AMIDO

METHANECH, *Marsh gas* *Methylhydride*

Light carburetted hydrogen. Mol w 16 (-155° to -160°) (-181°) at 6.7 atmospheres, -73.5° at 56.8 atmospheres (Wroblewsky, *C R* 99, 136) S G (air = 1) 553 (Regnault, *C R* 36, 676) S G (liquid) 415 at -164° (Olszewski, *P* [2] 31, 58) S H 593 (R) $\mu_0 = 1.000412$ (Crouillebois, *C R* 67, 692) H F p 21,750 H F v 21,170 (Thomson, *Th*) H C p 213,500 (Berthelot, *A Ch* [5] 23, 179) S 05449 at 0° (Bunsen, *A* 93, 18) S (alcohol) 523 at 0° *Critical temperature* -73.5° (Wroblewsky), -99.5° (Dewar, *P M* [3] 18) *Critical pressure* 56.8 atmospheres (Wroblewsky).

Occurrence—The bubbles of gas given off by decaying vegetable matter in stagnant pools consist of marsh gas, CO_2 , and nitrogen. It often escapes into coal mines, where it is known as fire damp, since it forms an explosive mixture with air. It escapes from the earth in various places, as in Italy, North America, and especially at Baku on the Caspian. It occurs among the products of distillation of wood, peat, coal, and bituminous shale, constituting 35 to 40 p c of coal gas. Methane occurs also in the intestinal gases. It occurs also among the products of the passage of ethylene (Norton & Noyes, *Am* 8,

862) and other gases through a red hot tube. The gaseous product obtained by heating ethylene at 400° contains 36 p c methane and 40 p c ethane (Day, *Am* 8, 153).

Formation—1 By the action of potassium-amalgam on COI , in presence of water (Regnault). 2 By passing a mixture of CHCl_3 or COI , and hydrogen through a red-hot tube (Berthelot)—3 By the action of powdered zinc on chloroform dissolved in aqueous alcohol (Sabanejeff, *B* 9, 1810)—4 By exposing a mixture of CO and hydrogen to the action of electricity in an induction tube (Brodie, *Pr* 21, 245)—5 By the action of water on zinc methide (Frankland)—6 By the action of sodium on MeI in presence of ether (Wanklyn & Buckeisen)—7 In small quantity by passing a mixture of CS_2 and H_2S over red hot copper (Berthelot, *A Ch* [3] 53, 69) 8 By heating CS_2 with PH_3I at 130° (Jahn, *B* 13, 127)—9 Among the products of the dry distillation of barium formate (Berthelot, *J* 1857, 426)—10 By distilling crystallised sodium acetate (2 pts) with KOH (2 pts) and quicklime (3 pts) (Dumas, *A Ch* [2] 73, 92) Von Schlegel (*A* 226, 140) recommends 1 pt of sodium acetate and 2 pts of soda lime (*cf* Schorlemmer, *C N* 29, 7)—11 When river mud is added to a solution of calcium acetate a slow evolution of a mixture of methane (2 vols) and CO_2 (1 vol) occurs, calcium carbonate being left. Calcium lactate undergoes a similar fermentation, the gases being evolved in the same proportion (Hoppe Seyler, *H* 11, 561).

Preparation—By the action of the copper-zinc couple on an alcoholic solution of MeI , the escaping gas being well washed by a scrubber containing a further quantity of copper zinc (Gladstone & Tribe, *C J* 45, 154).

Properties—Colourless gas. V sl sol. water. Much less soluble in alcohol than ethane. Its illuminating power is slight (*cf* L T Wright, *C J* 47, 200). May be liquefied by combined cold and pressure (Cailletet, *J* 1877, 221). Methane is not absorbed by aqueous KOH or by ammoniacal cuprous chloride. When compressed with water below 0° under a pressure of 30 atmospheres it forms a crystalline hydrate, the critical temperature of which is 21.5° (Villard, *C R* 106, 1602, 107, 395).

Reactions—1 When passed through a red-hot tube it is for the most part unaffected, but a little naphthalene is formed. *Electric sparks* partially convert it into carbon, hydrogen, and acetylene (Berthelot, *C R* 67, 1188). When passed over a red hot palladium spiral it is decomposed, if dry, into carbon and hydrogen, and, if moist, into CO and hydrogen (Coquillon, *C R* 86, 1197)—2 It is not attacked by sulphuric acid, by nitric acid, by a mixture of hot conc H_2SO_4 and HNO_3 , by POI , or by chlorine in the dark—3 A mixture of chlorine (2 vols) with methane (1 vol) when exposed to sunlight presently explodes. The explosion may also be brought about by an electric spark. If the mixture be first diluted with CO , and then exposed to sunlight, quiet chlorination takes place, and if excess of chlorine is present chloroform and CCl_4 are formed. A mixture of methane (1 vol) and chlorine (1 vol) exposed to diffused daylight gives methyl chloride. In presence of moisture, chlorine forms HCl , CO_2 , and CO .—

4 A mixture of air and methane passed over a red-hot platinum spiral yields formic acid (Coquillon, *C R* 77, 444)—5 When mixed with nitric oxide and fired by an electric spark CO, and oxygen are among the products (Cooke, *C N* 58, 130)

Constitution—That the four atoms of hydrogen in methane are of equal value may be inferred from the existence of only one set of methyl compounds The same thing may be shown thus—

From CHHHI we may obtain, by treatment with KCy, an acetonitrile CHHHCy which we may call 'α' From this we can obtain CHHH CO₂H, CHHCl CO₂H, CHHCy CO₂H, and 'β' acetonitrile CHHCyH successively But from CHHCy CO₂H we can obtain malonic ether CHH(CO₂Et)(CO₂Et), and thence we can get CHCl(CO₂Et)(CO₂Et), CHCy(CO₂H)(CO₂H), and 'γ' acetonitrile CHCyHH successively The three acetonitriles 'α', 'β', and 'γ' are then found to be identical, hence three at least of the atoms of hydrogen in methane are of equal value The fourth acetonitrile might probably be obtained from CH(CO₂Et), *via* CCl(CO₂Et)₃, and CCy(CO₂H)₃ (Henry, *C R* 104, 1106) Since

$\text{CO} \begin{smallmatrix} \text{Cl} \\ \text{OEt} \end{smallmatrix}$ the product of the action of CO $\begin{smallmatrix} \text{Cl} \\ \text{OEt} \end{smallmatrix}$ on isobutyl alcohol, is the same as CO $\begin{smallmatrix} \text{OEt} \\ \text{OC}_2\text{H}_5 \end{smallmatrix}$, obtained from CO $\begin{smallmatrix} \text{Cl} \\ \text{OC}_2\text{H}_5 \end{smallmatrix}$ and ethyl alcohol, the carbonyl group is united to two atoms of hydrogen of equal value Now from aldehyde CO $\begin{smallmatrix} \text{H} \\ \text{CH}_3 \end{smallmatrix}$ we may obtain

$\text{Cl} \begin{smallmatrix} \text{O} \\ \text{AcO} \end{smallmatrix} \text{C} \begin{smallmatrix} \text{H} \\ \text{CH}_3 \end{smallmatrix}$ whence silver propionate forms $\text{C}_2\text{H}_5\text{O} \begin{smallmatrix} \text{O} \\ \text{AcO} \end{smallmatrix} \text{C} \begin{smallmatrix} \text{H} \\ \text{CH}_3 \end{smallmatrix}$ But from aldehyde we

may also obtain $\text{Cl} \begin{smallmatrix} \text{O} \\ \text{C}_2\text{H}_5\text{O} \end{smallmatrix} \text{C} \begin{smallmatrix} \text{H} \\ \text{CH}_3 \end{smallmatrix}$ whence silver acetate yields $\text{AcO} \begin{smallmatrix} \text{O} \\ \text{C}_2\text{H}_5\text{O} \end{smallmatrix} \text{C} \begin{smallmatrix} \text{H} \\ \text{CH}_3 \end{smallmatrix}$

These two acetyl propionyl derivatives of ortho aldehyde are found to be identical, hence the two atoms of hydrogen in methane which are displaced by oxygen in forming carbonyl are of equal value It follows that there are two pairs of equivalent atoms of hydrogen in methane (Geuther, *A* 205, 203, 225, 290) And since Henry has shown that three of the atoms of hydrogen in methane are equivalent, it follows that the fourth is so also

References—TETRA BROMO, BROMO IODO, BROMO-NITRO, CHLORO IODO, CHLORO-NITRO, and TETRA IODO, METHANE, BROMOFORM, CHLOROFORM, IODOFORM, and METHYL, METHYLENE, and METHENYL COMPOUNDS In fact, all organic compounds may be regarded as derivatives of Methane (Kekulé)

METHANE CARBOXYLIC ACID IS ACETIC ACID

Methane dicarboxylic acid IS MALONIC ACID
Methane tricarboxylic acid C₂H₃O₆ *ie* CH(CO₂H)₃ Formyl tricarboxylic acid
Methenyl tricarboxylic acid

Ethyl ether CH(CO₂Et), [29°] (258°) at 760 mm (195°–205°) at 140 mm S G $\frac{1}{2}$ 1.100 From sodium malonic ether, benzene and ClCO₂Et (Conrad & Guthzeit, *A* 214, 81; B.

12, 1286, cf Claisen, *B* 21, 8897, 3567) Colourless oil V sol alcohol ether In a freezing mixture, it solidifies to long needles or prisms According to Michael (*J pr* [2] 87, 473) it dissolves readily in dilute NaOH forming ONa(CO₂Et)₃, which may be crystallised

Reactions—1 With aqueous KOH at 100° it forms HOEt, K₂CO₃ and potassium malonate—2 Dilute H₂SO₄ and alcoholic NaOH at 0° also form malonic acid; so that a salt of the acid CH(CO₂H)₃ has not been obtained—3 Chlorine forms CCl(CO₂Et)₃ (Conrad *B* 14, 618)

Anilide of the di-ethyl ether
CH(CO₂Et)₂(CONHPh) [124°] From sodium-malonic ether and an alcoholic solution of phenyl cyanate (Michael, *J pr* [2] 35, 452)

Nitrile of the di-ethyl ether v CYANO MALONIC ETHER

METHANE PHOSPHONIC ACID v METHYL PHOSPHINE

METHANE - TRI - QUINOLYL - HYDRO - IODIDE v QUINOLINE IODOFORM

METHANE SELINIC ACID v SELENIUM ORGANIC COMPOUNDS

METHANE SULPHINIC ACID CH₃SO₂ *ie* CH₃SO₂H From ZnMe₂ and SO₂ (Hobson, *A* 106, 287) The aqueous solution of the acid soon decomposes with deposition of sulphur—CaA', (dried at 100°) amorphous—BaA', (dried at 100°) cubes, v sol water, insol alcohol—MgA', aq (dried at 100°)—ZnA', amorphous

Derivative—TRI CHLORO METHANE SULPHINIC ACID

Methane di-sulphinic ether v METHYLENE DIPHENYL DISULPHONE

METHANE SULPHONIC ACID CH₃SO₃ *ie* CH₃SO₃H

Formation—1 By the oxidation of di methyl trisulphide (Cahours, *A Ch* [3] 18, 258), or of dimethyl disulphide (Muspratt, *A* 65, 251)—2 By oxidising methyl sulphocyanide with nitric acid (S G 1 25)—3 By treating tri chloro methane sulphonic acid with sodium amalgam (Kolbe, *A* 54, 174)—4 By heating MeI with aqueous K₂SO₄ at 120° (Colman, *A* 148, 101)

Properties—Syrup which decomposes above 130° Potash fusion forms K₂CO₃, hydrogen, and K₂SO₄ (Berthelot, *J* 1869, 336)

Salts—NH₄A' thin trimetric plates (from absolute alcohol)—LiA' aq—(NaA'), NaI (Colman)—KA' (dried at 100°)—KHA', (dried at 100°)—CaA', S 71 at 20° (Nithaok, *A* 218, 284)—SrA' aq S 83 at 22°—BaA' aq v e sol water, insol alcohol—MgA', 10 aq—FbA' aq—CuA', 5 aq—AgA'

Chloride CH₃SO₂Cl (160°) (N) S G 1.51 From the acid and PCl₅ (Carius, *A* 114, 142) Not attacked by H₂S, by chlorine, or by KCy (McGowan, *J pr* [2] 80, 280) Decomposes aqueous ammonia with evolution of nitrogen

Amide CH₃SO₂NH₂ Formed by passing NH₃ into a solution of the chloride in ether. Prisms (from benzene containing alcohol)

Anilide CH₃SO₂NHPh Large plates (from alcohol) (McGowan)

Derivatives v CHLORO-, and CHLORO-BROMO-METHANE SULPHONIC ACID

Methane disulphinic acid CH₃S₂O₂ *ie*. CH₃(SO₂H)₂ Methylene disulphonic acid. Methionic acid

Formation.—1. A product of the action of

SO₂ on ether (Liebig, *A* 13, 35, Wetherill, *A* 66, 122, Strecker, *A* 100, 199) — 2 From Et₂SO₂ and SO₂ (Hubner, *A* 223, 208) — 3 By the action of hot fuming H₂SO₄ on acetonitrile, acetamide, or sulpho-acetic acid (Buckton & Hofmann, *A* 100, 133, *C J* 9, 241) — 4 Together with sulpho-acetic acid, by the action of ClSO₃H on acetic acid (Baumstark, *A* 140, 82) — 5 By heating lactic acid with fuming H₂SO₄ (Strecker, *A* 118, 290) — 6. From methylene iodide and K₂SO₄ (Strecker, *A* 148, 90) — 7 By heating chloroform with aqueous K₂SO₄ at 180° (Strecker) — 8 From CCl₄, SO₂, K₂SO₄, water, and K₂SO₄ at 125° (Rathke, *A* 161, 152) — 9 By oxidising CH₃(SCy)₂ with HNO₃ (Lermontoff, *B* 7, 1282)

Properties — Very deliquescent needles Not attacked by chlorine or by nitric acid

Salts — (NH₄)₂A'' trimetric crystals, m. sol cold water — K₂A'' needles *S* 7.1 at 22° — BaA'' 2aq pearly tables — PbA'' 2aq prisms, v sol water, insol alcohol — CuA'' 5aq — Ag₂A'' thin plates

Derivative v BROMO METHANE DISULPHONIC ACID

Methane tri-sulphonic acid CH₃S₃O₆ v.e CH(SO₃H)₃ Obtained by heating CH₃O SO₃K (1 pt) with fuming H₂SO₄ at 100° (Theilkuhl, *A* 147, 134) Formed also by the action of aqueous K₂SO₄ at 100° on CH₃C(NO₂)(SO₃K)₂ or on CCl₃(NO₂) (Rathke, *A* 167, 219) Long needles, v.e sol water and alcohol — K₂A''' 2aq small prisms — Ca₂A''' 12aq small prisms, v sol water, insol alcohol — Ba₂A''' 9aq plates Not decomposed by HCl — Pb₂A''' 2aq

METHANE THIOSULPHONIC ACID

Methyl ether C₂H₅SO₂ v.e CH₃SO₂ SCH₃ *Dimethyl disulphoxide* Obtained by warming dimethyl disulphide with nitric acid (*S G* 12) diluted with four times its volume of water (Lukashevitch, *Z* 1868, 641) Oil Readily oxidised by HNO₃ to methane sulphonic acid

METHANTHRENE C₁₃H₁₂ [117°] An isomeride of methyl naphthalene obtained, together with other products, by distilling podocarpic acid with zinc dust (Oudemans, *B* 6, 1125) The product is crystallised from alcohol and sublimed White laminae with violet fluorescence It boils above 360° V.e sol. boiling alcohol, CS₂, and HOAc

Picric acid compound

C₁₃H₇C₆H₃(NO₂)₃OH [117°] Slender orange needles

Methanthrene-quinone C₁₃H₆O₂ [187°] From methanthrene and CrO₃ in HOAc Minute trimetric laminae May be distilled Insol water, sl sol ether, v sol alcohol Reduced to a hydride by aqueous SO₂

METHAZONIC ACID C₁₃H₈N₂O₄ v.e.

NO (CH₃)₂O? [*c* 60°] The sodium salt is formed by acting on nitro methane with alcoholic NaOH, the resulting crystalline pulp being heated on the water bath The upper (alcoholic) layer is poured off, and the lower layer deposits sodium methazonate on cooling It is dissolved in water and reprecipitated by alcohol (Friese, *B* 9, 304) According to Lecco (*B* 9, 706) it is best to add dilute H₂SO₄ and shake with ether If the ether is dried with Na₂SO₄ and evaporated over H₂SO₄, it leaves methazonic acid as large crystals, which may be recrystallised from benzene. It is v. sol

water, alcohol, and ether, m sol benzene, insol petroleum naphtha The Na salt crystallises from alcohol in long prisms

METHENYL-AMIDINE v FORMAMIDINE

METHENYL-AMIDOXIM v FORMAMIDOXIM

METHENYL-AMIDO-*o*-CRESOL C₇H₇NO v.e.

C₆H₄(OH)₂<N>CH [*s* 1 1/2] [39°] (200°) Colourless crystals Prepared by the action of formic acid on amido-*o* cresol (Hofmann & Miller, *B* 14, 570)

Methenyl-amido *p*-cresol C₇H₇NO v.e.

C₆H₄(CH₃)<N>CH [*s* 5 1/2] [46°] Formed from amido *p* cresol by distillation with formic acid (H & M) Crystals

METHENYL (α) AMIDO NAPHTHYL MER-

CAPTAN C₁₀H₇<N>CH [46°] Formed by heating formyl (α) naphthylamine with sulphur Colourless oily fluid Insol water Volatile with steam Base — B₂H₂Cl₂PtCl₄ yellow needles (Hofmann, *B* 20, 1799, 2265)

METHENYL-AMIDO-PHENOL C₇H₇NO v.e.

C₆H₄<N>CH [30 5°] (182 5°) V D (H—1) 112.5 (calc 119) Obtained by boiling formic acid with *o* amido phenol (Ladenburg, *B* 10, 1123) Prisms Hot conc HCl aq decomposes it, forming *o* amido phenol

METHENYL-AMIDO-PHENYL MERCAP-

TAN C₇H₇NS v.e [12] C₆H₄<S>CH (230°) Colourless fluid Sparingly soluble in water, easily in alcohol and CS₂. Has decided basic properties Heavier than water Volatile with steam Isomeric with phenyl mustard oil

Preparation — 1 By reduction of the chloro derivative (vol u p 78) with tin and HCl or with HI — 2 By the action of formic acid on amido phenyl mercaptan (Hofmann, *B* 13, 14) — 3 Formed together with aniline, methyl aniline, and a base C₆H₅NS₂ by boiling dimethyl aniline with sulphur (Mohlau & Krohn, *B* 21, 59)

Properties — Oil, smelling like quinoline and having a bitter taste It is related to thiazole as quinoline is to pyridine The sulphur cannot be displaced by the action of lead hydrate or even by heating with copper to 250° On fusion with KOH it gives formic acid and amido phenylmercaptan PCl₅ at 180° forms the chloro derivative C₆H₄(NS)CCl

Salts — (B'HCl).PtCl₄, sparingly soluble tables, or needles — (B'HCl)AuCl₃ — B₂H₂FeCy₄ *Methylo-iodide* C₇H₇NSMeI [210°]; needles, sl sol cold alcohol, v sol hot water

General character — This base stands in the same relation to quinoline as thiophene stands to benzene In consequence of this analogy the base and its derivatives exhibit great similarity in properties with the corresponding quinoline compounds thus the boiling-points of the methenyl- and ethenyl-*o* amido phenyl mercaptan do not differ much from those of quinoline and methyl-quinoline, and they form crystalline picrates and uncrystallisable chromates They also form colouring-matters analogous to the cyanines (obtained by the action of alkalis upon a mixture of the alkyl-iodides of quinoline and methyl quinoline) Thus by boiling an aqueous solution of the amyl-iodides of methenyl- and

ethenyl amido-phenyl mercaptans with NH_3 , a red colouring matter $\text{C}_6\text{H}_4\text{N}_2\text{S}_2\text{I}$ is obtained which crystallises in four sided violet red plates with green reflex, sol hot alcohol, sl sol cold alcohol, insol water. Neither of the amylo iodides when treated separately, in the same way, gives any colouring matter. In a similar manner to the amylo iodides, the mixed methyl iodides give a compound crystallising in reddish-golden plates, sol alcohol with a scarlet red colour. Similar colouring matters are also obtained from the corresponding derivatives of amido naphthyl mercaptan, or from the latter derivatives conjointly with those of amido phenyl mercaptan, or from those of amido phenyl-mercaptan conjointly with those of quinoline and methyl-quinoline. All the above products are stronger colouring matters than the cyanines (Hofmann, *B* 20, 2262).

Derivatives v AMIDO-, CHLORO-, and NITRO-METHENYL-AMIDO PHENYL-MERCAPTAN

METHENYL-*o*-AMIDO-PHENYL-MERCAPTAN α -CARBOXYLIC ACID $\text{C}_6\text{H}_4\langle\text{N}\rangle\text{C}\text{CO}_2\text{H}$ [108°]

Formation—1 By the action of cold alcoholic caustic potash upon amido imido ethenyl-*o*-amido phenyl mercaptan

$\text{C}_6\text{H}_4\langle\text{N}\rangle\text{C}\text{C}(\text{NH}_2)\text{NH}$, ammonia being eliminated—2 In small quantity by oxidation of ethenyl-*o*-amido phenyl mercaptan

$\text{C}_6\text{H}_4\langle\text{N}\rangle\text{C}\text{CH}_3$ with cold aqueous KMnO_4

Properties—White needles. Sol water and alcohol. It very readily splits off CO_2 giving methenyl amido-phenyl mercaptan (Hofmann, *B* 20, 2256)

METHENYL AMIDO-TOLYL MERCAPTAN

$\text{C}_6\text{H}_4(\text{CH}_3)\langle\text{N}\rangle\text{CH}$ [1 4] [15°] (255°) Prepared by boiling *p*-amido *m*-tolyl-mercaptan with formic acid (Hess, *B* 14, 492). Sol alcohol and ether. Weak base forming unstable salts— $\text{B}'\text{HCl}$, PtCl_4 , needles

METHENYL AMIDOXIM v FORMAMIDOXIM

METHENYL-AMIDO XYLIL MERCAPTAN

$\text{C}_6\text{H}_4(\text{CH}_3)\langle\text{N}\rangle\text{CH}$ From the thio-formyl derivative of *u m* xylylidine $\text{C}_6\text{H}_4\text{Me}_2(\text{NHCSH})$ by oxidising with alkaline K_2FeC_6 (Gudeman, *B* 21, 2549). Heavy oil

METHENYL-BROMO-TOLYLENE-*o*-DI-

AMINE [1 4 2] $\text{C}_6\text{H}_4(\text{CH}_3)(\text{Br})\langle\text{N}\rangle\text{CH}$ [187°]. Obtained by heating bromo-tolylene-diamine with formic acid (Hubner & Schupphaus, *B* 17, 776). Fine colourless needles. Sol alcohol, ether, and acetone, sl sol water and benzene

Salts— $\text{B}'\text{HCl}$ colourless soluble needles— $\text{B}'\text{H}_2\text{SO}_4$ long needles— $\text{B}'\text{HNO}_3$ sparingly soluble colourless needles— $\text{B}'\text{H}_2\text{Cl}_2\text{HgCl}_2$ needles— $\text{B}'\text{H}_2\text{Cl}_2\text{PtCl}_4$ orange crystalline pp— $\text{B}'\text{H}_2\text{CrO}_4$ glistening red needles— $\text{B}'\text{C}_2\text{H}_5(\text{NO}_2)_2\text{OH}$ this picrate forms yellow needles [229°]

METHENYL-TRICARBOXYLIC ACID v.

METHAN-TRICARBOXYLIC ACID

METHENYL FLUORIDE CHF_3 . *Fluoroform*. *V.D.* 250 (sale 244) δ (alcohol) 6

Formed by warming a mixture of iodoform (2 pts), chloroform (1 pt), and silver fluoride (2 pts) (Meslans, *C.R.* 110, 717). Colourless gas, condensing at 20° under 40 atmospheres' pressure. Smells like chloroform, and burns with blue flame forming HF . Sl sol water, chloroform, and benzene. Alcoholic KOH yields KF and potassium formate

METHENYL DI-PHENYL-DIAMINE v. DI-PHENYL FORMAMIDINE

METHENYL-*o*-PHENYLENE-DIAMINE

$\text{C}_6\text{H}_4\langle\text{NH}\rangle\text{CH}$ *Anhydro-formyl phenylene diamine* [167°] (above 360°). Prepared by heating *ortho* phenylene diamine for 5 or 6 hrs with formic acid. The yield is nearly theoretical (Wundt, *B* 11, 826). Large crystals. Monacidic base— $\text{B}'\text{HCl}$, PtCl_4 — $\text{B}'\text{HCl}$ AuCl_3

METHENYL TRI-PHENYL TRIKETONE

v TRI-BENZOYL METHANE

METHENYL DI-TOLYL-AMIDINE v. DI-TOLYL FORMAMIDINE

METHENYL-TOLYLENE-*o* DIAMINE

$\text{C}_6\text{H}_4(\text{CH}_3)\langle\text{NH}\rangle\text{CH}$ [1 2 3] [143°]. Obtained by reduction of methenyl bromo tolylene-*o* diamine (Hubner & Schupphaus, *B* 17, 777). Colourless glistening needles (from benzene). V e sol water and alcohol

Salts— $\text{B}'\text{HNO}_3$ long soluble needles— $\text{B}'\text{H}_2\text{Cl}_2\text{PtCl}_4$ 3aq sparingly soluble orange needles

Methenyl-tolylene-diamine

$\text{C}_6\text{H}_4(\text{CH}_3)\langle\text{NH}\rangle\text{CH}$ [1 3 4] [c 101°]. From the diamine and formic acid (Ladenburg, *B* 10, 1123)— $\text{B}'\text{H}_2\text{PtCl}_4$ yellow prisms

METHIONIC ACID v METHANE DISULPHONIC ACID

METHOXY-compounds v Methyl derivatives of Oxy. compounds

METHRONIC ACID $\text{C}_6\text{H}_4\text{O}$, *s.e.* $\text{C}_6\text{H}_4\text{O}(\text{CO}_2\text{H})_2$, $\text{C}(\text{CO}_2\text{H})_2\text{CH}$

or $\text{CMeO} \parallel \text{CCH}_2\text{CO}_2\text{H}$ (Knorr, *B* 22, 152)

Di methyl furfuranedicarboxylic acid *Sylvane carboxyacetic acid* *Pyrotritanic carboxylic acid* [205°] (F), [207°] (P)

Formation—1 By heating equimolecular weights of aceto acetic ether, Ac_2O , and sodium succinate, and saponifying the resulting mono ethyl methronate $\text{C}_6\text{H}_4\text{O}(\text{CO}_2\text{Et})(\text{CO}_2\text{H})$ with baryta (Fittig, *A* 250, 173, 182). 2 By treating a mixture of glyoxal and aceto acetic ether with a concentrated aqueous solution of ZnCl_2 , and extracting the product with aqueous KOH (Polonowsky, *A* 246, 6, Fittig & Hantzsch, *B* 21, 2135, 3189)

Properties—Needles (from water), v e sol. alcohol, m sol ether and HOAc , m sol hot, v. sl sol cold, water, almost insol CHCl_3 and CS_2 . The aqueous solution gives no colour with FeCl_3 . It is not attacked by reducing agents

Reactions—1 On heating it gives off carbonic acid gas leaving methyl-furfuryl acetic

$\text{OH} \parallel \text{CH} \parallel \text{CMeO} \parallel \text{CCH}_2\text{CO}_2\text{H}$ [188°]. — 2 Dilute

HCl aq at 200° gives CO_2 and acetyl acetone.

Salts— $(\text{NH}_4)'\text{A}'$ 3aq small needles— $\text{C}_2\text{H}_5\text{A}'$: heavy crystalline pp nearly insol.

hot water —CaA'' 2aq —BaA'' 2aq needles (P) —BaA'' amorphous (F) —BaH₂A'' needles, more soluble than the Ca salt (F) —Ag₂A'' aq bulky white pp

Mono-methyl ether MeHA'' [98°]; needles —AgMeA'' white pp (F)

Di methyl ether Me₂A'' Oil

Mono-ethyl ether EtHA'' [76°] Prepared as above Long pointed needles, v sol alcohol, ether, chloroform, and benzene, m sol CS₂, sl sol water —Ca(EtA'')₂ 2aq needles or prisms —Ba(EtA'')₂ 2aq needles, v sol hot, sl sol cold, water —AgEtA'' white matted needles

Di ethyl ether Et₂A'' (300°–305°) (F)

Phenyl hydrazide C₁₀H₁₁N₃O₂ 16 C₁₀H₉O₂N₂HCl₂ [212°] (F)

Phenyl hydrazide of the mono-ethyl ether C₁₀H₁₁N₃O₂ [134°], crystalline Insol cold aqueous alkalis

METHRONOL C₁₁H₂₀ 16

CH CH C CHPh CHMe
| CH CH O CH₂ CHMe ? *Phenyl di methyl*

naphthalene tetrahydride (323°) V D 80 (calc 82) Formed from phenyl methacrylic acid by heating with H₂SO₄ (40 cc) and water (60 cc) (H Erdmann, A 227, 250) Oil Chromic mixture oxidises it to o benzoyl benzoic acid, acetic acid, benzoic acid, anthraquinone, and CO₂

METHYL The radicle CH₃ The name methyl was also given by Frankland and Kolbe (C J 1, 60) to dimethyl C₂H₅, now called ethane The methyl derivatives of hydroxylic compounds are described under the compounds of which they are the ethers e.g. CH₃O C₂H₅, CO₂H₂, the methyl derivative of oxy benzoic acid is described under Oxy benzoic acid

DI-METHYL-ACETAL so called V vol i p 105

METHYL ACETAMIDE v *Acetyl derivative of METHYLAMINE*

METHYL-ACETANILIDE v *Acetyl derivative of METHYL-ANILINE*

METHYL ACETATE v *ACETIC ACID*

METHYL-ACETIC ACID is PROPIONIC ACID

Di-methyl acetic acid is Iso BUTYRIC ACID

Tri-methyl acetic acid v *VALERIC ACID*

METHYL-ACETO ACETIC ACID v vol i p. 22, vol ii p 78

METHYL ACETO-ACETATE v *ACETO ACETIC ACID*

METHYL DI-ACETONAMINE v *ACETON-AMINE*

METHYL-ACETONE v. *METHYL ETHYL KETONE*

Di-methyl acetone v *METHYL ISOPROPYL KETONE* and *DI ETHYL KETONE*

METHYL-TRI-ACETONE-ALCAMINE v. *ALCAMINE*

METHYL-TRI-ACETONINE v *ACETONINES*

METHYL-ACETO-PROPIONIC ACID v *ACE-TYL BUTYRIC ACID*

METHYL ACETO-SUCCINIC ETHER v *ACE-TYL-METHYL-SUCCINIC ETHER*

METHYL-ACETOTHIENONE v. *METHYL-THIENYL METHYL KETONE*

METHYL-ACETYL-BENZENE v. *TOLYL METHYL KETONE*

Di-methyl-acetyl benzene v. *XYLYL METHYL KETONE*

METHYL ACETYLENE v *ALLYLENE*.

Di methyl-acetylene v *BUTINENE*

Di methyl-di-acetylene C₆H₄ 22

CH₃ C C C C CH₃, *Hexinene* [64°] (180°). Formed by oxidising the copper derivative of allylene with alkaline K₂FeCy₄ (Griner, C R 105, 283) Solid, volatile with steam Does not react with ammoniacal cuprous chloride Combines with bromine in the cold, forming C₆H₄Br₂ [44°]

METHYL-ACETYLENE DI QUINOLINE

C₂₁H₁₆N₄ 22

CH₃ CH CH OH CH C N OMe
| N C CH OH C CH C CH OH
Diethyl-ene quinoline methyl quinoline [157 5°]

Formed on heating p amido (Py 3) styryl-quinoline with HCl at 150°, and then gradually adding paraldehyde (Bulach, B 22, 289)

DI-METHYL-ACETYLENE-DI-UREA

C₆H₁₀N₄O₂ 16 CO NH Me CH NH Me
| NH CH NH
CO NH CH NH Me CO *Glycol-di methyl urel.*

[210°] Formed by adding HCl to a mixture of glyoxal and methyl urea (Franchmonta Klobbie, R T C 7, 19) Needles, v sol water, insol ether and ligroin Forms with HNO₃ a di nitro-compound, which is not decomposed by boiling with water

Di-methyl-acetylene di urea C₆H₁₀N₄O₂ 16

CO NH Me NH Me CO Formed by heating di-

methyl diketone with urea in aqueous solution (Franchmont a Klobbie, R T C 7, 251) Prisms or needles, insol ether and CHCl₃, sl sol alcohol Does not melt below 290° Nitric acid converts it into a nitramine

CO NH Me N(NO₂) Me CO, which crystallises

from alcohol in plates, and is decomposed by boiling water into CO₂, N₂O, di methyl diketone, and urea

DI-METHYL β ACETYL PROPIONIC ACID v *MESITONIC ACID*

METHYL-ACETYL-PYRROLE v *METHYL-PYRROL METHYL KETONE*

v *Methyl-di-acetyl-pyrrole* v *METHYL-PYRROL-ENE DI-METHYL-KETONE*

METHYL-ACETYL UREA v *Acetyl derivative of METHYL UREA*

METHYL ACRIDINE C₁₄H₁₁N 16

C₆H₃ CH₃ CMe
| N
C₆H₄ [114°]. Formed by heating

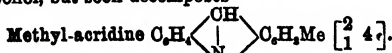
diphenylamine (50 g), glacial acetic acid (80 cc), and ZnCl₂ (85 g.) at 220° for 14 hours The yield is good (82 g) (Bernthsen, A 224, 85) Formed also by heating diphenylamine with acetone nitrile at 200° (Bernthsen, A 192, 29) Dimetric crystals, α = 1 2397 (Bernthsen a. Osann, B 19, 427) Aqueous KMnO₄ at 100° oxidises it very slowly, a product being perhaps quinoline tri carboxylic acid. HNO₃ (S G 138) boiled for 6 hours with methyl acridine (6 g) gives tri-nitro-acridine-carboxylic acid, which forms glittering

prisms, sparingly soluble in the usual menstrua —B'HCl.—B'H₂PtCl₆.

Methylo-iodide B'MeI [185°], red needles (from water). Sol hot water, less sol hot alcohol, least sol ether

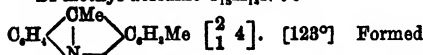
Methylo-chloride B'MeCl. Decomposes at 180°–185°.

Methylo-hydroxide B'Me(OH) From the methylo iodide by treatment with aqueous NaOH Grey powder, turning green in air Sol. alcohol, but soon decomposes



[132°] Formed by distilling diamido-di-*p* tolyl-phenyl methane PhCH(OH)(C₆H₄MeN₂H₂) with zinc dust (Ullmann, *J pr* [2] 86, 265) Yellow matted needles Smells aromatic, and causes sneezing V sol water, v sol alcohol, ether, benzene, and hot petroleum ether Its solution in dilute H₂SO₄ exhibits bluish green fluorescence

Di-methyl-acridine C₁₈H₁₁N + e



by heating phenyl *p* toluidine (12 g) with HOAc (66 g) and ZnCl₂ (18 g) at 230° (Bonna, *A* 239, 64) Needles or prisms (from alcohol) Volatile with steam V sol alcohol and benzene —B'HCl yellow needles —B'HI orange red needles —B'C₆H₄(NO₂)₂OH brown crystals —The sulphate is reddish-brown

Methyl-acridine chloral v **TRI CHLORO OXY-PROPYL-ACRIDINE**

DI-METHYL-ACRYLIC ACID v **PENTENOIC ACID**

TETRA-METHYL ADIPIC ACID C₁₀H₁₈O₄ + e CO₂H CH₂·OMe₂·CMe₂·CH₂·CO₂H? [210°] A product of the action of finely-divided silver on α -bromo-isovaleric ether CHMe₂·CHBr CO₂Et (O Hell a W Mayer, *B* 22, 62) Snowy-white granules (from benzene), v sol alcohol and chloroform. Forms no anhydride on heating (difference from di-isopropyl succinic acid which accompanies it in its preparation) May be sublimed —BaA'' 5aq S 12.5 at 15° —MnA'' 3aq S 5.9 at 25° —NiA'' 4aq S (of NiA'') 4.7 at 25° —CoA'' 3aq. S (of CoA'') 6.65 at 23° —ZnA'' S 195 at 18° —CdA'' 3aq S 2.29 at 81° —CuA'' S 0.24 —PbA'' S 0.45 at 18° —AgA'' S 0.46 at 18°

METHYL-ESCOULETIN v. **ESCOULETIN**.

METHYLAL v **FORMIC ALDEHYDE**

METHYLALACREATINE v **ALACREATINE**

METHYL ALCOHOL CH₃O·e·OH, **OH Oxy-methane** *Methyl hydroxide* *Wood spirit* *Wood naphtha* Mol w 32 (84°8') (R Schuff, *A* 220, 99), (65°9') (Perkin, *O J* 45, 465), (66°2') (Zander) S G $\frac{7}{8}$ 7953 (Brühl), $\frac{11}{16}$ 79726, $\frac{11}{16}$ 78941 (Perkin), $\frac{11}{16}$ 8111 (Zander) S V 42.7 (5), 42.6 (Lossen, *A* 254, 55) μ_D = 1.3832 n_D = 1.293 (Brühl) H F p 51,450 (Thomsen), 61,400 (Stohmann, *J pr* [2] 40, 358) H.F.v. 50,560 (Thomsen, *Th*). H O 170,600 (Stohmann). M.M.1 640 at 18.7°. *Compressibility*. 00010879 at 15° (Dupré, *Pr* 20, 386)

Occurrence.—Free in the juices of plants (Guthzeit, *J*, 1879, 905, Maquenne, *O R* 101, 1067) and in the aqueous distillate in the fruits of *Horaceum* (Guthzeit, *A*, 177, 844; 240, 248).

Methyl alcohol occurs also in the product of the distillation of wood (Taylor [A.D. 1812], *Tilloch's Phil Mag* 60, 315, Dumas a Peligot, *A* 15, 1; *A Ch* 58, 5, 61, 193) and of colophony (Kalbe a. Lwow, *B* 16, 851) It occurs as methyl salicylate in oil of winter-green (*Gaultheria procumbens*)

Formation.—1 From methane by chlorinating, and heating the resulting methyl chloride with aqueous KOH for 7 days at 100° (Berthelot, *A* 105, 241) The MeCl may also be treated with HOAc and NaOAc at 200°, and the resulting MeOAc saponified MeCl treated with Ag₂SO₄ and H₂SO₄ at 100° gives HMeSO₄, whence methyl alcohol can be readily obtained —2 From hydrogen cyanide by reduction with zinc and H₂SO₄, and treating the resulting methylamine with nitrous acid (Linnemann, *Z* [2] 4, 284) —3 By distilling calcium formate (Lieben a. Paternò, *G* 3, 290, *A*, 167, 293, Friedel a. Silva, *O R* 76, 1545)

Preparation.—The crude watery liquid obtained by the distillation of wood contains methyl alcohol, acetone, acetic acid, methyl acetate, and ammonium acetate It is separated from the tarry products and redistilled, the first tenth of the distillate is mixed with slaked lime and redistilled, a little sulphuric acid is then added, and the liquid redistilled and rectified over quicklime The crude wood spirit thus obtained has a strong aromatic odour and turns brown on keeping. It contains methyl acetate, acetone, di-methyl acetic orthoaldehyde CH₃CH(OMe)₂ (Daneer, *C J* 17, 222), allyl alcohol, methyl ethyl ketone, and other ketones (Grodzky a. Krämer, *B*, 9, 1920) Methyl alcohol may be obtained from wood spirit by saturating with fused CaCl₂ and heating on a water bath The residue is a compound of methyl alcohol with CaCl₂, which when distilled with water is decomposed, giving off the methyl alcohol The distillate is then rectified over quicklime (Kane, *A* 19, 164) According to Gould (*C J* 7, 811) it is best to distil the wood spirit with conc NaOHaq (to saponify MeOAc) and then to dehydrate with K₂CO₃ before saturating with CaCl₂ A better method for obtaining methyl alcohol from wood spirit is to distil the spirit (1 pt) with H₂SO₄ (1 pt) and potassium oxalate (2 pts), the crystalline di-methyl oxalate is then decomposed by distillation with water, and the MeOH dried over lime (Wöhler, *A* 81, 376) The oxalate may also be obtained by dissolving oxalic acid, dehydrated at 100°, in boiling MeOH, leaving the solution to cool, and washing the crystals with cold water as long as the filtrate exhibits the iodoform reaction (Erlenmeyer, *N Rep Pharm*, 28, 624). An alternative method consists in passing HCl into a solution of benzoic acid in wood spirit, distilling, ppg with water the portion collected above 100°, decomposing by boiling for several hours with aqueous NaOH, distilling, and rectifying over quicklime (Carius, *A*, 110, 210). The following modification of Wöhler's process is recommended by Britmar a. Fawcitt (*Tr E* 38, 509) 100 c.c of wood spirit are digested with 150 g of powdered NaOH, and then distilled on a water-bath, 500 g. of oxalic acid crystals are mixed with 200 c.c. of sulphuric acid, and then 400 c.c. of the alcohol, purified as above, is added, and the whole cautiously heated on a

water-bath The methyl oxalate thus obtained is thoroughly dried by pressure, and the alcohol regenerated by digestion with water at 70°. To dehydrate the alcohol thus obtained, digestion with baryta, lime, and dried copper sulphate is necessary. Even when prepared from the oxalate methyl alcohol is not quite pure, as it contains bodies which exhibit Lieben's iodoform reaction. It may be freed from these by dissolving in it one tenth of its weight of iodine, gradually adding NaOH until the liquid is distinctly alkaline, and distilling. The product, rectified over CaO, has a specific gravity of 81 at 15° (Regnault & Villejean, *A Ch* [6] 4, 430, *C R* 99, 82). Methyl alcohol may also be prepared by saponifying methyl formate (8 pts.) with NaOH (2 pts.) and water (6 pts.) (Grodzky, *A Kramer*, *B* 9, 1928, Baidy & Bordet, *A Ch* [5] 16, 564). Methyl alcohol may be obtained very nearly pure by distilling oil of winter green with potash, and rectifying the distillate over lime.

Properties—Colourless mobile liquid, with purely spirituous odour. Burns with a pale flame. Mixes with water, alcohol, and ether, dissolves fixed and volatile oils and most resins. Hence it is much used as a solvent. In aqueous solution it produces intoxication, in concentrated solution it is poisonous. When perfectly dry it dissolves a little CuSO_4 , acquiring a bluish green colour, but it does not dissolve crystallised $\text{CuSO}_4 \cdot 7\text{aq}$ (Klepl, *J pr* [2] 25, 526). Methyl alcohol is etherified by acetic acid more rapidly and completely than any other alcohol (Menschutkin, *Z P C* 1, 611). Dittmar & Fawsitt give the specific gravity of mixtures of methyl alcohol and water as follows:

Pct. of CH_3OH	Sp. gr. at 15°	Sp. gr. at 15.56°
0	99907	99907
10	98429	98262
20	97233	96808
30	96057	95467
40	9457	93697
50	92873	91855
60	90917	89799
70	88687	87487
80	86314	85035
90	83751	82396
100	81015	79589

Older determinations are by Ure (*P M* [3] 19, 51), Dupré (*Pr* 20, 336), and Duclaux (*A Ch* [5] 13, 87). The same observers find the vapour tension of methyl alcohol, measured in millimetres of mercury at 0°, to be

Temp	Tension	Temp	Tension
0	29.7	40	259.4
10	53.8	50	409.4
20	94.0	60	624.3
30	158.9	64.96	760.0

(cf. Kononoff, *P* [2] 14, 40). The following vapour tensions are given by Richardson (*C J* 49, 762)

Temp	Tension	Temp	Tension
-8.3	17.8	39.2	235.13
+0.7	32.06	49.2	370.26
11.2	55.82	59.9	584.24
22.7	108.96	65.7	753.05
30.2	155.32		

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The specific heat of aqueous methyl alcohol was found by Dupré to be as follows—

Weight p.c. of methyl alcohol	Specific heat	Weight p.c. of methyl alcohol	Specific heat
10	98582	60	80177
20	95914	70	75500
30	92658	80	69999
40	89219	90	64282
50	84645	100	54325

Estimation—Pure methyl alcohol does not give Lieben's iodoform reaction with iodine and potash.

The amount of methyl alcohol in commercial wood spirit may be estimated by adding 5 c.c. of the spirit, drop by drop, to 30 g. of PI_2 in a flask provided with inverted condenser. The flask is then heated for 5 minutes in boiling water, and the MeI then distilled off by inclining the condenser. The volume of MeI is read off in a graduated receiver 5 c.c. of methyl alcohol yield 7.19 c.c. of methyl iodide (Krell, *B* 6, 1310, cf. Grodzky & Kramer, *B* 7, 1495, 9, 1928). If it is desired to determine how much of the methyl alcohol is in the form of MeOAc , the spirit may be boiled with standard alkali, and the amount used up in the saponification determined.

Baidy and Bordet (*A Ch* [5] 16, 565) modify Krell's process, using only 15 g. of PI to 5 c.c. of wood spirit, but adding 5 c.c. of aqueous HI subsequently. They distil the liquid at 90°–90°, and at the end of the distillation add 5 c.c. of water and quickly distil. The methyl iodide is washed once or twice with water and measured. Di methyl ortho acetic aldehyde or methyl acetal $\text{CH}_3\text{CH}(\text{OMe})_2$ boils at 64° and gives MeI on treatment with PI_2 , but it does not react with aniline, so that when the wood spirit is to be used in preparing di methyl aniline, by heating with aniline hydrochloride, the estimation by PI_2 will give too high results.

The presence of acetone is objectionable in methyl alcohol that is to be used for preparing di methyl aniline. Its amount can be determined by shaking 1 or 2 c.c. of the alcohol with 20 or 30 c.c. of normal aqueous KOH, adding 20 or 30 c.c. of a decinormal solution of iodine, until the solution is clear. The liquid is then acidified with HCl (SG 1.025), excess of decinormal sodium thiosulphate added, and the excess determined by titrating back with iodine solution. If 1 c.c. of methyl alcohol be taken, the weight of acetone in 100 c.c. will be found by multiplying the amount of iodine used up by 7.612 (Messinger, *B* 21, 3366, cf. Kramer, *B* 13, 1002, Hintz, *Fr* 27, 182).

The estimation of methyl alcohol in ethyl alcohol can be effected by oxidation with a standard solution of $\text{K}_2\text{Cr}_2\text{O}_7$, for 1 pt. of ethyl alcohol reduces 4.278 pts. of the dichromate in becoming oxidised to acetic acid, while 1 pt. of methyl alcohol, in being oxidised to CO_2 , and water, reduces 9.224 pts. of $\text{K}_2\text{Cr}_2\text{O}_7$. A solution of 80 g. $\text{K}_2\text{Cr}_2\text{O}_7$ and 150 c.c. H_2SO_4 per litre is employed, and the unreacted chromate estimated volumetrically (O. Hehner, *An* 12, 25).

When 10 c.c. of pure ethyl alcohol are shaken with 1 c.c. of a 1 p.c. solution of KMnO_4 at 20°, no reduction takes place for 20 minutes, but if the ethyl alcohol contains crude wood

spirit, decolourisation takes place at once (Haber mann, *Fr* 27, 668)

Reactions — 1 Passage through a red hot tube yields acetylene and other products (Berthelot, *C R* 50, 805) — 2 In contact with platinum black and air it is oxidised to formic aldehyde and formic acid — 3 Heated with ammonium chloride in a sealed tube at 300° it yields mono-, di-, and tri methylamine (Berthelot) — 4 Bleaching powder does not yield chloroform (Goldberg, *J pr* [2] 24, 115) — 5 Potassium gives off hydrogen, forming KOMe — 6 When its vapour is passed over heated potash, hydrogen is given off, and potassium formate, oxalate, and carbonate are successively formed — 7 Conc H₂SO₄ forms HMeSO₄, which, when heated with excess of methyl alcohol, yields di methyl oxide When distilled with excess of H₂SO₄ di methyl sulphate passes over — 8 Converted into methyl chloride by treatment with HCl, with PCl₅, or with S₂Cl₂ The joint action of H₂SO₄ and HBr gives rise to methyl bromide and ethyl bromide (Niemilovitch, *M* 10, 820) — 9 SOCl₂ forms MeSH, Me₂SO, MeCl, and other products (Carius, *A* 111, 93) — 10 SO₂ forms CH(OH)(SO₂H)₂ (Max Muller, *B* 6, 1031) — 11 When acidulated with 5 p.c sulphuric acid and submitted to electrolysis it forms CO₂, CO, methyl formate, methyl sulphate, and methylal (Renard, *C R* 80, 236) — 12 Distillation over heated *vinc dust* splits it up into CO and hydrogen, a very little methane being also formed (Jahn, *B* 13, 983, *M* 1, 378) — 13 Zinc chloride forms CH₄, water, Me₂O, hexa methylbenzene, and traces of propylene, butylene, and other hydrocarbons (Le Bel a Greene, *C R* 87, 260, 89, 413) — 14 Aldehyde (1 vol), MeOH (2 vols), and HCl form CH₃CH(OMe)₂ (Claus a Trauer, *B* 19, 3004) — 15 Not affected by *B aceti* growing in solutions containing it (Brown, *C J* 49, 177) — 16 Converts *m* and *p* diazo benzoic acid into C₆H₅(OMe)CO₂H, while *o* diazo benzoic acid only gives benzoic acid (Griess, *B* 21, 978) — 17 When heated with hydrochlorides of aromatic bases (e.g. aniline, xylylidine, piperidine) it often displaces hydrogen in the nucleus by methyl (Hofmann, *B* 15, 2895, Ladenburg, *B* 16, 2057) — 18 Unlike EtOH, it hardly reacts when heated with phenol and ZnCl₂ (Auer, *B* 17, 669) — 19 Unlike butyl and amyl alcohols, it does not form a homologue of benzene when heated with benzene and ZnCl₂ at 270° (Goldschmidt, *B* 15, 1066) — 20 ClCONH₂ forms methyl allophanate NH₂CO NH CO₂Me [208°] (Gattermann, *A* 244, 40)

Metallic derivatives — KOMe Obtained by dissolving K in methyl alcohol — (KOH)₂(MeOH)₂ or (KOMe)₂(MeOH)₂ 3aq [c 110°] Obtained by evaporating a solution of KOH in methyl alcohol (Göttig, *B* 21, 1832) Large elastic plates, somewhat heavier than water, on which they move about rapidly, being decomposed — (NaOH)₂(MeOH)₂ or (NaOMe)₂MeOH aq Obtained by dissolving NaOH in dry MeOH and evaporating Crystalline mass, which moves about on the surface of water while decomposing and dissolving — NaOMe 13aq Obtained by evaporating a solution of NaOH in not quite dry methyl alcohol (Göttig, *B* 21, 561) — NaOMe(MeOH)₂ (Fröhlich, *A* 202, 295) CO passed over NaOMe at 160°

forms acetic acid CO passed into a hot mixture of NaOMe and NaOAc forms propionic acid (Geuther a Fröhlich, *A* 202, 312) When distilled with the Ba salt of a carboxylic acid it displaces the carboxyl by hydrogen Thus succinic acid may be converted into propionic acid (Mai, *B* 22, 2135) — TIOME From TIOEt and excess of MeOH (Lamy, *J* 1864, 466) — (Ba(OMe))₂BaO.H₂O aq White nacreous plates, obtained by evaporating at 135° a solution of BaO in MeOH (De Forcrand, *C R* 102, 1397, 1557) Perhaps identical with Ba(OMe) aq described by Dumas and Paillet (*A Ch* 58, 17) 1 litre of MeOH dissolves 200 g of BaO The heat of formation of the compound from solid 3BaO and liquid 4MeOH is 62,500 If a considerable quantity of water is added to the solution of BaO in MeOH the hydrate BaO 10aq is precipitated The compound BaO2Ba(OMe)₂ 2aq is readily formed, even if as much as 3 p.c water be present in the methyl alcohol, but by evaporating in the cold over H₂SO₄ crystals of Ba(OMe) 3aq are formed These are sol water and alcohol By passing HCN into a solution of baryta in methyl alcohol there is formed barium cyanomethylate Ba(OMe)CN.MeOH This is a crystalline powder, sol water, less sol methyl alcohol It loses at 100° the methyl alcohol of crystallisation

Combinations — CaCl₂ 4MeOH Methyl alcohol dissolves CaCl₂ with great rise of temperature, and on cooling this compound separates in large six sided tables (Kane, *A* 19, 168) It is not decomposed at 100°, but water readily liberates the MeOH — LiCl 3MeOH — MgCl₂ 6MeOH (Simon, *J pr* [2] 20, 377) — SbCl₅MeOH [81°] Slightly yellow crystals (from hot alcohol), decomposes at 130°, giving off MeCl and HCl (W C Williams, *C J* 30, 463) — CuSO₄ 2MeOH Minute bluish green crystals, obtained by shaking anhydrous CuSO₄ with methyl alcohol (Forcrand, *C R* 102, 551)

METHYL ALDEHYDE is FORMIC ALDEHYDE
METHYL - ALIZARIN v DI OXY METHYL

ANTHRAQUINONE

METHYL-ALLANTOIN C₄H₇N₃O₃ [225°] Formed by oxidising methyl uric acid with cold aqueous KMnO₄ (Hill, *B* 9, 1090) Monoclinic prisms, decomposed by fusion V sol hot water, sl sol alcohol, insol ether Conc HIAq splits it up into urea and methyl hydantoin — AgC₄H₃N₃O₃ prisms, sl sol cold water

METHYL-ALLOXAN C₄H₅N₃O₄ 1c

CO < NH CO > CO Formed by oxidising methyl uric acid with HNO₃ or with KClO₃ and HCl (Hill, *B* 9, 1092) Formed also by treating theobromine (2 pts) with KClO₃ (9 pts) and HCl (S G 106) at 50° (Maly a Andreasch, *M* 3, 108, cf Fischer, *A* 215, 304) It is converted by alkalis into methyl alloxanic acid, and by boiling HNO₃ into methyl parabanic acid H₂S forms di methyl alloxantin KHSO₄ forms a compound B'KHSO₄ aq, which crystallises in large monoclinic prisms

Di methyl alloxan C₄H₇N₃O₄ 1c

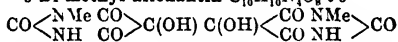
CO < NMe CO > CO A product of the action of KClO₃ and HCl on caffeine at 50° It is extracted by ether (E Fischer, *A* 215, 257, Maly a Andreasch, *M* 3, 92) Colourless six sided

tables (containing 2aq), which slowly turn red in air. V sol water, almost insol alcohol, insol ether. Dyes the skin red. With FeSO_4 and ammonia it gives a characteristic indigo colouration. After drying over H_2SO_4 , it is left as an amorphous powder (containing aq), sol alcohol and ether. Di methyl alloxan decomposes at 100° . It prevents the ppn of cupric and ferric salts by potash. Hydrogen sulphide converts it into amalic acid. KHSO_4 forms the compound $\text{C}_4\text{H}_4\text{N}_2\text{O}_4\text{KHSO}_4$, which crystallises in long tables, $S 7.2$ at 20° , almost insol alcohol, insol ether. This compound may be crystallised from warm water, and does not give the indigo colouration with FeSO_4 and ammonia.

METHYL ALLOXANIC ACID $\text{C}_4\text{H}_4\text{N}_2\text{O}_5$. Formed by the action of alkalis on methyl-alloxan. When the product of the action of HNO_3 on methyl uric acid is neutralised with CaCO_3 , and then mixed with alcohol and ammonia the salt $\text{CaC}_4\text{H}_4\text{N}_2\text{O}_5$ is ppd (Hill, *B* 9, 1092). This salt is gelatinous, and when boiled with water it gives off methylamine.

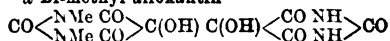
METHYL-ALLOXANTIN $\text{C}_4\text{H}_4\text{N}_2\text{O}_5$. Obtained by mixing solutions of diuric acid and methyl alloxan (Andreasch, *M* 3, 431). Crystallises from water in minute thin tables (containing 3aq).

s Di methyl alloxantin $\text{C}_{10}\text{H}_{10}\text{N}_4\text{O}_5$ &c



Formed by passing H_2S into an aqueous solution of methyl alloxan (Maly & Andreasch, *M* 3, 109). Thin plates (containing 4aq), almost insol alcohol and ether, sl sol cold water. Turns red in air. Gives a red colouration with ammonia, and a violet colouration with potash or baryta (Andreasch, *M* 3, 109).

u Di-methyl alloxantin



Formed by reducing di methyl alloxan to di methyl dialuric acid, and ppg the aqueous solution of this acid with alloxan (Andreasch, *M* 3, 428). Crystallises from hot water in minute four sided pyramids (containing aq), insol alcohol and ether.

Tetra methyl alloxantin v AMALIC ACID

METHYL ALLYL v BUTYLENE

METHYL-ALLYL ACETO ACETIC ACID v ACETO ACETIC ACID

DI METHYL-ALLYL-AMINE *Methylol codida* $\text{C}_4\text{H}_8\text{NMe}_2$. From allylamine and MeI (Bono, *B* 20, Ref 137). When distilled with KOH it yields NMe_3 and an aldehyde $\text{C}_4\text{H}_8\text{O}$ (130° - 135°).

p **METHYL-ALLYL BENZENE** $\text{C}_{10}\text{H}_{12}$ &c $\text{CH}_3\text{C}_6\text{H}_4\text{C}_3\text{H}_5$ p *Allyl toluene* (192°). Formed from cymene by chlorinating and heating the resulting $\text{CH}_3\text{C}_6\text{H}_4\text{C}_3\text{H}_5\text{Cl}$ with alcoholic KOH (Errera, *G* 14, 283, 505). Liquid. Combines with bromine KMnO_4 oxidises it to p toluic acid. HBrAq ($S 1.59$) at 200° forms a polymeride ($\text{C}_{10}\text{H}_{12}$) (350°), and this on keeping changes to an amorphous solid, which by distillation is reconverted into p methyl-allylbenzene.

METHYL-DI-ALLYL-CARBINOL v OCTINYL ALCOHOL

Di methyl allyl-carbinol v. HEXENYL ALCOHOL.

METHYL-ALLYL-CHLORACETOL v DI-CHLORO HEXYLENE

METHYL ALLYL ETHER v METHYL ALLYL OXIDE

METHYL ALLYL DIKETONE $\text{C}_8\text{H}_{10}\text{O}_4$ &c $\text{CH}_3\text{COCOCH}_2\text{C}_6\text{H}_5$ *Acetyl crotonyl* ($c 130^\circ$). Obtained by distilling its mono oxim with dilute H_2SO_4 (Otte & Von Pechmann, *B* 22, 2124). Yellow oil with irritating odour. Not obtained pure.

Mono oxim $\text{CH}_3\text{COO}(\text{NOH})\text{C}_6\text{H}_5$ *Nitroso-allylacetone* *Methyl nitroso butenyl ketone* (146°). From allyl aceto acetic acid and nitrous acid. White plates (from ligroin), v sol usual menstrua. Its solution in NaOHAq is yellow.

Di oxim $\text{CH}_3\text{C}(\text{NOH})\text{C}(\text{NOH})\text{C}_6\text{H}_5$ (135°). Formed from the mono oxim by warming with hydroxylamine. Small plates.

Phenyl hydrazide oxim $\text{CH}_3\text{C}(\text{N}(\text{HPh})\text{C}(\text{NOH})\text{C}_6\text{H}_5$ (137°). Brownish-red needles (from benzene and ligroin).

METHYL ALLYL OXIDE $\text{C}_4\text{H}_6\text{O}$ &c $\text{CH}_3\text{OCH}_2\text{C}_3\text{H}_5$ *Allyl methyl ether* (46°) $S 8.11$ 77 $H F p 34.080$ $H F v 32.050$. From allyl bromide and NaOMe (Henry, *B* 5, 455). With bromine it forms $\text{CH}_3\text{OCH}_2\text{CHBrCH}_2\text{Br}$ (185°). ICl gives $\text{CH}_3\text{OCH}_2\text{CHCl}$ (196°) (Silva, *B* 8, 1469).

METHYL ALLYL PROPYL CARBINOL v OCTENYL ALCOHOL

METHYL ALLYL SULPHIDE $\text{C}_6\text{H}_8\text{S}$ &c $\text{CH}_3\text{SC}_3\text{H}_5$ ($c 92^\circ$) $V D 43.36$. Formed by heating lead methyl mercaptide $(\text{CH}_3\text{S})_2\text{Pb}$ with allyl bromide and ether at 100° (Obermayer, *B* 20, 2925).

METHYL ALLYL THIO-UREA

$\text{CH}_3\text{NHCSNHCH}_2\text{C}_3\text{H}_5$ (50.5°). Formed from allylamine and methyl thio carbimide or from methylamine and allyl thio carbimide (O Hecht, *B* 23, 286). White mass.

METHYL AMIDO ACETIC ACID $\text{C}_4\text{H}_7\text{NO}_3$ &c $\text{CH}_3\text{NHCH}_2\text{CO}_2\text{H}$ *Methyl glycolic* *Sarcosine* $Mol w 89$ (210° - 215°). Formed by the action of boiling baryta water on creatine or caffeine (Liebig, *A* 62, 310, Rosengarten & Strecker, *A* 157, 1, Schilling, *C* C 1884, 811). Formed also by heating chloro acetic ether with aqueous methylamine at 125° (Volhard, *A* 123, 261).

Properties — Trimetric prisms, v e sol water, sl sol alcohol. Has a somewhat sweet taste. Heated to 210° - 220° the greater part evolves H_2O giving the anhydride $\text{C}_4\text{H}_7\text{N}_2\text{O}_2$, whilst a smaller portion splits up into di methyl amine and CO_2 (Mylus, *B* 17, 286). Chloride of cyanogen passed into the fused substance gives methyl hydantoin and sacosine-anhydride (Traube, *B* 15, 2110). When taken internally, the greater part passes as such into the urine (Baumann & Mering, *B* 8, 587, Salkowski, *H* 4, 107, Schiffer, *H* 5, 266). Nitrous acid passed into its hot aqueous solution forms a nitroso-derivative $\text{CH}_3\text{N}(\text{NO})\text{CH}_2\text{CO}_2\text{H}$, which is a thick liquid, forming the salt CaA^+ , aq crystallising in needles (Schultzen, *Z* 1867, 616). Gives methyl uric acid when heated with urea (Horbaczewski, *M* 6, 356).

Salts — $(\text{C}_4\text{H}_7\text{NO}_3)_2\text{Cu}$ 2aq ultramarine blue crystals (*E* Schmidt, *A* 217, 273) — $\text{C}_4\text{H}_7\text{NO}_3\text{HCl}$ needles (from alcohol) — $(\text{C}_4\text{H}_7\text{NO}_3)_2\text{ZnH}_2\text{O}$ 8 (alcohol) 038 V e. sol water (Bulginsky, *J*.

1867, 495) — $(C_2H_5NO)_2H_2PtCl_2$, 2aq monoclinic tables, $a b c = 1.0381 \ 1 \ 0.747$, $\beta = 75^\circ 27'$ — $C_2H_5NO \cdot H_2AuCl_4$, yellow needles, sl sol cold water — $C_2H_5NO \cdot HNO_3$, [c 70°] Very hygroscopic, v sol cold alcohol (Franchimont, *R T C* 2, 839) — $(C_2H_5NO)_2H_2SO_4$, aq four sided tables, v e sol water *S* (boiling alcohol) 10

Guanidine hydrochloride

$C_2H_5NO \cdot HClCH_2N_2$. Formed by heating methyl-amido acetic acid with guanidine hydrochloride (Baumann, *B* 7, 1151) Tables (from alcohol)

Anhydride $C_2H_5N_2O_2$, *e*

$CH_3 \cdot N \cdot CH_2 \cdot CO$

$\begin{array}{c} | \\ OC \cdot CH_2 \cdot N \cdot CH_3 \end{array}$ (?) *Sarcosine anhydride*

[150°] Formed by the action of cyanogen chloride on melted sarcosine Prepared by heating sarcosine to 210° – 220° and distilling the residue (Traube, *B* 15, 2112, Mylius, *B* 17, 286) Colourless prisms V sol water, al cohoh, and ether Bitter taste Weak base By boiling with water it again gives sarcosine On oxidation with $KMnO_4$ it yields *s di methyl oxamide*

Salts — $B'_2H_2Cl_2PtCl_2$, 2aq prisms — $B'_2H_2Cl_2PtCl_4$, 4aq large six sided tables — $B'_2HClAuCl_4$, 2aq prisms

Sarcosine-uric acid $C_5H_8O_8N_4$. Obtained by heating a mixture of sarcosine (3 pts) and uric acid (2 pts) at 210° , a good yield is obtained

Properties — Colourless prismatic crystals (containing 2aq) Sol hot water It shows the murexide reaction It slowly reduces alkaline solutions of copper It has weak acid and basic properties, dissolving in aqueous acids and alkalis By fusion with KOH at 110° it is resolved into uric acid and sarcosine Heated with bromine water it is converted into bromo sarcosine-mesouric acid $C_5H_8N_4O_8Br$

Salts — The acetate forms microscopic crystals, sl sol hot acetic acid, insol cold $HOAc$, v sol water The formate is very analogous to the acetate The ammonium salt is crystalline — $A''Ag$, insol white amorphous pp (Mylius, *B* 17, 518)

Bromo-sarcosine mesouric acid $C_5H_8N_4O_8Br$ Formed by digesting a warm aqueous solution of sarcosine uric acid with bromine, which gives a nearly theoretical yield (Mylius, *B* 17, 521) Heavy colourless tables Sl sol water By H_2S it is reduced to sarcosine mesouric acid It is readily decomposed by alkalis, but is stable towards acids

Sarcosine-mesouric acid $C_5H_8O_8N_4$ Obtained by reducing a hot aqueous solution of bromo sarcosine mesouric acid with H_2S (Mylius, *B* 17, 524) Trimetric tables or needles V sol water, insol alcohol It is a strong acid, but also possesses weak basic properties By bromine water it is reconverted into the bromo derivative, similarly chlorine water gives the chloro-derivative It is very oxidisable, reducing $AuCl_3$, alkaline, copper solution, $KMnO_4$, &c, very readily

Salts — The acetate forms an unstable crystalline pp, sl sol acetic acid — $A''HNH$, fine needles, v sol water — $A''Ag$, white amorphous pp

Tri-methyl-amido acetic acid internal anhydride *v* BETAIN.

DI-METHYL-AMIDO ACETIC ORTHALDE

HYDE *Methylo-hydroxide* C_2H_5NO , *e* $CH(NMe_2OH)CH(OH)_2$. *Muscarine* Occurs, together with neurine, in the fly agaric (Schmie deberg a Koppe, *J* 1870, 875) Formed from neurine by oxidation with conc HNO_3 (Schmie deberg a Harnack, *J* 1876, 804) Deliquescent crystalline mass, sol alcohol Alkaline in reaction Active narcotic poison Separated from neurine by placing the mixed hydrochlorides on filter paper, when the muscarine salt deliquesces and is absorbed by the paper (Harnack, *J* 1876, 803) — $B'(C_2H_5NOCl)_2PtCl_2$, 2an — $C_2H_5NO \cdot AuCl_3$

Di-ethyl derivative of the methylo hydroxide C_2H_5NO , *e*

$CH_3(NMe_2OH)CH(OEt)_2$ *Hydroxide of tri-methyl-amido-acetal* When chloro acetal $CH_2ClCH(OEt)_2$ is treated with tri-methyl amine there is formed $CH_3(NMe_2Cl)CH(OEt)_2$ and C_2H_5NOCl , which is probably $CH_3(NMe_2Cl)CHO$ The two salts are separated by fractional ppn by platinum chloride — $(C_2H_5NOCl)_2PtCl_2$, orange crystals, sl sol cold water — $C_2H_5NO \cdot AuCl_3$, lemon yellow needles

Di-methyl amido acetic aldehyde *Methylo hydroxide* C_2H_5NO , *e* $CH_3(NMe_2OH)CHO$ Formed by saponifying $CH_3(NMe_2OH)CH(OEt)_2$ with baryta (Berlnerblau, *B* 17, 1142) The hydrochloride C_2H_5NOCl is also formed as above Gives the aldehyde reactions — *Platinochloride* $(C_2H_5NOCl)_2PtCl_2$, octahedra (from dilute alcohol)

DI METHYL-AMIDO-ACETONE *Methylo-chloride* C_2H_5NOCl , *e* $CH_3COCH_2NMe_2Cl$ *Caprine chloride* Formed by the action of dry NMe_3 upon chloro acetone in the cold (Niementovitch, *M* 7, 241) Very deliquescent shining needles, sol alcohol Gives the alkaloidal reactions Acts physiologically like curare — $C_2H_5NOClAuCl_3$, [189°]

p DI-METHYL-AMIDO-ACETOPHENONE $C_6H_5(NMe_2)COCH_3$, [59°] Formed by methylation of *p* amido acetophenone (Klingner, *B* 18, 2694) Yellowish plates (from hot water) V sol alcohol, ether, and hot water

o DIMETHYL AMIDO ANISOL. *v* *o* Di-METHYL-AMIDO PHENOL *methyl ether*

METHYL AMIDO-AZO-COMPOUNDS. *Azo-*COMPOUNDS

METHYL AMIDO BENZAMIDE *v* *Amide of* METHYL AMIDO BENZOIC ACID

METHYL-AMIDO BENZENE *v* METHYL-ANILINE and TOLUIDINE

Di-methyl-amido-benzene *v* XALIDINE, METHYL TOLUIDINE, and DI METHYL-ANILINE

Tri-methyl amido benzene *v* MESIDINE and ψ CUMIDINE

Tetra-methyl amido-benzene *v* TETRA

METHYL PHENYL AMINE

Tetra methyl di amido benzene *v* TETRA

METHYL-PHENYLENE DIAMINE

Di-methyl-tri-amido-benzene

$C_6H_5(NMe_2)(NH_2)_2$, [184°] [44°] (298°) Needles. Readily turned blue in the air Prepared by reduction of dinitro dimethyl aniline [87°] Formed also by reduction of nitro-*p* dimethyl amido-phenyl oxamic ether

Acetyl derivative

$C_6H_5(NMe_2)(NH_2)(NHAc)$ [158°] Prisms (containing aq) (Wurster a Sendtner, *B* 12, 1805).

***u*-Tri-methyl-tri-amido-benzene**

$C_6H_3(NMe_2)(NH_2)(NHMe)$ [1 z 4] [90°] (294°)
Prepared by reduction of the nitrosamine derived from nitro trimethyl *p*-phenylene diamine (Wurster a Schobig, *B* 12, 1812) White needles Sol water Gives a blue colouration with HNO_3

Diacetyl derivative [184°] Leaflets

METHYL-AMIDO-BENZENE-AZO- COMPOUNDS v Azo- COMPOUNDS and Dis azo- COMPOUNDS**DI-METHYL-AMIDO-BENZENE PHOSPHINIC ACID** $NMe_2, C_6H_4, P(OH)_2$ [162°]

Formed by decomposing its chloride with water (Schenk a Michaelis, *B* 21, 1498) White needles, v sol hot water and alcohol On boiling its aqueous solution it is split up into di-methyl aniline and phosphorous acid When heated alone it gives phosphorus, PH_3 , and di-methyl aniline

Salt— $NaHA''$ 2aq large crystals (from alcohol)

Chloride NMe_2, C_6H_4, PCl_2 [66°] (250° at 120 mm) Formed by the action of PCl_3 (100 g) on di-methyl aniline (70 g) in presence of $AlCl_3$ (20 g) The mixture is boiled for eight hours with inverted condenser, and the product extracted with petroleum ether and distilled *in vacuo* Thin plates, v sol benzene, m sol ether, sl sol petroleum ether Converted by the action of chloro benzene and sodium into di-methyl amido tri phenyl phosphine

DI-METHYL AMIDO BENZENE PHOSPHONIC ACID $NMe_2, C_6H_4, PO(OH)_2$ [133°]
Formed by oxidising $NMe_2, C_6H_4, P(OH)_2$ with $HgCl_2$ (Schenk a Michaelis, *B* 21, 1500) Crystals, v e sol water and alcohol

METHYL AMIDO BENZENE SULPHONIC ACID $C_6H_4NSO_3, v e NHMe, C_6H_4, SO_3H$ Formed, together with methane disulphonic acid, by heating the acetyl derivative of methyl aniline with H_2SO_4 at 145° (Smyth, *B* 7, 1240) Crystals, which decompose at 182° without previous fusion — BaA', aq crystalline powder

Methyl-amido-benzene sulphonic acid
 $NHMe, C_6H_4, SO_3H$ Formed by heating methyl-aniline ethyl sulphate at 210° (Mundelius *B* 7, 1350) Plates (containing aq), insol alcohol and ether — $BaA', 3' aq$ small prisms — $CaA' 4aq$ — $PbA', 8aq$

Di-methyl-amido-benzene *p*-sulphonic acid
 NMe_2, C_6H_4, SO_3H [150°] (S), [257°] (V a G)

Formation —1 By heating di-methyl aniline with rather more than the equivalent quantity of sulphuric acid at 185° (Smyth, *B* 6, 344, 7, 1237, Armstrong, *B* 6, 663, Vignon, *C R* 107, 263) —2 From di-methyl aniline and $ClSO_2Et$ (Wenghoffer, *J pr* [2] 16, 448) —3 From bromo-di-methyl-aniline and H_2SO_4 at 180° (Michler a. Walder, *B* 14, 2177) —4 From di-methyl aniline and $SOCl_2$ followed by water (Michaelis a Godchaux, *B* 23, 555)

Properties —8 sided prisms (containing aq). Decomposes at 230°

Salts — $BaA', 3aq$ needles or plates — $BaA', 5aq$ (Vignon) — $BaA', 11aq$ efflorescent trichinic crystals (Laar, *J pr* [2] 20, 262) — $NaA' 2aq$ Long needles, m sol water, sl sol aqueous $NaOH$, insol water (M a G.)

Chloride $C_6H_4(NMe_2)_2SO_2Cl$

Ethyl ether EtA' [85°] Formed from alcohol and the chloride

Di-methyl-amido-benzene *p*-sulphonic acid Anhydride of the methylo hydroxide

$C_6H_3NSO_3, v e C_6H_4, < \begin{smallmatrix} NMe_2 \\ SO_2 \end{smallmatrix} > O$ Formed by allowing a mixture of amido benzene *p*-sulphonic acid, MeI , conc KOH aq, and $MeOH$ to stand for some time in the cold (Griess, *B* 12, 2116) Four sided plates, v sol cold water, almost insol alcohol, insol ether Decomposes before fusion Ppd as periodide by a solution of iodine in $HIAq$ —
 $(SO_2, H_2C_6H_4, NMe_2, Cl), PtCl_3, 8aq$ orange tables, v e sol cold water

METHYL-*o*-AMIDO-BENZOIC ACID

Amide $NHMe, C_6H_4, CONH_2$ [160°] Formed by heating *o* amido benzamide with MeI at 100° (Weddige, *J pr* [2] 36, 152) Plates, v sol hot alcohol, sol water In dilute solutions it shows blue fluorescence

Acetyl derivative

$NHMe, C_6H_4, CO NHAc$ [155°] Long needles, v sol hot alcohol, sl sol benzene and ether When heated above 155° it forms the anhydro derivative $C_6H_4, < \begin{smallmatrix} CO \\ NMe \end{smallmatrix} > CMe$ [199°] which is an oxy di-methyl quinoxaline

Nitroso derivative of the amide

$NO NMe, C_6H_4, CO NH_2$ [149°] Pale yellow prisms (from hot alcohol) (Finger, *J pr* [2] 37, 431)

Methyl-*m*-amido-benzoic acid $C_6H_4NO_2, v e NHMe, C_6H_4, CO_2H$ Formed by boiling (a)-benzcreatin $NH, C(NH)NMe, C_6H_4, CO_2H$ with baryta water (Griess, *B* 8, 325) Nodular groups of plates, v sol hot water Nitrous acid ppts a nitroso-derivative Salt — $C_6H_4NO_2, HCl$ six-sided plates

Di-methyl-*m*-amido-benzoic acid $C_6H_4NO_2, v e NMe_2, C_6H_4, CO_2H$ [151°] Obtained by saponifying its methyl ether Needles, sl sol hot water

Methyl ether NMe_2, C_6H_4, CO_2Me (270° uncor) Obtained by fusing the isomeric anhydride of the methylo hydroxide (*v infra*) (Griess, *B* 6, 587) Heavy yellowish oil, sol acids — $C_6H_4, MeNO_2, H_2SO_4$ very small prisms, sl sol dilute sulphuric acid, v sol water — $(C_6H_4, MeNO_2), H, PtCl_3$ spindle shaped laminae, v sl sol cold water

Anhydride of the methylo-hydroxide

$C_6H_3NO_3, v e C_6H_4, < \begin{smallmatrix} NMe_2 \\ CO \end{smallmatrix} > O$ *Benzetaine*
Formed by allowing a mixture of *m* amido benzoic acid with $MeOH$ (1 mol), aqueous KOH (3 mols), and MeI (3 mols) to stand in the cold, the product, after freeing from $MeOH$ by distillation, being saturated with HI , whereupon the iodide NMe_2, I, C_6H_4, CO_2H is ppd This iodide is then boiled with water and lead hydroxide (Griess) Small deliquescent needles (containing aq), which give up their water of crystallisation at 105° V e sol cold alcohol, insol ether Tastes bitter and is neutral in reaction Forms $(NMe_2, Cl, C_6H_4, CO_2H), PtCl_3, 4aq$ crystallising in large prisms, sl sol hot water

Methylo iodide NMe_2, I, C_6H_4, CO_2H Formed as above Small short prisms (containing aq), sl sol cold water

Methylo-chloride NMe_2, Cl, C_6H_4, CO_2H

Monoclinic crystals, a b c = 1.939 1 0.876, β = 88° 49' (Zingel, *Z* *K* 10, 414)

Di-methyl-*p*-amido-benzoic acid

$\text{NMe}_2\text{C}_6\text{H}_4\text{CO}_2\text{H}$ [235°] Formed by boiling its chloride with water. Prepared by boiling for three hours a mixture of *p* amido benzoic acid (1 mol), MeI (2 mols), and aqueous KOH (3 mols) dissolved in methyl alcohol (Michler, *B* 9, 400). Obtained also by heating tetra methyl-di amido benzophenone with soda lime at 340°, extracting the product with hot water, and ppg by acetic acid (E Bischoff, *B* 22, 341). Short needles (from alcohol), sol aqueous KOH and HClAq , insol dilute acetic acid. Nitrous acid forms $\text{NMe}_2\text{C}_6\text{H}_4(\text{NO})\text{CO}_2\text{H}$ [224°]. The Ca salt forms yellowish plates.

Methyl ether $\text{NMe}_2\text{C}_6\text{H}_4\text{CO}_2\text{Me}$ [103°] Silvery plates (from alcohol), *v* sol benzene, ether, and chloroform, *m* sol dilute alcohol. Nitrous acid forms $\text{NMe}_2\text{C}_6\text{H}_4(\text{NO})\text{CO}_2\text{Me}$ [101°].

Chloride $\text{NMe}_2\text{C}_6\text{H}_4\text{COCl}$. Obtained by heating di methyl aniline with COCl_2 at 50° in a sealed tube (Michler). Crystalline.

Nitrile $\text{NMe}_2\text{C}_6\text{H}_4\text{CN}$. From di methyl *p* phenylene diamine by Sandmeyer's reaction (Ahrens, *B* 20, 2958).

Anhydride of the methylo hydroxide

$\text{C}_6\text{H}_3\text{NO}_2$ i.e. $\text{C}_6\text{H}_3\text{<}\overset{\text{NMe}_2}{\text{CO}}\text{>O}$ [c 255°]. When *p* amido benzoic acid is mixed with MeI, KOH, and methyl alcohol, there is formed, even in the cold, di methyl *p* amido benzoic acid and the iodide $\text{NMe}_2\text{C}_6\text{H}_4\text{CO}_2\text{H}$ which crystallises in short yellow plates [233°] (Michael & Wing, *Am* 7, 195). The corresponding anhydride crystallises from alcohol in colourless plates (containing aq), *v* sol water. It loses its water of crystallisation at 100°. The periodide forms long dark prisms [200°]. The platinumochloride ($\text{NMe}_2\text{C}_6\text{H}_4\text{CO}_2\text{H}$), PtCl_4 , forms large red prisms.

Tetra-methyl-di-amido-benzoic acid Di-methylo di-hydroxide

$(\text{HO NMe}_2)_2\text{C}_6\text{H}_4\text{CO}_2\text{H}$ [135]. The iodide $(\text{INMe}_2)_2\text{C}_6\text{H}_4\text{CO}_2\text{H}$ is formed when di amido-benzoic acid (1 pt) is mixed with MeI (6 pts), methyl alcohol (10 pts), and twice as much of a concentrated solution of KOH as is required to neutralise the acid. The mixture should stand for some time, being kept alkaline by further additions of potash. The alcohol is then distilled off, and the iodide ppd by HIAq (Griess, *B* 7, 59). This iodide crystallises from hot water in six sided tables or plates, *v* sol hot water. With moist Ag_2O it yields a caustic alkaline liquid which, on evaporation, leaves the hydroxide as a hygroscopic mass of soft white plates. It rapidly absorbs CO_2 , and ppt metallic oxides from their salts. The corresponding chloride $(\text{NMe}_2\text{Cl})_2\text{C}_6\text{H}_4\text{CO}_2\text{H}$ aq, obtained by neutralising the hydroxide by HCl, crystallises in small six sided plates, *v* sol water, *sl* sol hot alcohol. The periodide forms brownish yellow needles. The carbonate $\text{CO<O NMe}_2\text{>C}_6\text{H}_4\text{CO}_2\text{H}$ 3aq, obtained from the iodide and Ag_2CO_3 , crystallises from water in very small soluble plates, having an alkaline reaction. The platinumochloride $(\text{ClNMe}_2)_2\text{C}_6\text{H}_4\text{CO}_2\text{HPTCl}_4$ aq is a pp composed

of very small pale yellow plates, usually grouped in stars.

Derivative v CHLORO METHYL AMIDO BENZOIC ACID

DI-METHYL-*p*-AMIDO-BENZOIC ALDEHYDE, $\text{C}_6\text{H}_3\text{NO}$ i.e. $\text{NMe}_2\text{C}_6\text{H}_4\text{CHO}$ [78°]. Formed, together with CHCl_3 , by heating $\text{NMe}_2\text{C}_6\text{H}_4\text{CH(OH)Cl}$, with alcoholic KOH (Bossneck, *B* 18, 1520, 19, 366). With di methyl aniline and hydrochloric acid it forms $(\text{NMe}_2\text{C}_6\text{H}_4)_2\text{CH}$.

Oxim $\text{NMe}_2\text{C}_6\text{H}_4\text{CHNOH}$ [144°]. Yellowish brown plates (Knöfler & Bossneck, *B* 20, 3195).

Phenyl hydrazide $\text{NMe}_2\text{C}_6\text{H}_4\text{CHNHPh}$ [148°]. Needles (from alcohol).

DIMETHYL AMIDO BENZOPHENONE

$\text{C}_6\text{H}_5\text{CO C}_6\text{H}_4\text{NMe}_2$ *Benzodimethylaniline Benzoyl di methyl aniline* *Benzoyl phenyl di methyl amine* [90°]. Formed by heating the methylo iodide to 181°.

Formed also by heating malachite green $(\text{NMe}_2\text{C}_6\text{H}_4)_2\text{C(OH)C}_6\text{H}_5$ with conc HClAq at 180°. Colourless plates (from alcohol). Insol water, *sl* sol cold alcohol, *v* e sol hot alcohol or ether. It is a feeble base, its solution in concentrated acids being ppd by water.

Methylo iodide $\text{C}_6\text{H}_5\text{CO C}_6\text{H}_4\text{NMe}_2\text{I}$. Large tables, *sl* sol cold water. Formed by heating *p* amido benzophenone with MeI at 100°. It decomposes at 181° (Doebner & Weiss, *B* 14, 1836, *A* 210, 270, 217, 257).

Di methyl-amido-benzophenone

$\text{C}_6\text{H}_5\text{CO C}_6\text{H}_4\text{NMe}_2$ [39°] (330°-340°). Obtained by heating benzoic acid with di methyl aniline and P_2O_5 at 185° (O Fischer, *A* 206, 88). Needles (from ligroin). Forms unstable salts. With nitrous acid it yields the oily nitroso derivative $\text{C}_6\text{H}_5\text{CO C}_6\text{H}_4(\text{NO})\text{NMe}_2$ (E Bischoff, *B* 22, 340).

Di-methyl-di-*p*-amido-benzophenone *Di-benzoyl derivative* $\text{C}_6\text{H}_3\text{N}_2\text{O}$ i.e. $(\text{NMeBz C}_6\text{H}_4)_2\text{CO}$ [102°]. Formed by heating tetra methyl di amido benzophenone with BzCl at 190° (Nathansohn & Muller, *B* 22, 1877). Small light brown plates, *sl* sol cold, *v* sol hot, alcohol, *sl* sol benzene, almost insol water and ether. Acids and alkalis at 100° do not saponify it.

Tri-methyl-di-amido-benzophenone

$\text{NHMe C}_6\text{H}_4\text{CO C}_6\text{H}_4\text{NMe}_2$ [156°]. Formed, together with di methyl aniline, by boiling penta methyl-tri-amido tri phenyl-carbinol with HClAq (Wichelhaus, *B* 19, 109). Nodular groups of needles (from alcohol).

Tetra-methyl-di-amido-benzophenone

$\text{NMe}_2\text{C}_6\text{H}_4\text{CO C}_6\text{H}_4\text{NMe}_2$ [172°] (above 360°) (Græbe, *B* 20, 3262).

Formation—1 By passing COCl_2 into di methyl aniline (Michler, *B* 9, 716, 1900).—2 By boiling hexa methyl tri amido tri phenol carbinol with HClAq (Wichelhaus, *B* 19, 109).—3 From CCl_4 , SO_2Cl and di methyl aniline (Michler & Moro, *B* 12, 1168).

Preparation—By boiling auramine with aqueous HCl till decoloured, and ppg with NH_3 .

Properties—White plates (from dilute alcohol), insol water, *m* sol alcohol, *v* e sol warm benzene, *v* *sl* sol ether.

Reactions—1 Heating with ZnCl_2 and

NH_4Cl at 160° produces auramine (Fehrmann, *B* 20, 2844) — 2 Boiling HNO_3 (S G 1 48) gives insoluble yellow crystals of tetra nitro di methyl-di nitramido benzophenone and tri nitro phenyl-methyl nitramine [127°], soluble in alcohol (Romburgh, *R T C* 6, 867) — 3 Excess of bromine added to a solution of the base in glacial acetic acid gives the tetra-bromo derivative ($\text{C}_6\text{H}_2\text{Br}_4\text{NMe}_2$), CO [172°] which crystallises in slender yellow needles from alcohol (Nathansohn a Muller, *B* 22, 1883) — 4 Heating with aniline hydrochloride yields phenyl auramine — 5 Combines with tri nitro benzene (2 mols) forming a compound which crystallises in long reddish violet needles [c 100°] With half the quantity of tri nitro benzene (1 mol) it forms small deep violet plates [123°] — 6 Combines with *m*-di-nitro benzene (2 mols) forming a red crystalline compound — 7 Nitrous acid in the cold forms the nitroso- derivative $\text{NMe}_2\text{C}_6\text{H}_4\text{CO C}_6\text{H}_4(\text{NOH})\text{NMe}_2$ [159°] crystal lising from alcohol in golden plates This substance is a weak base, its solution in HCl aq being pptd by water It gives Liebermann's reaction Stannous chloride reduces it to the original tetra methyl di amido benzophenone The nitroso derivative forms the following salts $\text{B}^+\text{H}_2\text{Cl}_2$ Obtained by passing HCl into a solution of the nitroso ketone in benzene — $\text{B}^+\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OH}$ [152°] Orange needles (from alcohol) The nitroso ketone gives also a phenyl hydrazide $\text{C}_6\text{H}_5\text{NHNH C}_6\text{H}_4\text{N}_2\text{O}_2$ [148°] (E Bischoff, *B* 21, 2452, 22, 337) — 8 In presence of dehydrating agents, such as PCl_5 or AlCl_3 , it condenses with secondary and tertiary bases Thus with di methyl aniline it yields 'crystal violet' $\text{C}(\text{OH})(\text{C}_6\text{H}_5\text{NMe}_2)_3$.

Salts — $\text{B}^+\text{H Cl}_2$ small radially grouped prisms (from alcohol) Decomposed by water, with separation of the base (Fehrmann, *B* 20, 2844) — $\text{B}^+\text{H PtCl}_4$ yellow granules, insol water, v sl sol alcohol — Picrate $\text{B}^+\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OH}$ [157°] Small, purple, radially grouped, prisms (from alcohol), insol cold, v sl sol hot, water, m sol alcohol

Di methyl di iodide $\text{B}^+\text{Me}_2\text{I}_2$ [105°] Light yellow plates (from alcohol), sl sol cold, v sol hot, water and alcohol Split up at 150° into MeI and the base (Nathansohn a Muller, *B* 22, 1876)

Di methylo-di-hydroxide $\text{B}^+\text{Me}_2(\text{OH})_2$ From the preceding and moist AgO Small yellow plates (from alcohol), quickly becoming dark and resinous (N a M)

Oxime $\text{HON C}(\text{C}_6\text{H}_5\text{NMe}_2)_2$ [233°], colourless crystals (Munohmeyer, *B* 19, 1852, 20, 1852)

Phenyl-hydrazide $\text{C}_6\text{H}_5\text{NHNH C}_6\text{H}_4\text{N}_2\text{O}_2$ ($\text{NMe}_2\text{C}_6\text{H}_4$), CN HPh [175°] Needles (from benzene-alcohol), m sol warm alcohol, v sol ether Coloured green by acid oxidising agents (Ziegler, *B* 20, 1111) Conc H_2SO_4 gives a red colouration

Imide ($\text{NMe}_2\text{C}_6\text{H}_4$), C NH Auramine base [136°] Formed by heating tetra-methyl-di-amido benzophenone with NH_4Cl and ZnCl_2 at 150° to 160° Formed also by treating a solution of tetra methyl-di-amido benzophenone in CS_2 with PCl_5 and subsequently adding ammonia (Caro a Kern) (cf AURAMINE in THORPE'S DICTIONARY OF APPLIED CHEMISTRY). Lemon-yellow

plates, insol water and ether, m sol alcohol Dilute HCl aq readily converts it, even in the cold, into tetra methyl di amido benzophenone

Reactions — 1 Sodium amalgam reduces it (in alcoholic solution) to leucauramine ($\text{C}_6\text{H}_5\text{NMe}_2$), CH NH_2 [185°] This substance forms colourless crystals which when treated with HOAc yield an intense blue solution (Græbe, *B* 20, 3265) — 2 H_2S in alcoholic solution at 60° forms tetra methyl di amido thio benzophenone [164°] — 3 CS_2 also forms ($\text{NMe}_2\text{C}_6\text{H}_4$), CS together with thiocyanic acid (Fehrmann, *B* 20, 2847)

Salts — B^+HCl aq Auramine Yellow six sided tables (from water at 70°) After expulsion of the water it melts (G) or decomposes (F) at 267° M sol cold water, m sol alcohol Boiling water converts it into tetra methyl di-amido benzophenone Dyes wool and silk greenish yellow Cotton prepared with tannin is also dyed yellow by auramine — $\text{B}^+\text{H}_2\text{PtCl}_4$ granules, insol water, sl sol alcohol — B^+HI [268°] Minute columns — B^+HSCy aq [200° – 210°] — $\text{B}^+\text{H.C}_2\text{O}_4$ [194°] Orange needles, sl sol water, m sol warm alcohol — $\text{B}^+\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OH}$ [230° – 236°] Plates, insol cold, v sl sol hot, water, m sol hot alcohol

Phenyl imide ($\text{NMe}_2\text{C}_6\text{H}_4$), C NPh Formed by heating the hydrochloride of the imide (auramine) with aniline at 180° (F). Small greyish-yellow radially grouped needles (from alcohol), decomposing at 80° Insol water and ether — B^+HCl reddish crystalline mass, sol water and alcohol Its aqueous solution gradually decomposes, yielding aniline and tetra methyl di-amido benzophenone — $\text{B}^+\text{H}_2\text{PtCl}_4$ — $\text{B}^+\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OH}$

p Tolyli imide ($\text{NMe}_2\text{C}_6\text{H}_4$), $\text{C NC}_6\text{H}_4\text{Me}$ Resembles the phenyl imide — $\text{B}^+\text{H.PtCl}_4$

Ethylene-di-amide $\text{C}_6\text{H}_5\text{N}_2\text{Me}$

($\text{NMe}_2\text{C}_6\text{H}_4$), $\text{C} \begin{array}{c} \text{NH CH}_2 \\ | \\ \text{NH CH}_2 \end{array} \text{C}$ Formed by heating auramine (hydrochloride) with ethylene diamine at 100° – 110° (Fehrmann, *B* 20, 2855) Yellowish plates (from alcohol), insol water, m sol warm alcohol On treatment with HOAc or dilute HCl aq it decomposes into ethylene diamine and ($\text{NMe}_2\text{C}_6\text{H}_4$), CO In alcoholic solution it is decomposed by H_2S into ethylene diamine and ($\text{NMe}_2\text{C}_6\text{H}_4$), CS — $\text{B}^+\text{H}_2\text{PtCl}_4$ yellow needles — $\text{B}^+\text{H}_2\text{PtCl}_4$ amorphous flakes, insol water and ether, v sol warm alcohol — $\text{B}^+\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OH}$ flakes, v sl sol water, v sol alcohol.

Tolylene di amide

($\text{NMe}_2\text{C}_6\text{H}_4$), $\text{C} \begin{array}{c} \text{NH} \\ | \\ \text{NH} \end{array} \text{C}_6\text{H}_4\text{Me}$ Formed by heating the hydrochloride of the imide (auramine) with (1, 2, 4) tolylene diamine at 160° , and treating the product with ammonia Small brown scales In dilute acetic acid solution it dyes cotton, mordanted with tannin, reddish-brown Dilute HCl aq regenerates the original ketone — $\text{B}^+\text{H}_2\text{PtCl}_4$ — $\text{B}^+\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OH}$

Tetra-methyl-tri-amido-benzophenone

$\text{C}_6\text{H}_5\text{N}_3\text{O}_2$ + $\text{NMe}_2\text{C}_6\text{H}_4\text{CO C}_6\text{H}_4(\text{NH}_2)(\text{NMe}_2)$ [82°] Obtained by reducing nitro-tetra methyl-di amido benzophenone (Nathansohn a Muller, *B* 22, 1884) Amorphous yellow powder, v sol hot, sl sol cold, alcohol and benzene, insol ether — $\text{B}^+\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OH}$ minute plates, v sol

hot alcohol, insoluble in benzene and ether — $B'H_3PtCl_6$, light red crystalline pp, v sol warm alcohol, sl sol benzene, insol water and ether

METHYL- α -AMIDO- n -BUTYRIC ACID

$C_6H_{11}NO_2$, *i.e.* $CH_3CH_2CH(NHMe)CO_2H$. Formed by heating α -bromo-butyric acid with a concentrated solution of methylamine in sealed tubes at 100° . The product is boiled with baryta to expel NH_4Me , the Ba exactly ppd by H_2SO_4 , and the HBr by Ag_2CO_3 (Duvillier, *A Ch* [5] 20, 188, *C R* 83, 425). Glistening leaflets (from alcohol), subliming above 120° without fusion. V sol water, m sol hot alcohol, insol ether. Has a sweet taste. Feebly acid in reaction. By the prolonged action (several months) of cyanamide (1 mol) in concentrated and slightly ammoniacal solution it is converted into butyrocresatinine $CH_3CH_2CH<\frac{CO\ NH}{N(CH_3)}>C\ NH$

which crystallises from alcohol in slender silky needles composed of small rectangular plates (Duvillier, *C R* 95, 456, *Bl* [2] 39, 539).

Salts — $HA'HCl$ [150°]. Badly defined crystals, sol alcohol, insol ether. Gives off HCl on fusion — $H_2A'H_2PtCl_6$, orange-red crystals, v sol water and alcohol, insol ether — $H_2A'H_2PtCl_6$, 5 aq. Formed at 0° — $HA'HAuCl_4$, aq. transparent yellow prisms, sol water, alcohol, and ether — The nitrate and sulphate crystallise with difficulty in long needles — $CuA', 2aq$, light-blue prisms, sol alcohol.

Di methyl α -amido-butyric acid *Anhydride of the methylo-hydroxide* $C_6H_{11}NO_2$, CH_3CH_2CH-CO

i.e. $(CH_3)_2N-O$ **Tri-methyl- α -amido-butyro-betaïne** Prepared by the action of bromo butyric ether on an excess of tri methylamine in alcoholic solution (Duvillier, *C R* 104, 1520). Large transparent crystals, containing aq, which they lose at 120° , becoming opaque, v sol. water and alcohol, insol ether. Has a bitter taste — $B'HCl$. Uncrystallisable — $B'H_3PtCl_6$, orange prisms, sl sol alcohol. The aurochloride forms yellowish-white crystals.

METHYL AMIDO-CAPROIC ACID *v* **METHYL-AMIDO HEXOIC ACID**

METHYL-AMIDO-CHLORO- *v* **CHLORO-METHYL-AMIDO**

METHYL-AMIDO-CYANURIC ACID *v* **CYANIC ACID**.

METHYL-AMIDO-ETHANE SULPHONIC ACID $C_6H_{11}NSO_3$, *i.e.* $CH_3NHCH_2CH_2SO_3H$ *Methyl-taurine* [242].

Formation — From methyl-ethylene- ψ thio-urea and bromine-water (Gabriel, *B* 22, 1148).

Preparation — Silver chloro-ethane sulphonate is heated with 8 times its weight of a solution of methylamine, saturated at 0° , for 5 hours at 120° . The product is boiled with baryta to expel excess of methylamine, freed from excess of baryta by H_2SO_4 , and evaporated to crystallisation. The crystals are washed with alcohol and recrystallised from water. The yield is 25 p.c. (E. Dittrich, *J pr* [2] 18, 66).

Properties — Transparent triclinic crystals. Sol water, insol alcohol and ether. Its aqueous solution is acid to litmus. Crystallises un-

altered from strong HCl. Does not form salts with acids and alkalis.

Reactions — 1 N_2O_5 forms isethionic acid, $(HO)CH_2CH_2SO_3H$ — 2 With cyanamide it combines to methyl taurocyamine or methyl guanido-ethane sulphonic acid.

Di methyl amido ethane sulphonic acid $NMe_2CH_2CH_2SO_3H$. Formed by heating di methylamine β chloro ethane sulphonate (20 g) with aqueous (53 p.c.) dimethylamine (5 g) at 160° for 10 hours (James, *J pr* [2] 31, 416, *C J* 47, 370). Large tabls (from water), v sol water, insol ether. Decomposes at 270° — 280° without fusion. Does not combine with HCl.

Anhydride of the methylo hydroxide $<\frac{CH_2NMe_2}{CH_2SO_3}>O$. Formed by heating β chloro-ethane sulphonic acid with aqueous tri methylamine. Slender prisms (from water), v sol water, insol alcohol and ether. Neutral in reaction. Tastes sweet. Not decomposed at 300° . Not acted upon by cyanamide. Baryta gives NMe_2 and isethionic acid (James, *C J* 49, 489).

DI-METHYL-AMIDO-ETHYL ALCOHOL *v* **DI-METHYL-OXYETHYL AMINE**

DI-METHYL AMIDO ETHYL BENZENE *v*

DI-METHYL AMIDO PHENYL ETHANE

METHYL-AMIDO-ETHYL KETONE

$CH_3COCH(NH_2)CH_3$. Perhaps formed by reducing methyl nitroso ethyl ketone, but if so it quickly changes to *s*-tetra methyl pyrazine.

METHYL-AMIDO-FORMIC ACID *v* **METHYL-CARBAMIC ACID**

METHYL- α -AMIDO n HEXOIC ACID

$C_6H_{11}NO_2$, *i.e.* $CH_3CH_2CH_2CH_2CH(NHMe)CO_2H$ *Methyl-amido caproic acid* S 102 at 11° . Formed by heating α -bromo-hexoic acid (1 mol) with aqueous methylamine (2 or 3 mols) at 100° for several hours (Duvillier, *C R* 90, 822, *A Ch* [5] 29, 165). Silky needles (from water) or pearly plates (from alcohol). Sl sol cold alcohol, insol ether. Neutral in reaction. Volatilises above 100° . Does not reduce silver or mercurous nitrate. Gives with ferric chloride an intense red colouration and a yellowish brown pp. An equivalent quantity of cyanamide in cold concentrated solution containing a few drops of NH_3 forms in a few weeks crystals of 'hexoic

creatinine' $C_6H_7CH<\frac{NMe\ C\ NH}{CO\ NH}>$ Sl sol.

cold water, v sol alcohol (Duvillier, *C R* 96, 1583, *Bl* [2] 40, 307) — $HA'HCl$ transparent plates, v sol water and alcohol, insol ether — $H_2A'H_2PtCl_6$, orange crystals, v sol water, v sol alcohol, v sl sol ether — The aurochloride forms golden needles, the sulphate forms very indistinct crystals — $CuA', 2aq$, blue scales, S 1.

Di methyl amido-hexoic acid *Methylo-hydroxide* $HO\ NMe_2\ C_6H_{11}\ CO_2H$. Formed by heating leucine (1 mol) derived from casein with aqueous KOH (3 mols) and MeI (3 mols). The resulting $INMe_2\ C_6H_{11}\ CO_2K$, which crystallises from alcohol in needles, being decomposed by moist Ag_2O (Körner a Menozzi, *G* 13, 359). Strongly alkaline. Decomposes at 120° — 130° into trimethylamine, an acid, $C_6H_{11}O_2$, and oxy-hexoic acid. Forms the platinochloride

(CINMe_2 , C_6H_5 , CO_2H). PtCl_2 , aq., aurochloride $\text{Cl}_2\text{AuNMe}_2$, C_6H_5 , CO_2H [163°], and periodide. The periodide forms lustrous green crystals, decomposed by H_2S , yielding INMe_2 , C_6H_5 , CO_2H [191°], crystallising in small prisms

DI-METHYL-AMIDO-HYDROQUINONE

$\text{C}_6\text{H}_3(\text{NMe}_2)(\text{OH})_2$ [2 4 1] *Di methyl derivative* $\text{C}_6\text{H}_3(\text{NMe}_2)(\text{OMe})$

Methylo iodide B'MeI [202°] Formed by heating the di methyl derivative of amido hydroquinone with methyl iodide (Bässler, *B* 17, 2122) White needles, v sol water, sl sol absolute alcohol, nearly insol benzene, chloro form, and ligroine

Methylo hydroxide B'MeOH soluble needles

Methylo chloride B'MeCl white easily soluble needles [172°] — (B'MeCl), PtCl_2 , yellow crystalline pp

TRI METHYL-AMIDO IMIDO IMIDO DI-PHENYL SULPHIDE

Methylo chloride $\text{C}_6\text{H}_3\text{N}_3\text{S}_2\text{Cl}_2$ $\text{N} \begin{array}{c} \text{C}_6\text{H}_3(\text{NMe}) \\ \text{C}_6\text{H}_3 \end{array} \text{S} \text{Methylene}$
 NMe_2Cl

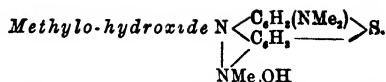
blue Chloride of tetra methyl thionine

Formation — 1 By the action of H_2S on an acid solution of nitroso di methyl aniline and oxidation of the resulting leuco methylene blue. In this reaction di methyl *p* phenylene diamine is an intermediate product (Caro, *E P* 1877, 3751, Koch, *B* 12, 593, Bernthsen, *A* 230, 137) — 2 By dissolving nitroso di methyl aniline in H_2SO_4 (S G 14), treating with sulphide of zinc, and oxidising the resulting leuco methylene blue (Oehler, *G P* 1882, 24,125, Muhliauser, *D P J* 262, 371) — 3 By oxidation of di methyl *p* phenylene diamine in presence of $\text{Na}_2\text{S}_2\text{O}_8$, and further oxidation of the resulting $\text{C}_6\text{H}_3(\text{NMe}_2)(\text{NH}_2)\text{S SO}_3\text{H}$ by chromic acid (Bernthsen) — 4 By reducing the compound $\text{C}_6\text{H}_3(\text{NMe}_2)(\text{NH}_2)\text{S SO}_3\text{H}$ to $\text{C}_6\text{H}_3(\text{NMe}_2)(\text{NH}_2)\text{SH}$ and oxidising a mixture of this mecaptan with di methyl aniline with HCl and $\text{K}_2\text{Cr}_2\text{O}_7$, the resulting 'soluble green' $\text{C}_6\text{H}_3\text{N}_3\text{S}$ or $\text{N} \begin{array}{c} \text{C}_6\text{H}_3(\text{NMe}_2) \\ \text{C}_6\text{H}_3 \end{array} \text{S}$ slowly changing in aqueous

solution to methylene blue (Bernthsen, *A* 251, 10)

Properties — Minute dark blue laminae (containing 3aq), v sol water and alcohol. Dyes mordanted cotton blue. Its aqueous solution is blue and not affected by HCl or ammonia. NaOH gives a violet colour and, when added in large quantity, a dirty violet pp. Conc H_2SO_4 gives a yellowish green solution, becoming blue on dilution. The aqueous solution is decolourised by H_2S or $\text{Na}_2\text{S}_2\text{O}_3$, the colour is restored even by feeble oxidising agents. Sodium hyposulphite $\text{Na}_2\text{S}_2\text{O}_4$ reduces it to the leuco base. Fuming HCl at 260° decomposes it, giving off H_2S . Boiling dilute HNO_3 oxidises it, forming H_2SO_4 . Boiling aqueous KOH forms 'thionol', the anhydride of di-oxo-imido-di phenyl sulphide $\text{N} \begin{array}{c} \text{C}_6\text{H}_3(\text{OH}) \\ \text{C}_6\text{H}_3 \end{array} \text{S}$

Salt — ($\text{C}_6\text{H}_3\text{N}_3\text{S}_2\text{Cl}_2$), ZnCl_2 , aq.



Formed by treating methylene blue with moist Ag_2O . Amorphous, dark blue mass, v e sol. water and alcohol, insol ether

Methylo iodide $\text{C}_6\text{H}_3\text{N}_3\text{SI}$ bronzy needles (from hot water). The di chloro derivative of this iodide $\text{C}_6\text{H}_3\text{Cl}_2\text{N}_3\text{SI}$ is formed by successive treatment of di chloro di methyl *p* phenylene diamine $\text{C}_6\text{H}_3\text{Cl}_2(\text{NMe}_2)(\text{NH}_2)$ with H_2S , FeCl_3 , and KI (Mohlau, *B* 19, 2012)

TRI METHYL-AMIDO IMIDO IMIDO DI-PHENYL-SULPHONE

Methylo-iodide $\text{C}_6\text{H}_3\text{N}_3\text{SO}_2\text{I}$ $\text{N} \begin{array}{c} \text{C}_6\text{H}_3(\text{NMe}_2) \\ \text{C}_6\text{H}_3 \end{array} \text{SO}_2 \text{Methyl-}$
 NMe_2I

ene azure Formed, together with methylene violet, by boiling methylene blue with Ag_2O and a large quantity of water (Bernthsen, *A* 230, 175). Slender needles with green lustre, sol water, forming a violet solution from which it is pptd by KI . Alkalis destroy the colour of the solution. The corresponding $\text{C}_6\text{H}_3\text{N}_3\text{SO}_2\text{Cl}$ crystallises in needles, v e sol water. It dyes silk like methylene blue

TETRA METHYL-DI AMIDO-IMIDO DI-PHENYL METHANE v Imide of Tetra-METHYL-DI AMIDO-BENZOPHENONE

METHYL-DI-AMIDO IMIDO DI-PHENYL SULPHIDE $\text{C}_6\text{H}_3\text{N}_3\text{S}$ $\text{NMe} \begin{array}{c} \text{C}_6\text{H}_3(\text{NH}_2) \\ \text{C}_6\text{H}_3 \end{array} \text{S}$

Methyl di amido thio diphenylamine Formed by reduction of di nitro methyl imido-di phenyl sulphide with tin and HCl (Bernthsen, *A* 230, 130). Insol water, sol ether. The dilute solution of its hydrochloride is coloured blue by FeCl_3 — $\text{B'H}_4\text{Cl}_2$. Needles, v sol water, almost insol HCl aq

Tetra methyl-di-amido-imido-di phenyl sulphide $\text{C}_6\text{H}_3\text{N}_3\text{S}$ $\text{NH} \begin{array}{c} \text{C}_6\text{H}_3(\text{NMe}_2) \\ \text{C}_6\text{H}_3 \end{array} \text{S}$

Leuco methylene blue Formed by treating methylene blue with sodium hyposulphite $\text{Na}_2\text{S}_2\text{O}_4$ (Bernthsen, *A* 230, 147). Needles (from alcohol), sl sol water, m sol cold alcohol. Gives an acetyl derivative, v sol alcohol, ether and benzene. With MeI it forms $\text{NMe} \begin{array}{c} \text{C}_6\text{H}_3(\text{NMe}_2\text{I}) \\ \text{C}_6\text{H}_3 \end{array} \text{S}$ crystallising in plates, v sl sol water and alcohol, insol ether. Oxidised by air to methylene blue — $\text{B'H}_4\text{ZnCl}_2$, prisms

Di-methyl amido imido phenyl disulphide *Methylo chloride* $\text{C}_6\text{H}_3\text{N}_3\text{S}_2\text{Cl}_2$ S

$\text{NMe}_2\text{Cl} \begin{array}{c} \text{C}_6\text{H}_3 \\ \text{N-S} \end{array} \text{S} \text{Methylene red}$ A by product in the manufacture of methylene-blue from di methyl *p* phenylene diamine and H_2S followed by FeCl_3 (Bernthsen, *A* 230, 165, 251, 22, Koch, *B* 12, 594). Small green glittering prisms (from alcohol), v e. sol. water, insol ether. Its aqueous and alcoholic solutions are purple. Not affected by HCl . Alkalis decompose it — Zinc salt ($\text{C}_6\text{H}_3\text{N}_3\text{S}_2\text{Cl}_2$), ZnCl_2 , aq

DI-METHYL-AMIDO-JUGLONE v. JUGLONE.

METHYL-AMIDO-MESITYLENE v. METHYL-MESIDINE.

METHYL AMIDO METHYL-BENZENES v.
METHYL-PHENYL METHYL AMINES

DI-METHYL-AMIDO-METHYL-PHEN.

AZINE $C_6H_5(NMe_2) \begin{smallmatrix} \diagup N \diagdown \\ | \\ C_6H_5Me \end{smallmatrix}$ Formed by

elimination of the NH group from 'tolylene red' by means of nitrous acid and alcohol (Bernthsen & Schweitzer, *B* 19, 2605) Red greenish glistering needles or flat prisms Has basic properties Dissolves in dilute acids with a violet colour, in conc H_2SO_4 with a reddish brown, which on dilution first becomes green, then blue, and finally violet Soluble in ether with a yellowish-red colour and yellow fluorescence Sublimable

Di methyl di-amido methyl phenazine

$C_{13}H_{16}N_4$ v e $C_6H_5(NMe) \begin{smallmatrix} \diagup N \diagdown \\ | \\ C_6H_5Me(NH_2) \end{smallmatrix}$

Tolylene red Formed by oxidation of nitroso-amido dimethyl aniline together with tolylene *m* diamine The leuco base has the formula $C_6H_5(NMe_2) \begin{smallmatrix} \diagup NH \diagdown \\ | \\ C_6H_5Me(NH) \end{smallmatrix}$ (Bernthsen & Schweitzer, *B* 19, 2604)

DI METHYL AMIDO METHYL-QUINOLINE TETRAHYDRIDE *Di-methylo di-iodide* $C_8H_8Me_2N NMe_2Me_2I_2$ [171°] Formed by heating (*Py* 1) amido quinoline tetrahydride with methyl alcohol and MeI (Ziegler, *B* 21, 862) Crystals

METHYL-AMIDO METHYL-THIAZOLE

$C_5H_5N_2S$ v e $\begin{smallmatrix} S C(NHMe) \\ | \\ CH=CMe \end{smallmatrix} \begin{smallmatrix} \diagup N \diagdown \end{smallmatrix}$ *Methyl thiazyl-*

amine [42°] Formed by the action of chloroacetone on methyl thio urea (Traumann, *A* 249, 44) Crystalline but extremely hygroscopic, *m* sol ether Strongly alkaline in reaction When heated with $HClAq$ in a sealed tube it yields methylamine Br completely decomposes it

Salts— B^+H^- small white needles, melts, when anhydrous, at 136° (*T*)—Platinochloride orange yellow plates [167°]

Acetyl derivative $C_5H_5AcN_2S$. [110°] White needles

Methyl-amido-methyl thiazole

$S C(NH) \begin{smallmatrix} \diagup NMe \diagdown \\ | \\ CH CMe \end{smallmatrix}$ *Imido di methyl thiazole*

[47.5°] Formed from chloro acetone by treatment with ammonium sulpho cyanide (Tehernia & Norton, *B* 16, 345) and heating with MeI the resulting amido methyl thiazole (so-called propinine sulphocyanide) (Hantzsch & Weber, *B* 20, 3122, 3336) Crystalline, but very hygroscopic Strongly alkaline When heated with $HClAq$ in sealed tubes it yields ammonia Conc $KOHAq$ has no action

Salts— B^+H^- Tables [164°]—Platinochloride orange tables [193°]

Acetyl derivative $C_5H_5AcN_2S$ [113°] White needles (containing 6 aq)

Di-methyl-amido-methyl-thiazole $C_5H_{10}N_2S$ $S \cdot C(NMe) \begin{smallmatrix} \diagup NMe \diagdown \end{smallmatrix}$

v e $\begin{smallmatrix} CH=CMe \end{smallmatrix} \begin{smallmatrix} \diagup NMe \diagdown \end{smallmatrix}$ *Methyl imido dimethyl-thiazole* [96°] Formed by adding conc $KOHAq$ to its hydro-iodide (Hantzsch & Weber, *B* 20, 3123) White needles, sol alcohol, water, and

ether Bromine reacts with formation of $C_5MeBr(NMe_2)SN$ [114°]

Salts— B^+H^- [54°] From MeI and $C_5MeH(NHMe)SN$ Melts at 155° when anhydrous

Methylo-iodide B^+MeI [85°] White needles, v e sol water and alcohol

DI-METHYL-AMIDO-NAPHTHOIC ACID $C_{10}H_7(NMe_2)(CO_2H)$ [14] [165°] Formed by heating di methyl (a) naphthylamine with $COCl_2$ at 70° for 4 hours (Friedlander, *B* 21, 3126) Needles (from dilute alcohol), sol dilute acids and alkalis When acted upon by diazo compounds the azo-group displaces the COH —(HA^+) H^+PtCl_6 yellow needles

DI METHYL AMIDO-NAPHTHO-PHEN

AZINE $C_{10}H_7 \begin{smallmatrix} \diagup N \diagdown \\ | \\ C_6H_5 NMe_2 \end{smallmatrix}$ *Di methyl*

naphth eurhodine [205°] Formed from nitroso di methyl aniline hydrochloride, (β) naphthylamine, and $AcHO$ (Witt, *B* 21, 720) Rhombic tables (from toluene or xylene), red by transmitted, and red or green, according to the faces, by reflected light, sol alcohol, ether and benzene, forming yellow solutions with yellow fluorescence Is volatile with slight decomposition, and sublimes readily in woolly flocks The violet red solution in conc H_2SO_4 becomes successively black, green, grey, and blue violet on dilution The salts crystallise readily, have a bronzy lustre, and are dissociated by water $AcHO$ dissolves it with red violet colour Conc HNO_3 with violet, soon becoming bright yellow and depositing the nitro compound

METHYL-AMIDO NAPHTHOQUINONE

$C_{11}H_9NO_2$ v e $C_{11}H_9O_2(NHMe)$ [232°] Formed by adding a solution of methylamine acetate to an alcoholic solution of (a) naphthoquinone, evaporating nearly to dryness, adding water, and crystallising the pp from alcohol (Plimpton, *C J* 37, 639) Glittering red needles, v sol alcohol Aqueous SO_2 at 150° forms an unstable colourless reduction product

Di-methyl-amido-naphthoquinone

$C_{10}H_7O_2(NMe_2)$ [118°] From (a) naphthoquinone and di methyl amine in alcoholic solution (Plimpton) Red needles

DI-METHYL AMIDO-NAPHTHYLAMINE v

NAPHTHYLENE DI METHYL DIAMINE

TETRA-METHYL-DI-AMIDO-DI NAPHTHYL PHENYL-METHANE

$Ph CH(C_6H_4NMe_2)_2$ [189°] Formed by the action of benzoic aldehyde on di methyl (a) naphthylamine in presence of $ZnCl_2$ (Friedlander, *B* 21, 3128) Colourless crystals, v sol $HOAc$, benzene, CS_2 , and dilute mineral acids, sl sol alcohol, ether Does not yield a colour on oxidation

Hexa methyl tri amido-di-naphthyl phenyl methane $HC(C_6H_4NMe_2)_3C_6H_4NMe_2$ [179°] Formed by condensing di-methyl *p* amido benzoic aldehyde with di methyl (a) naphthylamine in presence of $ZnCl_2$ (Friedlander, *B* 21, 3129) White needles Does not yield a colour on oxidation

METHYL-AMIDO-NITRO- compounds v **NI TRO-METHYL-AMIDO-** compounds

METHYL-AMIDO-OXY- compounds v **OXY METHYL-AMIDO-** compounds

METHYL-AMIDO PEREZONE *v.* **METHYL-**

AMIDO PIPITZARIC ACID

METHYL o AMIDO-PHENOL *Methyl derivative* C_6H_4NO *is* $C_6H_4(NHMe)(OMe)$ [1 2]*Methyl anisidine* (219°) Formed by mixing the methyl derivative of *o* amido phenol with MeI at 0° (Mühlhauser, A 207, 247) Oil — B'H PtCl₅ short yellow prisms, *m* sol water**Methyl-p amido phenol** *Ethyl derivative* C_6H_4NO *is* $C_6H_4(NHMe)(OEt)$ [1 4] (251°) Formed by heating the ethyl derivative of *p* oxy phenyl amido acetic acid at 260° (Bischoff & Nastvogel, B 22, 1789) Sl sol water, *v* sol alcohol and ether — B'HCl needles**Di methyl o amido phenol** C_6H_3NO *is* $C_6H_3(NMe_2)(OH)$ [1 2] [45°] Obtained, together with MeCl, by the dry distillation of the methyl chloride (Griess, B 13, 248) Small white prisms, *v* sl sol hot water, *v* sol alcohol, ether, HOAc, and aqueous KOH FeCl₃ gives a reddish violet colour Its hydrochloride is gummy**Methylo hydroxide** $C_6H_4(NMe_2OH)(OH)$ or $C_6H_4\begin{smallmatrix} O \\ | \\ NMe_2 \end{smallmatrix}$ aq Prepared by the action ofMeI and KOH on a solution of *o* amido phenol in methyl alcohol (Griess, B 13, 246) Prisms,which become $C_6H_4\begin{smallmatrix} O \\ | \\ NMe \end{smallmatrix}$ at 105° *V* solwater and alcohol, *insol* ether Has an intensely bitter taste On distillation it is converted into $C_6H_4(NMe)(OMe)$ With acids it forms the following salts — $C_6H_4(NMe_2I)(OH)$ aq white soluble prisms Its solution, neutralised by ammonia, deposits sparingly soluble needles orprisms of $C_6H_4(NMe_2I)(OH)C_6H_4\begin{smallmatrix} O \\ | \\ NMe_2 \end{smallmatrix}$ — $C_6H_4(NMe_2Cl)(OH)$ 2aq long soluble prisms — $C_6H_4(NMe_2Cl)(OH)$ PtCl₅ yellowish red needles, sl sol cold water — The periodide forms brown insoluble leaflets — The nitroprusside (C_6H_3NO), H FeCy₃NO forms crystals, sl sol cold water**Methyl derivative** C_6H_3NO *is* $C_6H_3(NMe)(OMe)$ [1 2] (211°) SG 2³ 1 016 Formed by an isomeric change by distilling $C_6H_4\begin{smallmatrix} O \\ | \\ NMe \end{smallmatrix}$ (Griess, B 13, 248) Formed alsofrom MeI and the methyl derivative of *o* amido-phenol (Mühlhauser, A 207, 248) Colourless liquid, with burning taste — B'H₂PtCl₅ sparingly soluble golden yellow prisms**Methylo iodide** of the *methyl derivative* $C_6H_3(NMe_2I)(OMe)$ From the preceding and MeI Long white needles (G) or tables (M), sol hot water and hot alcohol With moist Ag₂O it yields a strongly alkaline hydroxide It also yields a platinumchloride ($C_6H_3(NMe_2Cl)(OMe)$ PtCl₅), which crystallises in sparingly soluble yellow plates or tables**Di-methyl-m-amido-phenol** $C_6H_4(OH)NMe_2$ Prepared by fusing di methyl amido benzene *m* sulphonic acid with KOH Also by heating resorcin with di methyl amine under pressure**Ethyl ether** $C_6H_4(OEt)(NMe_2)$ [1 3] (247°) Obtained by boiling a solution of *m* amido-phenol and methyl iodide and slowly runningin the calculated quantity of potash The base is distilled over with steam (P Wagner, J pr [2] 32, 77, Baur & Stadel, B 16, 32) Converted by HCl and amyl nitrite into the nitroso compound $C_6H_3(NO)(OEt)(NMe)$ **Di-methyl-p-amido-phenol** *Methylohydroxide* The anhydride $C_6H_4\begin{smallmatrix} O \\ | \\ NMe_2 \end{smallmatrix}$ *is* formed by the action of MeI and conc KOHAq upon *p* amido phenol in the cold (Griess, B 13, 250) Prisms or plates Changes on distillation into the isomeric $C_6H_4(NMe_2)(OMe)$ [48°]**Methyl derivative** $C_6H_3(NMe)(OMe)$ [1 4] [49°] Formed as above (Griess, B 13, 249) Prisms or plates (from alcohol)**Methylo iodide** $C_6H_3(NMe_2I)(OMe)$ Formed by the action of MeI on either the methyl derivative or the methylo hydroxide Tables or plates With moist Ag₂O it yields the methylo hydroxide as a strongly alkaline mass. The platinumchloride ($C_6H_3(NMe_2Cl)(OMe)$ PtCl₅), forms small yellow prisms, sl sol water**Di-methyl-di-amido-phenol** *Anhydride* of the *methylo hydroxide* $C_6H_3N_2O$ *is* $C_6H_3(NH_2)\begin{smallmatrix} O \\ | \\ NMe_2 \end{smallmatrix}$ [4 1] [4 2] Prepared by reductionof the methylo hydroxide of nitro di methyl amido phenol with tin and HCl (Griess, B 13, 648) — B'H₂Cl₂ 4aq very soluble white plates — B'H PtCl₅ 2aq small prisms, sl sol water**TETRA-METHYL-DI-p-AMIDO-DIPHENYL** $NMe_2.C_6H_4.C_6H_4.NMe_2$ *Tetra methyl benzidine* [195°] (above 360°)**Formation** — 1 By heating di methyl aniline (1 pt) with H₂SO₄ (4 pts) at 200° Formed also in small quantity by oxidising di methyl aniline by boiling with PbO₂ and dilute H₂SO₄ (Miehler & Pattinson, B 14, 2161, 17, 115) — 2 By heating di methyl aniline with AlCl₃ in presence of air (Gnaud, Bl [3] 1, 692) — 3 By methylation of benzidine**Properties** — Colourless needles, sol hot, sl sol cold, alcohol Not volatile with steam Gives a green colouration with FeCl₃ or CrO₃ — B'H₂Cl, sparingly soluble needles — B'H₂Br, needles — B'H₂L white needles — B'H PtCl₅**Methylo iodide** B'MeI [263°] Needles, sl sol water and alcohol Loses MeI when distilled with soda lime**Methylo chloride** B'MeCl [228°] Crystals, very soluble in water and alcohol — B'MeClHPtCl₅ yellow pp**Tetra methyl op di amido diphenyl** (2 1) $NMe_2.C_6H_3.C_6H_3.NMe_2$ [1 4] *Tetra methyl diphenylamine* [52°] (333°-345°) Formed by heating diphenylamine hydrochloride with MeOH in sealed tubes at 180° (Reuland, B 22, 3015) Prisms Gives with platinum chloride an unstable pp Chloranil produces a blue colouration — B'C₆H₃(NO₂)₂OH [200°] Red needles**Methylo iodide** B'MeI [184°] Needles, *v* sol water, alcohol, and ether**Di methylo di-iodide** B'Me₂I₂ [196°] Crystalline, *v* sol water and alcohol**Tetra-methyl tetra-amido diphenyl** $NMe_2.C_6H_3(NH_2).C_6H_3(NH_2).NMe_2$ *Tetra-methyl benzidine* [168°] Prepared by reduction of di-

nitro tetra methyl diphenyl (Michler & Pattinson, *B* 14, 2165, 17, 118) White silvery plates, ν sol hot alcohol, sl sol cold alcohol, insol water. FeCl_3 gives a violet colouration. $\text{K}_2\text{Cr}_2\text{O}_7$ and H_2SO_4 produce a brownish red colour— $\text{B}^*\text{H}_2\text{Cl}_2$ (dried at 110°) sparingly soluble colourless needles— $\text{B}^*\text{H}_2\text{I}_2$ sparingly soluble needles— $\text{B}^*\text{H}_2\text{PtCl}_6$ yellow pp

Methyl α amido phenyl acetic acid $\text{C}_6\text{H}_4\text{NH}_2\text{CO}_2\text{H}$ $\text{C}_6\text{H}_4\text{CH}(\text{NHMe})\text{CO}_2\text{H}$ Formed from the nitrile of mandelic acid $\text{C}_6\text{H}_4\text{CH}(\text{OH})\text{CN}$ by digesting with alcoholic NH_4Me at 70° , and decomposing the resulting nitrile with HCl (Tiemann & Piest, *B* 14, 1982) Slender needles (from hot water) Sublimes at 274° Sl sol cold water, insol alcohol and ether

Amide $\text{C}_6\text{H}_4\text{CH}(\text{NHMe})\text{CONH}_2$ [155°] Slender needles— B^*HCl Needles, sol alcohol, insol ether

DI-METHYL-AMIDO-PHENYL- ω -AMIDO-CRESOL *Methyl derivative*

[4 1] $\text{NMe}_2\text{C}_6\text{H}_4\text{NHCH}_2\text{C}_6\text{H}_4\text{OMe}$ [1 4] [104°] Formed by reducing $\text{NMe}_2\text{C}_6\text{H}_4\text{NCH}_2\text{C}_6\text{H}_4\text{OMe}$ with sodium (Steinhart, *A* 241, 343) Light green plates, ν sol acids, forming red solutions. Its alcoholic solution decomposes rapidly

DI-METHYL-AMIDO-DI-PHENYL-AMINE $\text{Me}_2\text{NCH}_2\text{C}_6\text{H}_4\text{NHPH}$ [130°] One of the products formed by the action of phenyl hydrazine on nitroso dimethylamine in an alcoholic solution (O Fischer, *B* 21, 2612) White needles (from petroleum-ether), ν sol dilute HCl , m sol dilute SO_2 . Gives a blue colouration with FeCl_3 . Dissolves with a red colour in nitric acid

Nitrosamine $\text{C}_6\text{H}_4\text{N}_2\text{O}$ [116°] Yellow needles (from alcohol)

Tetra-methyl-di-amido-diphenyl-amine $(\text{NMe}_2\text{C}_6\text{H}_4)_2\text{NH}$ [119°] Obtained by oxidising a mixture of di methyl aniline (1 mol) and di methyl *p* phenylene diamine (1 mol) and reducing the resulting 'dimethyl-phenylene green' (Bundscheidler, *B* 16, 864) Yellowish dimetric tables

Hexa-methyl-tri amido triphenylamine *Tri-methylo trichloride* $(\text{NMe}_2\text{ClC}_6\text{H}_4)_3\text{N}$ Obtained by heating tri amido tri phenyl amine hydrochloride with MeOH at 190° (Heydrich, *B* 19, 758) White needles— $(\text{NMe}_2\text{ClC}_6\text{H}_4)_3\text{N} \cdot 3\text{PtCl}_6$

DI-METHYL AMIDO-PHENYL-BENZYL-AMINE $\text{NMe}_2\text{C}_6\text{H}_4\text{NHCH}_2\text{Ph}$ [48°] Obtained by reducing benzylidene di-methyl phenylene diamine [101°] with sodium amalgam (Kohler, *A* 241, 861) Yellowish plates, ν sol dilute mineral acids, alcohol, ether, benzene, and petroleum ether

Nitrosamine $\text{NMe}_2\text{C}_6\text{H}_4\text{N(NO)CH}_2\text{Ph}$ [128°] Slender yellow needles, sol alcohol

DI-METHYL-AMIDO-DI-PHENYL-CARBINOL $\text{NMe}_2\text{C}_6\text{H}_4\text{CH(OH)C}_6\text{H}_4$ *Di-methyl-amido di-phenyl carbinol* [70°] Formed by reducing di methyl amido benzophenone with sodium amalgam, or by the action of benzoic aldehyde on di methyl aniline (Albrecht, *B* 21, 3292) Thin white needles, insol water, ν e sol ordinary solvents, sl sol petroleum ether

Di-methyl-di-amido di-phenyl-carbinol $\text{NMe}_2\text{C}_6\text{H}_4\text{CH(OH)C}_6\text{H}_4\text{NH}_2$ [165°] Formed by carefully reducing ν nitro di methyl amido di-phenyl-carbinol with zinc dust and HCl (Al-

brecht, *B* 21, 3295) Dissolves in HOAc with blue colouration. Crystallises from benzene in needles containing benzene and melting at 142° . Gives off water (1 mol) when heated above its melting point. Boiling with zinc dust and HCl reduces it to di methyl diamido di phenyl methane [98°]

Tetra methyl di amido-di phenyl carbinol $\text{C}_6\text{H}_4\text{N}_2\text{O} \cdot 2 (\text{NMe}_2\text{C}_6\text{H}_4)_2\text{CH(OH)}$ [96°] Obtained by reducing tetra methyl di amido benzophenone in hot alcoholic solution with sodium amalgam (Michler & Dupertuis, *B* 9, 1899, Nathansohn & Muller, *B* 22, 1879) Colourless trichmic prisms, ν sol alcohol, HOAc , benzene, and ether. Its solution in HOAc is blue, the benzene solution is colourless

Salts— B^*HCl Formed by passing HCl into a solution of the base in ether. Small colourless slender radially grouped needles. In air it turns blue and deliquesces. It is dissociated by water— $\text{B}^*\text{H}_2\text{PtCl}_6$ minute yellow needles, ν sol hot alcohol— $\text{B}^*\text{C}_6\text{H}_5(\text{NO}_2)_2\text{OH}$ dark green crystalline mass, ν sol hot alcohol, sl sol benzene, insol ether

Di-methylo di-iodide $\text{B}^*\text{Me}_2\text{I}_2$ [195°] Small plates (from alcohol), sl sol cold, ν sol hot, alcohol and water, insol benzene and ether

Tetra-methyl di amido-tri-phenyl-carbinol $\text{C}_6\text{H}_4\text{N}_2\text{O} \cdot 2 (\text{C}_6\text{H}_4\text{C(OH)(C}_6\text{H}_4\text{NMe}_2)_2$ [132°] *Malachite green* *Benzaldehyde green*

Formation—1 By the action of dimethyl aniline on benzotrichloride in presence of a metallic chloride (Doebner, *B* 11, 1238, 13, 2222)—2 By the oxidation of a slightly acid solution of tetra methyl di amido tri phenyl-methane with MnO or PbO_2 (E & O Fischer, *B* 12, 796), or with tetra chloro quinone (O Fischer, *A* 206, 130)—3 By heating di methyl aniline (4 pts) with BzCl (2 pts) and ZnCl_2 (3 pts) (Fischer)

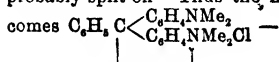
Preparation—1 From di methyl aniline (2 mols), ZnCl_2 (half its weight), sand, and benzo trichloride at 100° . The product is distilled with steam and the dye ppd from the aqueous residue by NaCl . The pp is the zinc double chloride, which may be converted by KOH into the base. This is converted into the oxalate which may be purified by crystallisation from water and then decomposed by ammonia (Doebner, *A* 217, 250)—2 By heating benzoic aldehyde (40 g) with dimethylamine (100 g) and 93 p c alcohol (40 g) over a water bath. POCl_3 (65 g) is then added gradually, and when cool the mass is extracted with warm water and the base ppd with NaOH . The yield is nearly theoretical (Nencki, *M* 9, 1148)—3 By heating benzoic aldehyde with ZnCl_2 and di methyl-aniline, and oxidising the resulting leuco base with PbO (Muhlhauser, *D P J* 263, 249)

Properties—Nearly colourless cubes. Insol water. ν sol alcohol forming a green solution. When freshly ppd it is ν sol ether, but when crystalline it is sl sol ether, m sol CS , acetone, benzene or light petroleum

Reactions—1 HClAq at 250° splits it into dimethylamine and di methyl *p* amido benzo phenone—2 On reduction it yields leuco malachite green (tetra methyl di amido tri phenyl-methane) [101°]—3 Fuming HNO_3 in HOAc forms an amorphous hexa nitro derivative

Salts—The salts of organic acids and

neutral salts of mineral acids are green and dye emerald-green. They are v sol water, the oxalate being m sol water and the picrate sl sol water. Concentrated mineral acids turn the solutions orange, forming acid salts. Diluting with water restores the green colour. In the cold, dilute acids dissolve the base, forming a nearly colourless solution, which turns deep green when heated, a molecule of water being probably split off. Thus the hydrochloride becomes



$(\text{C}_{22}\text{H}_{24}\text{N}_2\text{HCl})_2(\text{ZnCl}_2) \cdot 2\text{aq}$ [c 130°] *Malachite green*. Thick, dark green prisms, sol water (Fischer, B 14, 2520) — $\text{C}_{22}\text{H}_{24}\text{N}_2\text{ZnCl}_2 \cdot \text{aq}$ — $\text{C}_{22}\text{H}_{24}\text{N}_2\text{H}_2\text{SO}_4$ — $\text{C}_{22}\text{H}_{24}\text{N}_2\text{H}_2\text{SO}_4 \cdot \text{aq}$ lustrous green prisms — $(\text{C}_{22}\text{H}_{24}\text{N}_2)_2\text{H}_2\text{C}_2\text{O}_4$ *Malachite green*. Large green tables, sol water and alcohol — Picrates $\text{C}_{22}\text{H}_{24}\text{N}_2\text{C}_6\text{H}_5(\text{NO}_2)_3\text{OH}$ (Golden needles (from benzene), insol water — $\text{C}_{22}\text{H}_{24}\text{N}_2\text{C}_6\text{H}_5(\text{NO}_2)_3\text{OH}$ golden needles

Di-methylo-di-iodide $\text{C}_{22}\text{H}_{24}\text{N}_2\text{Me}_2\text{I}_2 \cdot \text{aq}$ [172°] Formed by heating the base with MeI and MeOH at 100° (Doebner). Green plates, sl sol alcohol, ether, benzene, CS_2 , and cold water, v sol hot water. The same compound is obtained by heating di p amido tri phenyl carbinol with MeI and alcohol at 120° (Doebner, B 15, 236)

Ethyl derivative

$\text{C}_6\text{H}_5\text{C}(\text{OEt})(\text{C}_6\text{H}_5\text{NMe}_2)_2$ [162°] From the base by heating with alcohol at 110° (O Fischer, B 12, 1686)

Sulphonic acid $\text{C}_{22}\text{H}_{24}(\text{SO}_3\text{H})\text{N}_2\text{O}$ Green needles with reddish brown lustre, v e sol hot water, forming a green solution — $\text{NaA}^+ - \text{MgA}^+, 4\text{aq} - \text{CaA}^+, 3\text{aq}$

Derivatives v BROMO, CHLORO, NITRO, and OXY TETRA METHYL-DI AMIDO TRI-PHENYL-CARBINOL

Tetra-methyl-ppo-tri-amido-tri phenyl-carbinol $\text{C}_{22}\text{H}_{24}\text{N}_2\text{O} \cdot \text{e}$

$\text{NH}_2\text{C}_6\text{H}_4\text{C}(\text{OH})(\text{C}_6\text{H}_5\text{NMe}_2)_2$ [191°] Formed by oxidation of the acetyl derivative of the leuco base (tetra methyl di p amido o amido tri phenyl methane) with lead peroxide and dilute H_2SO_4 (Fischer a Schmidt, B 17, 1892). Glistening prisms (from ether). The salts are soluble in water with a bluish-green colour.

Tetra methyl tri p amido-tri phenyl carbinol $\text{NH}_2\text{C}_6\text{H}_4\text{C}(\text{OH})(\text{C}_6\text{H}_5\text{NMe}_2)_2$ *Tetra methyl pararosamine*. Obtained by oxidising the acetyl derivative of tetra methyl tri p amido tri phenyl methane with PbO_2 , and boiling the resulting acetyl derivative with HCl (O Fischer a G Körner, B 16, 2904). Small crystals (from ether).

Penta-methyl tri amido tri phenyl carbinol $\text{C}_{22}\text{H}_{24}\text{N}_3\text{O} \cdot \text{e}$ $(\text{NMe}_2\text{C}_6\text{H}_4)_2\text{C}(\text{OH})\text{C}_6\text{H}_5\text{NHMe}$ *Methyl violet* [180°] Prepared by oxidising dimethylamine with SnCl_4 , with ICl , with HgO_2 and KClO_4 , with KClO_4 , and CuSO_4 , or with NaCl and $\text{Cu}(\text{NO}_3)_2$ (Lauth, *Rep Chim app* 1861, 345, Poirrier a Chappat, *Bt* [2] 6, 502, Hofmann, B 6, 357). Formed also by heating dimethyl aniline with $\text{C}_6\text{H}_5\text{SO}_2\text{Cl}$ at 100° (Hassencamp, B 12, 1275), and, together with formic aldehyde, by shaking hexa methyl tri amido tri phenyl carbinol with MnO_2 , and dilute H_2SO_4 (E a O

Fischer, B 11, 2097). Commercial methyl-violet may be freed from admixed hexa methyl-tri amido tri phenyl carbinol by boiling with ligroin (Wichelhaus, B 16, 2006, 19, 108). Methyl violet occurs in Hofmann's violet.

Properties — Brown powder, melting under water. Insol water, ether, and ligroin. Its alcoholic solution is violet. Its solution in HCl is reddish violet, and is ppd by NaOH , but not by ammonia. Reduced by ammonium sulphide to its leuco base. Tin and HCl reduce it, forming a substance melting at 155°. Boiling HCl splits it up into di methyl aniline and tri-methyl-di amido benzophenone.

Salts — Chloride $\text{C}_{22}\text{H}_{24}\text{N}_3\text{Cl} \cdot \text{e}$ $(\text{NMe}_2\text{C}_6\text{H}_4)_2\text{C}_6\text{H}_5\text{NMeHCl}$ Amorphous mass,

with green metallic lustre. Its aqueous solution is violet, but on adding HCl it becomes first green, then deep yellowish brown. It dissolves in alcohol. Conc H_2SO_4 forms a yellow solution, which on dilution changes through greenish blue to violet. It dyes silk, wool, and mordanted cotton violet — $\text{C}_{22}\text{H}_{24}\text{N}_3\text{I}$ minute needles.

Picrate $\text{C}_{22}\text{H}_{24}\text{N}_3\text{C}_6\text{H}_5(\text{NO}_2)_3(\text{OH})$ Bronzed needles (from alcohol).

Acetyl derivative Acetate $(\text{NMe}_2\text{C}_6\text{H}_4)_2\text{C}_6\text{H}_5\text{NMeAc OAc}$ [225°] From

methyl violet, Ac O , and NaOAc (O Fischer a G Körner, B 16, 2905).

Hexa-methyl tri amido-tri phenyl carbinol $\text{C}_{22}\text{H}_{24}\text{N}_3\text{O} \cdot \text{e}$ $(\text{NMe}_2\text{C}_6\text{H}_4)_3\text{COH}$ *Crystal violet*. *Hexa methyl para rosaniline* [195°]

Formation — 1 Together with its methylo-iodide, by heating methyl violet with MeI and MeOH at 120° (Hofmann, B 6, 363). — 2 By the action of dimethylaniline on tetra methyl-di-amido benzophenone in presence of dehydrating agents (Kern a Caro). In this reaction tetra methyl di amido thio benzophenone may also be used. — 3 By the action of COCl_2 , of CICO_2Et , or of CICO CCl_2 , in presence of ZnCl_2 on di-methylaniline. — 4 By condensation of tetra-methyl di amido-di phenyl carbinol with dimethylaniline and oxidation of the resulting leuco base. — 5 By gradually adding tetra chloroquinone (1 pt) to dimethylaniline (2 pts), and heating the product to 65° (Weister, Lucius, a Brunning, B 13, 212, 2100, Wichelhaus, B 16, 2005). Perhaps the substance formed in this case is wholly or partially the penta methyl compound. — 6 By the action of COCl_2 or of CICO_2Et on dimethylaniline in presence of AlCl_3 (Hofmann, B 18, 767, Wichelhaus, B 19, 109).

Properties — Dark reddish violet monoclinic tables. Needles containing benzene (from benzene). Insol water, sol ether, acetone, and ligroin, sl sol alcohol, v e sol chloroform and benzene. Boiling HCl splits it up into dimethylaniline and tetra methyl di amido benzophenone. Aqueous ammonium sulphide reduces it to hexa-methyl tri amido tri phenyl methane.

Salts — Chloride $\text{C}_{22}\text{H}_{24}\text{N}_3\text{Cl} \cdot \text{e}$ $(\text{NMe}_2\text{C}_6\text{H}_4)_2\text{C}_6\text{H}_5\text{NMeCl}$ Hexagonal crystals,

with greenish brown lustre (Wada, B 18, 768). Sol alcohol. Forms a violet solution in water, which on adding HCl becomes first blue, then green, and finally yellow. NaOH gives a violet pp. Conc H_2SO_4 forms a yellow solution,

changed on dilution through green and blue to violet. Dyes silk, wool, and mordanted cotton bluish-violet— $C_{22}H_{20}N_2Cl$ 8aq crystals, with bronze lustre— $(C_{22}H_{20}N_2Cl)_3PtCl_4$, brick-red crystalline pp, decomposed by water— $C_{22}H_{20}N_2OH_2$, green crystals. At 100° it gives off MeI, becoming the iodide of penta-methyl-tri-amido tri phenyl-carbinol—Picrate $C_{22}H_{20}N_2 \cdot 2C_6H_4(NO_2)_3OH$ yellowish green prisms, with coppery lustre.

Methylo-iodide $C_{22}H_{21}N_2MeI$. From pararosaniline, MeI, and MeOH at 115° (Hofmann, B 6, 365).

TETRA-METHYL-DIAMIDO-DIPHENYL-CUMYL-METHANE $C_{24}H_{26}N_4$, *re* (NMe_2, C_6H_5), $CH_2C_6H_4CH_2$. Prepared by heating cumino aldehyde with dimethylaniline and $ZnCl_2$ to 120° , the yield is about 80 p c. On oxidation it gives a dye stuff closely resembling malachite green.

Salts— $B''H_2Cl$, white crystalline powder— $B''(C_6H_5(NO_2)_2OH)_2$, green crystals, $[156^\circ]$, explodes at 220° — $B''H_2Cl_2PtCl_4$, yellow crystals.

Methylo iodide $B''MeI_2$, white needles $[220^\circ]$. Sl sol cold, v sol hot, water (Zeigler, B 13, 786).

DI-METHYL-AMIDO-PHENYLENE-DI-PHENYL-DIKETONE $NMe_2, C_6H_4(CO C_6H_5)_2$, $[55^\circ]$. From dimethylaniline and $BzCl$ at 180° . Crystals, v sol alcohol and ether.

Hexa-methyl-tri-amido-phenylene-diphenyl-diketone $C_{28}H_{30}N_6O_2$, *re* $NMe_2, C_6H_5(CO C_6H_5, NMe_2)_2$, $[122^\circ]$. From boiling di-methyl aniline and $COCl_2$ (Michler, B 9, 716, Michler a Dupertuis, B 9, 1899). Monoclinic crystals, $a, b, c = 587, 714, \beta = 125^\circ 18'$.

METHYL-AMIDO-PHENYL-ETHANE Nitroso derivative $[41]Et C_6H_5, NMeNO$, $[162^\circ]$. Formed from di-methyl-amido-phenyl-ethane, HCl, and $NaNO_2$ (Heumann a Wiernik, B 20, 2423). Needles, insol water, sl sol ether and cold alcohol. Zinc and HOAc reduce it to ethylphenyl-methyl hydrazine, the acetyl derivative of which melts at 68° .

Di-methyl-amido-phenyl-ethane $[41]Et C_6H_5, NMe_2$, $[89^\circ]$. Formed, together with hexa-methyl-tri-amido-tri-phenyl-methane, by heating glycol (1 mol) with dimethylaniline (2 mols) and $ZnCl_2$ at 100° — 120° (H a W). Needles or prisms (from alcohol), v sol ether, warm alcohol, and benzene, insol water. Its salts are deliquescent, and its platinochloride is unstable. Oxidising agents colour it blue.

Methylo-iodide $Et C_6H_5, NMeI$. From p-amido-phenyl-ethane and MeI (Hofmann, B 7, 527).

Tetra-methyl-di-amido-di-phenyl-ethane $C_{24}H_{28}N_4$, *re* $NMe_2, C_6H_5, CH_2, CH_2, C_6H_5, NMe_2$, $[50^\circ]$ (over 800°). Prepared by heating ethylene bromide with dimethylaniline at 100° (Schoop, B 13, 2196). Slender needles, sol ether, ligroin, hot wood spirit, and alcohol, insol water. With $FeCl_3$ it gives a green colouration, and finally quinone. But it does not yield a dye on oxidation— $B''H_2I_2$. Sol water and alcohol—Oxalate $B''2H_2C_2O_4$ —Picrate $B''C_6H_4(NO_2)_3OH$ yellow pp, sol hot alcohol.

Methylo iodide $C_{24}H_{29}N_4MeI$. From di-amido-di-phenyl-ethane, MeI, and a little KOH at 150° — 180° (Heumann a Wiernik, B 20, 909).

Tetra-methyl-di-amido-tri-phenyl-ethane $C_{26}H_{30}N_4$, $CMc(C_6H_5, NMe_2)_3$. This is the chief product of the action of acetophenone on dimethyl aniline in presence of $ZnCl_2$ (Doebner a Petschoff, A 242, 339). Yellow oil, v sol ether, benzene, petroleum ether, and hot alcohol. It boils above 360° with partial decomposition. It is not volatile with steam.

Hexa-methyl-tri-amido-tri-phenyl-ethane $NMe_2, C_6H_5, CH_2, CH(C_6H_5, NMe_2)_2$, $[125^\circ]$. Formed by heating $CH_3Cl, CHCl_3$ with dimethylaniline and $ZnCl_2$ at 110° — 120° (Heumann a Wiernik, B 20, 2424). White needles, insol water, v sl sol cold, sl sol hot, alcohol, v sol ether. With PbO_2 and HOAc it gives a greenish blue colour.

Octo-methyl-tetra-amido-tetra-phenyl-ethane $C_{28}H_{34}N_8$, *re* (NMe_2, C_6H_5), $CH, CH(C_6H_5, NMe_2)_2$, $[90^\circ]$, (300°) . Formed by heating acetylene tetra-bromide with dimethylaniline at 100° (Schoop, B 13, 2199). Prisms, sol alcohol, ether, and benzene, insol water. With $FeCl_3$ or CrO_3 it produces quinone— $B''2H_2, PtCl_4$, yellow amorphous pp—Picrate $B''C_6H_4(NO_2)_3OH$ yellow plates, sol hot water.

Deca-methyl-penta-amido-penta-phenyl-ethane $C_{30}H_{38}N_{10}$, $[181^\circ]$. Formed by heating a mixture of dimethylaniline (50 g), chloral hydrate (20 g), and $ZnCl_2$ (10 g) at 100° , the yield being 10 g (O Fischer, B 11, 951, A 206, 120, Boessneck, B 18, 1516). Colourless needles (containing aq) (from alcohol), v sol chloroform. Sol benzene, v sl sol alcohol and ether. On oxidation it gives a greenish blue dyestuff.

DI-METHYL- α -AMIDO- ω -DI-PHENYL ETHYL ALCOHOL $NMe, CHPh, CHPh, OH$, $[110^\circ]$. From $NH_2, CHPh, CHPh, OH$, MeI, and EtOH (Goldschmidt a Polonowska, B 20, 494). White needles— $B''_2H_2, PtCl_4, 3aq$.

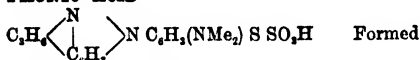
DI-METHYL-AMIDO-PHENYL-ETHYL-DITHIO-CARBONATE EtO, CS, CH_2, NMe_2 , $[51^\circ]$. Formed from di-methyl-p-phenylene diamine by diazotising and heating the product with aqueous potassium xanthate at 70° (Leuckhart, J pr 42, 41, 206). Light yellow crystals, insol water, sol ordinary menstrua. With alcoholic potash it gives $S(C_6H_5, NMe_2)_2$.

TETRA-METHYL-DI-AMIDO-DI-PHENYL-FURFURYL-METHANE $C_{24}H_{28}O, CH(C_6H_5, NMe_2)_2$, $[83^\circ]$. Prepared by the action of furfuraldehyde on dimethylaniline (O Fischer, B 11, 950).

DI-METHYL-AMIDO-PHENYL GLYOXYLIC ACID $C_{21}H_{21}NO_4$, *re* NMe_2, C_6H_5, CO, CO, H , $[187^\circ]$. Obtained by saponifying its ethyl ether which is produced by adding Cl, CO, CO, Et to dimethylaniline at 100° (Michael a Hanhardt, B 10, 2081). Small plates or needles. Sol water and alcohol— NaA' (dried at 150°). Small needles— BaA_2 (dried at 150°). Plates.

Ethyl ether EtA' , $[95^\circ]$. Yellow plates (from alcohol). Cannot be distilled.

TETRA-METHYL-DI-AMIDO-DI-PHENYL-HEPTANE (NMe_2, C_6H_5), CH, C_6H_5 , $[59.5^\circ]$, (275°) at 15 mm. Formed from dimethylaniline, heptonic aldehyde (menthol) and $ZnCl_2$ (Krafft, B 19, 2987). Crystalline solid, not readily oxidised— $B''H_2, PtCl_4$, yellow crystalline pp, v sl sol water and ether alcohol.

DI METHYL - AMIDO - PHENYL - IMIDO - QUINOLINE TETRAHYDRIDE THIOSULPHONIC ACID

by oxidising a mixture of quinoline tetrahydride (1 mol) with di methyl *p* phenylene diamine thio sulphonic acid (1 mol) by $\text{K}_2\text{Cr}_2\text{O}_7$ (Lellmann & Boye, *B* 23, 1374). Small lustrous green needles, sl sol water. Changes after some time to a brown powder. Zinc dust and HCl reduce it, and the product yields a blue dye on oxidation.

DI METHYL AMIDO PHENYL HEXYL KE

TONE $\text{C}_6\text{H}_5\text{NO} \cdot 12 \text{ NMe C}_6\text{H}_5\text{CO C}_6\text{H}_{11}$, [49°] (190° at 20 mm). Formed, together with the leuco base $\text{C}_{13}\text{H}_{15}\text{N}$, by adding heptyl chloride to a mixture of ZnCl_2 and dimethylaniline (Krafft, *B* 19, 2987, Auger, *Bl* [2] 47, 44). Colourless needles. Gives with conc HNO_3 a nitro compound $\text{C}_{13}\text{H}_{15}\text{CO C}_6\text{H}_4(\text{NO}_2)(\text{NMe}_2)$ [65°].

Oxim $\text{NMe C}_6\text{H}_5\text{C}(\text{NOH})\text{C}_6\text{H}_{11}$, [99°]. Colourless leaflets.

DI METHYL DI AMIDO DI PHENYL KE-TONE**DI METHYL AMIDO PHENYL MERCAP**

TAN $\text{NMe C}_6\text{H}_5\text{SH}$ [28°] (260°). Obtained by reducing tetra methyl di amido di phenyl sulphide with tin and HCl (Merz & Weith, *B* 19, 1575, Leuckart, *J pr* [2] 41, 207). Oil. Dissolves in NaOH aq. Re oxidised by air to $(\text{NMe C}_6\text{H}_5)_2\text{S}$ [118°]— $\text{Pb}(\text{SC}_6\text{H}_5)_2$ plates.

Di methyl di amido phenyl mercaptan

$\text{C}_6\text{H}_5\text{N.S} \cdot 12 \text{ C}_6\text{H}_5(\text{NMe})(\text{NH})(\text{SH})$ [112]. Formed by reducing methylene red (Jacobsen, *B* 21, 3104, Bernthsen, *A* 251, 1). Gives a diazo sulphide $\text{C}_6\text{H}_5(\text{NMe})\text{N}=\text{N}$ [78°].

1 ccI, gives a blue colouration. With AcCl in benzene it gives the hydrochloride of ethenyl-amido dimethylamido phenyl mercaptan $\text{NMe C}_6\text{H}_5\text{N}=\text{N}=\text{CMe} - \text{Zn}(\text{C}_6\text{H}_5)_2\text{N.S}$.

DI-METHYL AMIDO - DI PHENYL METHANE $\text{NMe C}_6\text{H}_5\text{CH C}_6\text{H}_5$. This is perhaps the base, boiling at 335°, produced by heating di methyl aniline benzylo chloride in a sealed tube at 230° (Michler, *B* 10, 2079).

Di methyl di amido-di phenyl methane

$\text{NMe C}_6\text{H}_5\text{CH}_2\text{C}_6\text{H}_5\text{NH}_2$ [93°]. Formed by boiling *p* nitro di methyl amido di phenyl carbinol with zinc dust and HCl (Albrecht, *B* 21, 3296). Colourless needles, coloured bluish violet by PbO_2 or by tetra chloro quinone.

Tetra methyl-di *p* amido di phenyl methane

$\text{C}_6\text{H}_5\text{N} \cdot 12 \text{ CH}_2(\text{C}_6\text{H}_5\text{NMe}_2)$, [91°].

Formation—1 By heating methylene iodide, CHCl_3 or CCl_4 with dimethylaniline (Hanhart, *B* 12, 680, Doebner, *B* 12, 810, cf Hannuman, *B* 10, 1235).—2 By heating methylal $\text{CH}_2(\text{OMe})_2$ (1 mol) with dimethylaniline (2 mols) in presence of ZnCl_2 to 120° (O Fischer, *B* 12, 1689, A 206, 117), or by acting on a mixture of methylal and dimethyl aniline with conc HCl aq (Trüger, *J pr* [3] 86, 237).—3 One of the products of the action of acetophenone or of di ethyl ketone on dimethylaniline in presence of ZnCl_2 (Doebner & Petschhoff, *A* 242, 338).—4 By heating dimethyl-

aniline with CCl_4 , SO_2Cl at 100° (Michler & Moro, *B* 12, 1170).—5 One of the products of the action of (a) naphthylamine sulphonic chloride on dimethylaniline (Michler & Salathé, *B* 12, 1789).—6 From C_6Cl_6 , dimethylaniline and ZnCl_2 (Heumann & Wiernik, *B* 20, 2426).—7 One of the products of the action of zinc and HCl on a mixture of dimethylaniline and CS_2 (Wiernik, *B* 21, 3204, cf Trüger, *J pr* [2] 36, 241).—8 By distilling tetra methyl di amido benzophenone (1 pt) with zinc dust (10 pts) (Nathansohn & Muller, *B* 22, 1882).—9 A product of the action of AcO on dimethylaniline (Reverdin & De la Harpe, *B* 22, 1006).—10 The chief product of the action of methyl hexyl ketone on dimethylaniline (Doebner & Petschhoff, *A* 242, 312).—11 By distilling tetra methyl di amido thio benzophenone with zinc dust (Baither, *B* 20, 1737).

Properties—Four sided tables or glistening leaflets (from alcohol). Insol water, sl sol cold, m sol hot, alcohol, v sol benzene, ether, and CS. Not volatile with steam. On oxidation with HOAc and MnO_2 it gives a fugitive blue colour. When boiled with MnO and H_2SO_4 it yields quinone. Its alcoholic solution is turned blue by boiling with tetra chloroquinone. It may be oxidised to tetra methyl di amido benzophenone (N & M). Conc HNO_3 in HOAc forms a tetra-nitro derivative which decomposes at 218°. With tri nitro benzene it forms a compound $\text{C}_6\text{H}_5\text{N}(\text{NO}_2)_3$, [114°], and with *m* di nitro benzene a compound $(\text{C}_6\text{H}_5\text{N})_2\text{C}_6\text{H}_3(\text{NO}_2)_2$, [74°] (Van Romburgh, *R T C* 7, 226).

Salts— $\text{B}^+\text{H}^-\text{I}^-$ sparingly soluble tables. $\text{B}^+\text{H}^-\text{PtCl}_6^-$ yellow precipitate—Picrate $\text{B}^+\text{C}_6\text{H}_5(\text{NO}_2)_3\text{OH}$ [178°].

Methylo iodide B^+MeI^- , [211°]. Yellow plates, v sol hot water and alcohol, insol ether (D & P).

Tetra - methyl - *cro* - *pp* - tri - amido - di-phenyl methane $(\text{NMe C}_6\text{H}_5)_3\text{CHNH}$, [135°]. Obtained by reducing the imide of tetra methyl di amido benzophenone (auramine) with sodium amalgam (Græbe, *B* 20, 3265). Colourless crystals, v sl sol water, m sol alcohol. Gives an intense blue colouration with HOAc.

Methyl *cro* amido tri phenyl methane

$\text{C}_6\text{H}_5\text{N} \cdot 12 \text{ C}_6\text{H}_5\text{CNHMe}$ [73°]. Obtained by passing methylamine into a hot solution of Ph_3CBr in benzene (Hemlin & Silberstein, *B* 17, 745). Prisms (from ligron). Insol water, v sol alcohol— $\text{B}^+\text{H}^-\text{PtCl}_6^-$ 6aq— B^+I^- Ppd as lustrous blue black needles by adding iodine to its solution in CS.

Di methyl-*cro* amido tri phenyl-methane

$\text{C}_6\text{H}_5\text{N} \cdot 12 \text{ C}_6\text{H}_5\text{CNMe}_2$, [97°]. From *cro*-bromo tri phenyl methane and NHMe_2 in benzene (H & S). Insol water, v sol alcohol— $\text{B}^+\text{H}^-\text{PtCl}_6^-$.

Di-methyl amido tri-phenyl-methane $\text{C}_6\text{H}_5\text{N} \cdot 12 \text{ C}_6\text{H}_5\text{CH C}_6\text{H}_5\text{NMe}_2$, [132°]

Formation—1 By heating di phenyl carbinol with di methyl aniline and P_2O_5 at 150° (O Fischer, *B* 11, 951, 12, 1690, A 206, 114).—2 By heating benzophenone chloride Ph_2CCl_2 with dimethylaniline and ZnCl_2 (F., cf Pauly, *A* 187, 209).

Preparation—Benzophenone (10 g), dimethylaniline (13 g), and zinc chloride (10 g)

are heated in sealed tubes for ten hours to 190°. After driving over excess of dimethylaniline and benzophenone the residue is extracted with ether, and after distilling this off the base is recrystallised from alcohol (Doebner & Petschow, *A* 242, 342).

Properties—Colourless needles, sl sol alcohol, v sol ether and benzene. Does not give a colouring matter on oxidation. Is a weak base, and does not combine with HOAc.

Salts— B^+HCl^- , $B^+H_2PtCl_6^-$, needles, sl sol hot water.

Methylo-iodide $C_{21}H_{21}NMeI$ [186°] Large white plates.

Tetra-methyl-di-amido-tri-phenyl methane $C_{25}H_{25}N_2$, *ie* $(NMe_2.C_6H_5)_3CH.C_6H_5$. *Leuco base of malachite green* [102°] and [94°].

Formation—1 By heating a mixture of benzoic aldehyde (10 pts) and dimethylaniline (23 pts) with $ZnCl_2$. The yield is 90 p.c. (O Fischer, *B* 10, 799, 1624, 11, 950, 2274, 12, 1685, *A* 206, 122).—2 From benzylidene chloride, dimethylaniline, and $ZnCl_2$.—3 By heating di-amido tri phenyl methane with MeI and MeOH at 130° (F).—4 By heating its carboxylic acid (di methyl-aniline phthalin) $(NMe_2.C_6H_4)_2CH.C_6H_4.CO_2H$ with barium hydride.—5 From tetra-methyl di amido tri-phenyl-carbinol (malachite green) by reduction with zinc-dust and HCl (Doebner, *B* 11, 1239, *A* 217, 256).—6 A by-product in the action of phthalyl chloride and in that of benzoyl chloride on dimethylaniline.—7 By heating phenyl-glyoxylic acid with dimethylaniline and $ZnCl_2$ (Peter, *B* 18, 539).—8 A by-product in the action of $ZnCl_2$ on a mixture of dimethylaniline and acetophenone (Doebner & Petschow, *A* 242, 333).

Preparation—A mixture of benzoic aldehyde (40 g), dimethylaniline (100 g), and 93 p.c. alcohol (40 g) is heated on a water bath, and $POCl_3$ (56 g) added slowly. The product is extracted with water, the filtrate ppd. by soda and the pp recrystallised from alcohol (Nencki, *M* 9, 1148).

Properties—Crystallises from benzene in [apparently triclinic] needles, melting at 102°, and from alcohol in triclinic leaflets, melting at 94° (E & O Fischer, *B* 12, 796). Insol water, v sol ether and benzene, sl sol petroleum. In small quantities it can be distilled. Its salts are oxidised by MnO_2 or PbO_2 to malachite green. Nitric acid forms a hexa nitro derivative [200°]. When distilled with zinc dust it is reduced to aniline, dimethylaniline, and *p* amido di-phenyl-methane (Manns, *C* 1888, 1363).

Salts— $B^+H_2Cl_2^-$, colourless hygroscopic needles, v sol water. Gives off HCl at 100°, becoming B^+HCl^- . $B^+H_2PtCl_6^-$, white pp, soon becoming yellowish green.—The aurochloride is a golden-yellow flocculent pp.—Picrate $B^+2C_6H_5(NO_2)_3OH^-$ [220°]. Aggregations of needles, m sol water.

Di-methylo-di-iodide $B^+Me_2I_2^-$ [218°–222°] (F), [231°] (Doebner, *B* 13, 2228). Plates, tables, or leaflets, v sol water. Decomposed on fusion into MeI and the base.

Tetra-methyl-ppo tri-amido-tri-phenyl-methane $([4]1)NMe_2.C_6H_5)_3CH.C_6H_5.NH_2$ [12] [185°] *o*-Amido-leuco-malachite green. Formed by reducing, by means of zinc-dust and HCl, the

product of the condensation of *o* nitro benzoic aldehyde with dimethylaniline (Fischer & Schmidt, *B* 15, 683, 17, 1891). Colourless crystals (containing C_6H_5). Gives a reddish brown dye stuff on oxidation.

Acetyl derivative $C_{25}H_{25}(NMe_2)_2(NHAc)$. [186°] Glistening crystals. Gives on oxidation $(NMe_2.C_6H_5)_3C(OH).C_6H_5.NHAc$.

Tetra-methyl-tri-amido-tri-phenyl-methane $C_{25}H_{25}N_3$, *ie* $(NMe_2.C_6H_5)_3CH.C_6H_5.NH$ [65°] Prepared by dissolving tetra methyl di *pp* amido di phenyl carbinol (20 pts) in $HClAq$ (12 pts of $S.G.$ 1.18) and water (100 pts), heating to 100°, and adding aniline hydrochloride (10 pts) (Nathansohn & Muller, *B* 22, 1886). Small needles (from warm alcohol), v sol alcohol, ether, and benzene, insol water. According to analogy it should be identical with the preceding body. With MeI it gives $CH(C_6H_5.NMe_2)_3I$, [172°].

Salts— $B^+H_2Cl_2^-$, yellowish green crystalline pp, got by adding HCl to an ethereal solution of the base. V sol hot alcohol, sl sol ether, insol benzene.— $B^+H_2PtCl_6^-$, light yellow flocculent pp, v sl sol water and alcohol.—Picrate $B^+C_6H_5(NO_2)_3OH^-$, light green flocculent pp, v sol hot alcohol, insol benzene and ether.

Benzoyl derivative $(NMe_2.C_6H_5)_3CH.C_6H_5.NHBz$ [128°] Small, slender blue plates (from warm alcohol), v sol hot alcohol and benzene, sl sol ether, insol water.

Tetra-methyl-ppm-tri-amido-tri-phenyl-methane $([4]1)NMe_2.C_6H_5)_3CH.C_6H_5.NH$ [13] [130°] Prepared by reducing *m* nitro tetra methyl-di amido tri phenyl carbinol (E & O Fischer, *B* 12, 803). Colourless prisms or needles. Gives a green dye on oxidation.

Tetra-methyl tri *p* amido tri phenyl methane $C_{25}H_{25}N_3$, *ie* $([4]1)NMe_2.C_6H_5)_3CH.C_6H_5.NH$ [14] [152°] Prepared by reducing the hydrochloride of *p* nitro tetra methyl di-*p* amido tri phenyl methane with zinc dust (Fischer, *B* 14, 2527). Colourless plates, sl sol alcohol. Gives on oxidation tetra methyl pararosaniline, a reddish violet dye.

Acetyl derivative [108°] Needles. Yields on oxidation the corresponding carbinol, a splendid green dye (Fischer & German, *B* 16, 708).

Penta methyl tri *p* amido tri phenyl methane $C_{25}H_{25}N_4$, *ie* $(NMe_2.C_6H_5)_3CH.C_6H_5.NMeH$ [116°] Obtained by heating tetra methyl pararosaniline with alcoholic ammonium sulphide at 100° (Hofmann, *B* 6, 360, E & O Fischer, *B* 12, 799, Fischer & Körner, *B* 16, 2906). Large colourless needles (from alcohol). Sl sol hot water, v sol alcohol and ether.— $B^+3H_2PtCl_6^-$.

Acetyl derivative [143°] **Hexa-methyl-tri-amido-tri-phenyl methane** $C_{25}H_{25}N_6$, *ie* $HC(C_6H_5.NMe_2)_3$. **Hexa methyl-paraleucaniline** [173°].

Formation—1 By treating chloral with dimethylaniline and $ZnCl_2$ (E & O Fischer, *B* 11, 2097). The base so prepared melted, however, at 250°.—2 From orthoformic ether (1 pt) and di methylaniline ($\frac{3}{2}$ pts) at 100° (Fischer & Knorr, *B* 17, 98).—3 By condensation of *p* di methyl-amido benzaldehyde with di methyl aniline in presence of dry HCl gas or $ZnCl_2$ (Boess-

neck, *B* 19, 566) — 4 From glycol, dimethyl-aniline, and ZnCl_2 at $100^\circ\text{--}120^\circ$ (Heumann a Wiernik, *B* 20, 2421) Needles or prisms When oxidised with MnO_2 and H_2SO_4 it gives formic aldehyde and methyl violet (Fischer)

Tri-methylo-tri-iodide $\text{C}_{12}\text{H}_{12}\text{N}_3\text{I}_3$ s.e. $\text{CH}(\text{C}_6\text{H}_5, \text{NMe}_2)_3$ [185°] From tetra- and hexa methyl-tri-*p*-amido-tri-phenyl methane, MeI, and MeOH at 100° (Hofmann a Girard, *B* 2, 448, Fischer, *B* 12, 2844) Yellow needles (containing aq), decomposes and becomes dark blue on fusion Gives $(\text{C}_{12}\text{H}_{12}\text{N}_3\text{Cl}_3) \cdot 3\text{PtCl}_2 \cdot 2\text{aq}$

Hexa-methyl opp (?) tri-amido-tri-phenyl-methane Tri-methylo-tri-iodide $\text{CH}(\text{C}_6\text{H}_5, \text{NMe}_2)_3$ [172°] Obtained by heating the corresponding tetra methyl tri-amido tri phenyl methane with MeI (Nathansohn a Muller, *B* 22, 1987) Small brown needles (from warm alcohol), v sol alcohol and hot water, almost insol ether and benzene

Hexa-methyl mpp tri-amido-tri-phenyl-methane Tri-methylo-tri-iodide $(3 \frac{1}{2}) \text{NMe}_2 \cdot \text{I} \cdot \text{C}_6\text{H}_5, \text{CH}(\text{C}_6\text{H}_5, \text{NMe}_2)_3$ [141°] Formed by heating the base $\text{C}_{12}\text{H}_{12}\text{N}_3$ or the corresponding $\text{CH}(\text{C}_6\text{H}_5, \text{NH}_2)_3$ with MeI and MeOH at 120° (Fischer, *B* 12, 802, 13, 673) Crystallises with difficulty, and is v sol water Gives rise to $8\text{PtCl}_2 \cdot 2\text{CH}(\text{C}_6\text{H}_5, \text{NMe}_2)_3$

References—CHLORO-, CHLORO-NITRO-, and NITRO-, METHYL-AMIDO-PHENYL-METHANES

TETRA METHYL DI-AMIDO TRI-PHENYL-METHANE CARBOXYLIC ACID $\text{C}_{12}\text{H}_8\text{N}_2\text{O}_4$ s.e. $(\text{NMe}_2, \text{C}_6\text{H}_5)_2\text{CH} \cdot \text{C}_6\text{H}_4 \cdot \text{CO}_2\text{H}$ *Di-methyl aniline phthalin* [200°] Obtained by treating di-methyl-amido phthalide with zinc-dust and HCl (Fischer, *A* 206, 101) Formed also by treating *p*-aldehyde benzoic acid with dimethylaniline and ZnCl_2 (Löw, *A* 231, 367) Plates (from alcohol), v sol ether, sl sol ligroin, m sol alcohol Conc H_2SO_4 forms a bluish violet solution Its zinc salt melts at 147° (L) —Platinochloride $\text{C}_{12}\text{H}_8\text{N}_2\text{O}_4 \cdot \text{H}_2\text{PtCl}_6$ —Picrate $\text{C}_{12}\text{H}_8\text{N}_2\text{O}_4 \cdot \text{C}_6\text{H}_5(\text{NO}_2)_3\text{OH}$

TETRA METHYL DI-AMIDO TRI-PHENYL-METHANE CARBOXYLIC ALDEHYDE $\text{C}_{12}\text{H}_8\text{N}_2\text{O}_3$ s.e. $(\text{NMe}_2, \text{C}_6\text{H}_5)_2\text{CH} \cdot \text{C}_6\text{H}_4 \cdot \text{CHO}$ From terephthalic aldehyde, dimethylaniline, alcohol, and ZnCl_2 (Löw, *A* 231, 381) Needles (from chloroform) Sl sol alcohol, m sol benzene, v sol chloroform Its phenylhydrazide melts at 22° — $\text{B}''\text{H}_2\text{PtCl}_6$

DI-METHYL-AMIDO-PHENYL (a)-NAPHTHYL SULPHONE $\text{C}_{12}\text{H}_8\text{N}_2\text{SO}_2$ s.e. $\text{NMe}_2, \text{C}_6\text{H}_5, \text{SO}_2, \text{C}_{10}\text{H}_7$ Formed, together with tetra methyl di-amido di-phenyl methane, by heating dimethylaniline (2 mols) with the chloride of naphthalene (a) sulphonic acid (Michler a Salathé, *B* 12, 1789) Crystals, v sol alcohol and ether Conc HClO_4 at 180° splits it up into naphthalene, H_2SO_4 , aniline, and MeCl Zinc and H_2SO_4 give dimethylaniline and naphthyl mercaptan Fuming HNO_3 gives tetra nitro methyl aniline and nitro naphthalene sulphonic acid

Di-methyl-amido-phenyl (b)-naphthyl sulphone Resembles the preceding body in its mode of preparation, properties, and decompositions (M a S)

DI-METHYL-*p*-AMIDO-PHENYL-OXAMIC ACID $\text{C}_{12}\text{H}_{12}\text{N}_2\text{O}_4$ s.e. $\text{NMe}_2, \text{C}_6\text{H}_5, \text{NH} \cdot \text{CO} \cdot \text{CO}_2\text{H}$ [192°] The ethyl ether is formed by boiling di-

methyl-*p*-phenylene diamine with oxalic ether, and separated by solution in alcohol from the accompanying tetra methyl di-amido-di-phenyl oxamide (Sendtner, *B* 12, 530) The ether is then saponified by alcoholic KOH. Needles (from water) or plates (from alcohol)

Ethyl ether EtA' [177°] Yellow plates or needles, v sol warm alcohol

DI-METHYL-AMIDO-PHENYL-OXAMIDE $\text{C}_{12}\text{H}_{12}\text{N}_2\text{O}_3$ s.e. $\text{NMe}_2, \text{C}_6\text{H}_5, \text{NH} \cdot \text{CO} \cdot \text{CO} \cdot \text{NH}_2$ [259°] Formed by treating di-methyl-amido-phenyl-oxamic ether with alcoholic NH_3 (Sendtner, *B* 12, 532) Nodules (from alcohol) — $\text{B}''\text{H}_2\text{SO}_4$ crystals

Tetra-methyl-di-amido-di-phenyl oxamide $\text{NMe}_2, \text{C}_6\text{H}_5, \text{NH} \cdot \text{CO} \cdot \text{CO} \cdot \text{NH} \cdot \text{C}_6\text{H}_5, \text{NMe}_2$ Formed as described under di-methyl-amido-phenyl-oxamic acid (Sendtner) Small yellow needles, insol water, sl sol boiling alcohol Does not melt at 270° Diacid base, forming salts soluble in water

TETRA-METHYL-DI-AMIDO-DI-PHENYL OXIDE $\text{C}_{12}\text{H}_8\text{N}_2\text{O}$ s.e. $(\text{NMe}_2, \text{C}_6\text{H}_5)_2\text{O}$ [119°] Formed by boiling the corresponding sulphide ('thiodimethylaniline') with an ammoniacal alcoholic solution of AgNO_3 (Holzmann, *B* 21, 2056) Stellate groups of colourless needles, insol water, sl sol cold alcohol, ether, and benzene Readily soluble in acids Conc HClO_4 at 200° gives MeCl and aniline — $\text{B}''\text{H}_2\text{PtCl}_6$, minute bright yellow plates, sl sol hot alcohol —Picrate $\text{B}''\text{C}_6\text{H}_5(\text{NO}_2)_3\text{OH}$ [150°] Small yellow needles, sl sol cold alcohol and benzene

Tetra methyl di-amido di-phenyl-di-oxide $(\text{NMe}_2, \text{C}_6\text{H}_5)_2\text{O}_2$ *Di-oxo di-methyl aniline* [91°] Obtained by adding (4 mols of) silver nitrate to an alcoholic solution of tetra methyl di-amido di-phenyl-di-sulphide $(\text{NMe}_2, \text{C}_6\text{H}_5)_2\text{S}_2$ (1 mol) treated with conc NH_3 It is also formed by the action of FeCl_3 upon the conc HCl solution of the same base Thin silky needles, or plates V sol alcohol, ether, and benzene, sol hot water It dissolves in acids, but its salts are not crystalline (Merz a Weith, *B* 19, 1573)

TETRA METHYL DI-AMIDO DI-PHENYL-OXINDOLE $\text{C}_{12}\text{H}_8\text{N}_2\text{O}$ s.e. $\text{C}_6\text{H}_5 - \text{C}(\text{O}, \text{H}, \text{NMe}_2)_2$

Di-methyl aniline-NH-CO isatin [234°] Formed by heating isatin with dimethylaniline and ZnCl_2 (Baeyer a Lazarus, *B* 18, 2642) Glistening colourless prisms Sl sol ether, alcohol, and ligroin, insol water Dissolves in acids On oxidation it gives a splendid bluish green dye stuff

DI-METHYL-AMIDO TRI-PHENYL-PHOSPHINE $\text{C}_{12}\text{H}_{12}\text{NP}$ s.e. $\text{NMe}_2, \text{C}_6\text{H}_5, \text{P}(\text{C}_6\text{H}_5)_2$ [152°] Formed by the action of sodium on a mixture of chloro benzene and $\text{NMe}_2, \text{C}_6\text{H}_5, \text{PCl}_2$ (Schenk a Michaelis, *B* 21, 1502) Colourless crystals, v sol benzene, sl sol alcohol and ether Weak base, being almost entirely ppd by water from its solution in HClO_4

Hexa methyl tri-amido tri-phenyl-phosphine $(\text{NMe}_2, \text{C}_6\text{H}_5)_3\text{P}$ [273°] Formed by heating dimethylaniline with PCl_3 in a sealed tube (Hannemann, *B* 9, 845) Formed also as a by product, in the action of PCl_3 on dimethylaniline in presence of AlCl_3 (S a M.) Colourless needles,

which turn blue on exposure to air V e sol chloroform, v sol dilute HCl aq, m sol hot alcohol

DI-METHYL-AMIDO-PHENYL PHOSPHINOUS ACID ν **DI-METHYL AMIDO BENZENE PHOSPHINIC ACID**

TETRA-METHYL-DI-AMIDO-DI-PHENYL-PHTHALIDE $C_{24}H_{24}N_4O_4$, $t.c$

$(NMe_2C_6H_4)_2C \begin{smallmatrix} O \\ \diagup \diagdown \\ O \end{smallmatrix} CO$ *Dimethylaminophthalate* [191⁷] Prepared by heating dimethylaniline with $ZnCl_2$ and phthalyl chloride or phthalic anhydride (O Fischer, *B* 9, 1758, 10, 952, 12, 1691, *A* 206, 92) Colourless pointed crystals, insol water, v sol benzene, v sl sol ligrom Reduced by zinc dust and $HOAc$ to tetra methyl di amido tri phenyl methane carboxylic acid (dimethylaniline phthalin) Potash-fusion gives dimethylaniline, $HOBz$, and phthalic acid HNO_3 gives a hexa-nitro-derivative which decomposes at 230°

Salts— $B'HCl$ small needles, m sol water — $B'H_2Cl_2$ hygroscopic crystalline mass, got by passing HCl into the ethereal solution Gives off HCl (1 mol) at 100° — $B''H_2PtCl_6$ crystal line pp — $B''H_2PtCl_6$ aq prisms — Picrate $B''2C_6H_3(NO_2)_3OH$

Methylo-vodide $B''Me_2I_2$ [c 185°]

Phthal green $C_{24}H_{24}N_4O_4$ This substance, isomeric with the preceding, is also formed in the action of phthalyl chloride on di methylaniline in presence of $ZnCl_2$ (Fischer) Its hydrochloride, $B'HCl$, forms greenish yellow needles, m sol water Its zinc double salt crystallises in brass yellow needles which form a green solution in water Zinc and HCl reduce it to a leuco-base $C_{24}H_{24}N_4O_4$ which crystallises in small prisms [236°], and is easily re-oxidised to phthal green

TETRA METHYL-DI-AMIDO PHENYL-DI-PHENYLENE-OXIDE-CARBINYL CHLORIDE

$C_6H_5C \begin{smallmatrix} C_6H_4NMe_2 \\ \diagup \diagdown \\ C_6H_4NMe_2Cl \end{smallmatrix} O$ *Tetra methyl-rosamine*

me Formed from benzotrichloride and dimethyl-*m* amidophenol at 50° – 60° (Heumann a Rey, *B* 22, 3002) Dark red flocculent pp, strongly fluorescent in acid or neutral solution H_2SO_4 dissolves it giving an orange yellow colour, becoming dark-red on addition of water Silk and wool are dyed in feebly acid bath rose to bluish red

Salts— $B'HCl$ dark red needles with steel-blue reflex, v sol water and $EtOH$ forming solutions with a splendid blue red colour, and yellow fluorescence — $(C_6H_4N_2OCl)_2PtCl_6$ dark-red pp

DI-METHYL-AMIDO-PHENYL-PROPANE $C_{11}H_{17}N$ $t.c$ $NMe_2C_6H_4CH_2CH_2CH_3$ *Di methyl amido propyl benzene* (230°)

From p bromo-di methyl-aniline, propyl bromide, and sodium (Claus a Howitz, *B* 17, 1327) Oil

Methylo-vodide $NMe_2IC_6H_4Pr$ [168°]

Tetra-methyl-di-amido-di-phenyl-propane $(NMe_2C_6H_4)_2CMe_2$ [88°] From acetone (1 mol), dimethylaniline (2 mols), and $ZnCl_2$ (Doebner, *B* 12, 813) Long needles

DI-METHYL-AMIDO-PHENYL-QUINON-

IMIDE $C_{14}H_9N_2O$ $t.c$ $C_6H_4 \begin{smallmatrix} O \\ \diagup \diagdown \\ N C_6H_4NMe_2 \end{smallmatrix}$ *Phenol-blue*. Formed by adding quinone chlor-

imide to a solution of di methyl aniline in concentrated aqueous oxalic acid (Fogh, *B* 21, 889). Formed also by the action of $NaOH$ on di methyl phenylene green (Möhlau, *B* 18, 2914) Black crystals (containing $\frac{1}{2}$ aq) Yields, when treated with hot dilute HCl aq, quinone and dimethyl phenylene p diamine May be reduced to a leuco-base

Sulphonic acid

$SO_3H C_6H_4 \begin{smallmatrix} O \\ \diagup \diagdown \\ M C_6H_4NMe_2 \end{smallmatrix}$ Formed by passing

chlorine into amido phenol sulphonic acid suspended in water, and then adding di methyl-aniline (F) Slender needles, v sl sol hot alcohol, insol cold water and ether Its alkaline solutions are blue Conc H_2SO_4 forms a cherry red solution

TETRA-METHYL DI-AMIDO DI PHENYL SULPHIDE $(NMe_2C_6H_4)_2S$ *Thiodimethylaniline* [126°] Formed by heating dimethyl aniline with persulphocyanic acid or SCl_2 (Tur sin, *B* 17, 584, Holzmann, *B* 20, 1640, 21 2056, Michaelis a Godehaux, *B* 23, 554) Light-yellow needles, sl sol alcohol and benzene

Reactions — 1 Silver nitrate converts it into $(NMe_2C_6H_4)_2O$ — 2 On heating with reduced copper (10 pts) at 300° it yields $NPhMe$

Salts — $B''H_2Cl_2$ [176°] White mass becoming coloured in the air, extremely sol water — $B''H_2PtCl_6$ 2aq — $B''H_2FeCl_6$ 6aq white powder, m sol water — $B''C_6H_3(NO_2)_3OH$ [142°] Yellow needles (from hot alcohol) — $B''2C_6H_3(NO_2)_3OH$ Amorphous [146°] — $B''HNCs$ [168°], pearly plates

Tetra-methyl-di-amido-di-phenyl-di-sulphide $(NMe_2C_6H_4)_2S_2$ *Di sulphido- or di thio di methyl aniline* [118°] Formed by adding

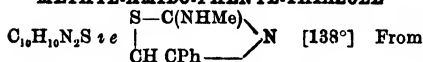
S_2Cl_2 , diluted with petroleum ether, to a dilute solution of dimethylaniline in the same solvent (Merz a Weith, *B* 19, 1570) Formed also by heating di methyl p amido phenyl ethyl di thio carbonate with an alcoholic solution of aniline at 200° (Leuckart, *J pr* [2] 41, 208) Small yellow needles, e sol CS_2 , more sparingly in hot benzene, alcohol, and petroleum ether, nearly insol water Copper powder removes the sulphur at c 230° forming dimethylaniline and other products By the action of Fe_2Cl_6 , or of alcoholic NH_3 and silver nitrate, it is converted into tetra methyl di amido di phenyl di oxide $(NMe_2C_6H_4)_2O_2$ By tin and HCl , or by sodium amalgam, it is reduced to di methyl-amido phenyl mercaptan, which is readily re-oxidised to the di sulphide Its salts are gummy and amorphous, they are decomposed by water

Tetra-methyl-tetra-amido-di phenyl-disulphide $(NMe_2C_6H_4(NH_2))_2S_2$ Formed by atmospheric oxidation from di methyl-di-amido phenyl mercaptan $NMe_2C_6H_4(NH_2)SH$ (Bernthsen, *A* 251, 1) Thick oil, sol ether, alcohol, and benzene Dissolves in acids, but reppd by alkalis In benzene solution it combines with sulphur forming a persulphide [97°], apparently $C_{24}H_{24}N_8S_3$

DI-METHYL AMIDO-DI-PHENYL SULPHONE $C_{14}H_9SO_2C_6H_4NMe_2$ [78°] (R), [82°] (M a M) Formed by heating dimethylaniline with benzene sulphonic chloride (Möhler, *B* 10, 1742, Van Romburgh, *R T C* 2, 805, Michler a. Meyer, *B* 12, 1791, Hassencamp, *B* 12,

1275) Needles (from alcohol), insol water, v e sol alcohol, ether, and benzene HClAq at 180° splits it up into MeCl, aniline, and benzene Zinc and H₂SO₄ reduce it to phenyl mercaptan and dimethylamine HNO₃ gives yellow crystals of tetra nitro-methyl aniline [127°]

METHYL-AMIDO-PHENYL-THIAZOLE



ω bromo acetophenone and methyl thio urea (Traumann, *A* 249, 46) Yellow plates (from ether), insol water, m sol alcohol With HClAq at 220° it yields methylamine

TETRA METHYL DI AMIDO-DI-PHENYL-THIENYL METHANE

$\text{C}_{24}\text{H}_{24}\text{N}_4\text{S}_2$ i.e. (NMe₂, C₆H₄), CH C₆H₅ *Thiophene leuco-malachite green* [93°] Formed by heating thiophene aldehyde with dimethylaniline, ZnCl₂, and a little alcohol (Peter, *B* 18, 538, Levi, *B* 20, 514) Formed also by heating dimethyl aniline with thienyl glyoxylic acid C₆H₅ CO CO₂H and ZnCl₂ (P) Needles, insol water, v sol alcohol and ether The alcoholic solution turns green in the air MnO₂ and dilute H₂SO₄ oxidise it to the carbinol (NMe₂, C₆H₄), C(OH) C₆H₅, which is a dark brown oil, sol alcohol and ether, and forming salts which dye a splendid yellowish green (*Thiophene green*)—Platinichloride C₂₄H₂₄N₄SH₂PtCl₆—Picrate C₂₄H₂₄N₄S₂C₆H₃(NO₂)₃(OH) [c 208°] Yellowish green needles, sl sol cold water, v sol alcohol

Di-methylo di iodide

C₆H₅S CH(C₆H₅, NMe₂)₂ [212°] White plates

TETRA METHYL-DI-AMIDO DI-PHENYL-THIOKETONE v TETRA METHYL DI AMIDO-THIO BENZOPHENONE

DI-METHYL DI AMIDO PHENYL THIO-SULPHURIC ACID NMe₂, C₆H₄(NH₂)₂S SO₂H [193°-204°] Obtained by adding an alkali to a $\frac{1}{10}$ p.c solution of methylene red until the colour is destroyed, then acidulating with HOAc and exposing to the air Formed also by the action of a strong solution of SO₂ on di-methyl di-amido phenyl mercaptan (Bernthsen, *A* 251, 1) Crystals, sl sol water and alcohol Its dilute aqueous solution gives a purple colour with traces of iodine or of FeCl₃ The hydrochloride crystallises in prisms

Reactions—1 A mixture of di-methyl di-amido phenyl thio sulphuric acid with dimethyl aniline when oxidised by K₂Cr₂O₇ and HOAc

forms the indamine $\text{C}_8\text{H}_7\text{N} \begin{array}{c} \text{N} \\ | \\ \text{C}_6\text{H}_4(\text{NMe}_2) \end{array} \text{S}$, an

emerald green powder (containing 3aq), and yields NMe₂, C₆H₄NH C₆H₄(NMe₂) S SO₂H on reduction, a body which is v sol hot alcohol and acids—2 A mixture of the thio-sulphuric acid with aniline hydrochloride gives on oxidation

$\text{C}_{11}\text{H}_{11}\text{N}_2\text{S}_2\text{O}_2$ i.e. $\text{C}_6\text{H}_5 \begin{array}{c} \text{N} \\ | \\ \text{C}_6\text{H}_4(\text{NMe}_2) \end{array} \text{S} (?)$ an

insoluble green compound decomposed by water Prolonged boiling with dilute FeCl₃ gives

$\text{C}_6\text{H}_5 \begin{array}{c} \text{N}(\text{NMe}_2) \\ | \\ \text{NH} \end{array} \text{S}$, a blue powder with bronze

lustre—3 A mixture of di-methyl-di-amido-phenyl thiosulphuric acid with o-toluidine yields

on oxidation the homologous C₁₃H₁₁N₂S₂O₄ as a bluish grey powder

Tetra-methyl-di-amido phenyl thiosulphuric acid C₆H₄(NMe₂)₂S SO₂H [175°-182°] Obtained by dissolving tetra methyl p phenylene diamine (27 g) in HCl (156 c.c of 38 p.c) and adding aluminum sulphate (10 g), and sodium thiosulphate (66 g) dissolved in water (36 c.c) A cold solution of K₂Cr₂O₇ (25 c.c of a 6.67 p.c solution) is then added, and the resulting liquid left to stand The precipitate which then separates is recrystallised from alcohol (Bernthsen, *A* 251, 60) Plates, sol hot water and acids Zinc and HCl reduce it to the mercaptan C₆H₄(NMe₂)₂SH

TETRA-METHYL DI p-AMIDO-DI-PHENYL

THIO UREA C₁₂H₁₂N₂S i.e. (NMe₂, C₆H₄, NH)₂S [186°] Formed by boiling di-methyl-p phenylene diamine with alcoholic CS₂ (Baur, *B* 12, 533) White needles, insol water and cold alcohol—B'H₂Cl₂ crystalline powder

Acetyl derivative [71°] White plates

TETRA-METHYL-DI-AMIDO-DI-ω-PHE-

NYL-TOLUIC ALDEHYDE C₂₄H₂₄N₄O i.e. (NMe₂, C₆H₄), CH C₆H₄CHO *Aldehyde of the leuco base of malachite green* [143°] Obtained by boiling an alcoholic solution of terephthalic aldehyde C₆H₄(CHO)₂ with di-methyl aniline and ZnCl₂ (W Löw, *A* 231, 381) Prismatic needles (from chloroform) Forms a crystalline compound with NaHSO₃ Its phenyl hydrazide melts at 225°—B'H₂PtCl₆

TRI METHYL-TRI AMIDO-DI-PHENYL-

TOLYL CARBINOL C₂₄H₂₄N₃O Formed by heating rosaniline chloride (1 pt) with MeI (2 pts), MeOH (8 pts), and KOH (1 pt) (Hofmann) Formed also by allowing an alcoholic solution containing rosaniline and 'iodine green' to stand in the cold (Girard a Willm, *Bl* [2] 25, 200)

The salt NHMe C₆H₃Me C $\begin{array}{c} \text{C}_6\text{H}_4\text{NHMe} \\ | \\ \text{C}_6\text{H}_4\text{NHMe} \end{array}$ Cl is one

of the substances known as Hofmann's violet (Hofmann, *C R* 54, 428, 56, 945, 1033, 57, 1131) Its absorption spectrum has been studied by Hartley (*C J* 51, 172)

Penta-methyl-tri-amido di-phenyl-tolyl-carbinol

NMe₂, C₆H₄Me C(OH)(C₆H₄NMe₂)(C₆H₄NMeH) The chloride NMe₂, C₆H₄Me C $\begin{array}{c} \text{C}_6\text{H}_4\text{NHMe} \\ | \\ \text{C}_6\text{H}_4\text{NHMe} \end{array}$ Cl

is probably the chief constituent of Hofmann's violet obtained by heating rosaniline with MeCl It has a green lustre and forms a violet solution in water The solution is decolourised by zinc dust HCl turns the solution first green, then yellow NaOH gives a brownish-red pp Conc H₂SO₄ gives a brownish-yellow solution changed, on dilution, through green to blue It dyes wool, silk, and mordanted cotton violet

Hexa-methyl-tri-amido-di-phenyl-tolyl carbinol NMe₂, C₆H₄Me C(OH)(C₆H₄NMe₂)₂ The zinc double salt of the methylo chloride of this body C₂₄H₂₄N₃Cl₂Zn or

NMe₂, C₆H₄Me C $\begin{array}{c} \text{C}_6\text{H}_4\text{NHMe} \\ | \\ \text{C}_6\text{H}_4\text{NHMe} \end{array}$ Cl appears to

constitute the dye known as 'iodine green,' which is obtained by heating rosaniline with MeCl, or by the action of MeCl on Hofmann's violet The

corresponding iodide $C_{12}H_{13}N_3I$, aq is obtained by heating MeI (2 pts) with MeOH (2 pts) and rosaniline (1 pt) at 100° (Hofmann a Girard, *B* 2, 440). The zinc double chloride forms a bluish-green aqueous solution, turned reddish yellow by hydrochloric acid, and rendered colourless by caustic soda. It dyes silk green. When heated strongly in the dry state it becomes violet. Its absorption-spectrum has been studied by Hartley (*C J* 51, 175). Iodine green may be reduced to penta methyl tri amido-di phenyl tolyl methane $NMe, C_6H_4Me, CH(C_6H_5, NMe_2)(C_6H_4, NHMe)$ [173°] (O Fischer a G Körner, *B* 16, 2910). Besides the bodies here described other methylated rosanilines are doubtless formed in the methylation of rosaniline.

DI-METHYL-AMIDO-PHENYL-p-TOLYL SULPHONE $C_{15}H_{17}NSO_2$ [95°] From $4[C_6H_4Me, SO_2, C_6H_4NMe_2]$ [95°] From C_6H_4Me, SO_2Cl and dimethylaniline (Michler a Meyer, *B* 12, 1793). Split up by HCl at 180° into MeCl, aniline, H_2SO_4 , and toluene. Zinc and H_2SO_4 reduce it to *p* tolyl mercaptan and dimethylaniline.

DI-METHYL-p-AMIDO-PHENYL-UREA $C_{12}H_{13}N_3O$ [179°] From potassium cyanate and di methyl-*p*-phenylene-diamine (Binder, *B* 12, 536). Long needles, sol hot water — $B''H_2PtCl_6$, yellow leaflets.

Tetra-methyl-di-*p* amido-di-phenyl-urea $C_{12}H_{13}N_3O$ [262°] (B), [246°] (M a Z). Obtained by heating urea with di-methyl *p* phenylene diamine (Binder, *B* 12, 535). The same body is apparently obtained by the action of COCl₂ on di methyl *p* phenylene diamine (Michler a Zimmermann, *B* 14, 2179) — $B''H_2Cl_2$, soluble crystalline powder — $B''H_2SO_4$, sl sol water — $B''H_2PtCl_6$.

METHYL- α -AMIDO-PROPANE SULPHONIC ACID $NHMe, CH_2, CHMe, SO_3H$ [220°-223°] From methyl propylene ψ -thio-urea and bromine-water (Gabriel, *B* 22, 2989). Colourless columns, v e sol water.

METHYL- α -AMIDO-PROPIONIC ACID $C_6H_5NO_2$ [260°] From α -chloro propionic ether and aqueous methylamine at 130° (Lundenburg, *J pr* [2] 12, 244). Prisms. Decomposed on fusion. Tastes sweet. Its copper salt crystallises in dark blue prisms — $HA'HCl$ [110°]. Deliquescent prisms $H_2A''H_2PtCl_6$, trichinic prisms — $HA'HNO_3$, [126°]. Monoclinic prisms.

Di-methyl α -amido-propionic acid *Methylochloride* $CH_3, CH(NMe_2, Cl), CO_2H$ Formed by treating α -chloro propionic ether with trimethylamine, saponifying the resulting ether with baryta, and adding HCl (Brühl, *B* 9, 34). Very hygroscopic, forming $(C_6H_5, NO_2, Cl)_2PtCl_6$ crystallising in roseate prisms, and $C_6H_5, NO_2, AuCl$, crystallising in golden needles. The corresponding *Anhydride* of the *Methylo-hydroxide* $CH_3, CH \begin{smallmatrix} CO \\ NMe_2 \end{smallmatrix} O$ or 'betaine' is formed on adding baryta to the methylo-chloride and subsequently neutralising with H_2SO_4 . It forms extremely deliquescent crystals, and is neutral in reaction. The iodide $(C_6H_5, NO_2)_2HI$ forms long colourless prisms, v sol alcohol and hot water.

DI-METHYL-AMIDO-PROPYL ALCOHOL v. DI-METHYL OXY-PROPYL AMINE

Tetra-methyl- β -di-amido-isopropyl alcohol C_6H_5, N_3O [170°-185°] Formed from di-chloro-isopropyl alcohol (*s* di-chlorohydrin) and NMe_2H at 60° (Berend, *B* 17, 510). Liquid, v e sol water — $B''H_2PtCl_6$; yellow plates.

Benzoyl derivative $(NMe_2, CH_2)_2CH, OBz$. Forms a platino chloride $B''H_2PtCl_6$, crystallising in tables.

DI-METHYL-AMIDO-PROPYLENE GLYCOL $C_6H_5, NO_2, NMe_2, CH_2, CH(OH), CH_2, OH$ [217°] From dimethylamine and chloro propylene glycol (chlorohydrin) (Roth, *B* 15, 1153). Thick syrup, v e sol water, alcohol, and ether — $B''H_2PtCl_6$.

Methylo-chloride $NMe_2, Cl, CH_2, CH(OH), CH_2, OH$ From the chlorohydrin and NMe_2 at 100° (V Meyer, *B* 2, 186; Hanriot, *A Ch* [5] 17, 99). Needles, v e sol water — $(C_6H_5, NO_2, Cl)_2PtCl_6$, crystalline tables — $C_6H_5, NO_2, AuCl$, [190°] Orange crystals, m sol alcohol.

Di-benzoyl derivative $NMe_2, CH_2, CH(OBz), CH_2, OBz$ oil — Picrate $B''O_2H, (NO_2)_2, OH$ [100°] Laminæ (Roth, *B* 15, 1153).

DI-METHYL-(B 2)-AMIDO-QUINOLINE $NMe_2, C_6H_4, CH, C, CH, CH$ [56°] $C_6H_5, (NMe_2), N$ [56°] CH, CH, C, N, CH

(C), [58°] (O) (e 335°) Prepared by boiling a mixture of *u*-di methyl *p* phenylene diamine (25 pts), glycerine (60 pts), nitro-benzene (15 pts), and H_2SO_4 (50 pts) (La Coste, *B* 16, 672). Colourless crystals. V sol alcohol, ether, and benzene.

Picrate $B''(C_6H_5, NO_2)_2, OH$ very fine orange needles [215°].

Methylo iodide $B'MeI$ long red needles — $(BMeCl)_2PtCl_6$.

Methylo-chloride $B'MeCl$ [244°] Long scarlet hygroscopic needles (Ostermayer, *B* 18, 596).

Methylo chloride of the tetrahydride $NMe, C_6H_5, NMeCl$ [220°], fine colourless needles. With ICl it forms $NMe, C_6H_5, NMeCl, ICl$ [127°], separating in small yellow crystals (Ostermayer, *B* 18, 596).

TETRA-METHYL-DI-AMIDO-QUINONE $C_6H_5, (NMe_2)_2, O_2$ [174°] Red tables. Formed by dissolving ordinary quinone in aqueous dimethylamine (Mylus, *B* 18, 467).

DI-METHYL-AMIDO-SULPHO-BENZOIC ACID $C_6H_5, (NMe_2)_2, (SO_3H), CO_2H$ [4 2 1] Obtained by heating the silver salt of amido sulpho benzoic acid with MeI and MeOH at 100° (Hedrick, *Am* 9, 413). Easily soluble crystals — CaA'' — $BaEt_2A''$.

TETRA-METHYL-DI-AMIDO-THIOBENZO-PHENONE C_6H_5, N_3, S [194°] (B), [202°] (G) S (alcohol) 072 at 18° S (ether) 27 at 18° S (chloroform) 4 53 at 18° . Formed by passing H_2S into an alcoholic solution of the hydrochloride of tetra methyl di-amido benzophenone imide (auramine) at 60° (Fehrmann, *B* 20, 2857; Baither, *B* 20, 1731, 3289). Formed also by the action of CS_2 on auramine (Graebe, *B* 20, 3266). Obtained also from $CSCl_2$ and dimethylaniline. Ruby red crystals, with blue lustre. Insol water and light petroleum, sl sol other solvents. Its

solutions in benzene and CS_2 are dark red with green fluorescence

Reactions — 1 Hot dilute hydrochloric acid gives H_2S and tetra-methyl di amido benzophenone — 2 Water at 120° under pressure also forms $\text{CO}(\text{C}_6\text{H}_4\text{NMe}_2)_2$ — 3 Boiling concentrated nitric acid forms $\text{CO}(\text{C}_6\text{H}_3(\text{NO}_2)_2\text{NMeNO}_2)_2$, whence hydroxylamine hydrochloride forms $\text{CO}(\text{C}_6\text{H}_3(\text{NO}_2)_2\text{NMeH})_2$ [c 196°] — 4 By heating with excess of benzyl chloride it is converted into $\text{CO}_2(\text{C}_6\text{H}_4\text{NMe}_2)_2$, a greyish green powder, decomposed by water with production of $\text{CO}(\text{C}_6\text{H}_4\text{NMe}_2)_2$ — 5 Aniline hydrochloride at 160° forms phenyl auramine $\text{NPh C}(\text{C}_6\text{H}_4\text{NMe}_2)_2$ [171°] — 6 Phenylhydrazine at 100° forms $\text{CO}(\text{C}_6\text{H}_4\text{NMe}_2)_2$ Aniline at 150° also forms this ketone — 7 Hydroxylamine gives the oxim of tetra methyl di amido benzophenone — 8 Distillation over red hot zinc dust yields di methyl aniline and tetra methyl-di amido di phenyl methane — 9 Acetyl chloride in the CS_2 solution forms $\text{C}_{11}\text{H}_{22}\text{N}_4\text{SACl}$, which forms a red alcoholic solution and a greenish yellow solution in benzene It begins to decompose at 160° — 10 Benzoyl chloride in CS_2 forms a similar compound $\text{C}_{11}\text{H}_{22}\text{N}_4\text{SBzCl}$, which is insol water, but decomposed by solution in alcohol — 11 Boiling with Ac_2O and NaOAc forms $\text{C}_{11}\text{H}_{22}\text{N}_4\text{SO}_4$, a dark grey powder — 12 Methyl iodide appears to give $\text{C}_{11}\text{H}_{22}\text{N}_4\text{SMel}$, which forms green plates with coppery lustre, decomposes at 108° , and yields a deep bluish green aqueous solution with red fluorescence It dyes silk green — 13 CSCl_2 acting on its CS_2 solution forms a black powder, probably $\text{C}_{11}\text{H}_{22}\text{N}_4\text{SCSCL}_2$. In chloroform solution, CSCL_2 forms white crusts of $\text{CCl}_2(\text{C}_6\text{H}_4\text{NMe}_2)_2\text{CHCl}$

METHYL-AMIDO-THYMOQUINONE

$\text{C}_{11}\text{H}_{17}\text{NO}_2$, $\pm \text{C}_6\text{H}_5\text{C}_6\text{HMe}(\text{NHMe})\text{O}_2$ [74°] From methylamine and thymoquinone in alcoholic solution (Zincke, B 14, 97) Dark violet laminae (from very dilute alcohol) Volatile with steam M sol water, v e sol alcohol The solutions are deep violet Alcoholic HCl converts it into methylamine and oxythymoquinone

Di methyl amido thymoquinone $\text{C}_{12}\text{H}_{19}\text{NO}_2$, $\pm \text{C}_6\text{H}_5\text{C}_6\text{HMe}(\text{NMe}_2)\text{O}_2$ Prepared by mixing dimethylamine with thymoquinone in alcoholic solution (Schulz, B 16, 899) Dark coloured oil Volatile with steam Split up by heating with dilute HCl into oxythymoquinone and di methylamine

Di methyl di-amido-thymoquinone

$\text{C}_{12}\text{H}_{19}\text{N}_2\text{O}_2$, $\pm \text{C}_6\text{H}_5\text{C}_6\text{HMe}(\text{NHMe})_2\text{O}_2$ [203°] Formed, together with methyl amido thymoquinone, by treating a cold concentrated alcoholic solution of thymoquinone with methylamine (Zincke) Formed also by the action of methylamine on di bromo thymoquinone Long reddish violet needles (from alcohol) Decomposed by KOH or H_2SO_4 in alcoholic solution into methylamine and di oxy thymoquinone

METHYL-AMIDO-TOLUENE v METHYL-TOLUIDINE

Methyl-di-amido-toluene v. METHYL-TOLYL-DIAMINE

Methyl-tri-amido-toluene $\text{C}_{11}\text{H}_{17}\text{N}_3$, $\pm \text{C}_6\text{H}_4\text{Me}(\text{NHMe})_3(\text{NHMe})$ The hydrochloride $\text{B}^+\text{H}_2\text{Cl}_2\text{aq}$ is prepared by reducing the nitramine of di-nitro-methyl-o-toluidine with tin and

HCl (Van Romburgh, R T C 3, 400) It forms small crystals

Di-methyl-amido-toluene sulphonic acid $\text{C}_6\text{H}_4\text{N}_2\text{SO}_3$, $\pm \text{C}_6\text{H}_4\text{MeSO}_3\text{H}$ [214°] Formed by heating di methyl-o toluidine (1 pt) with H_2SO_4 (4 pts) at 180° to 210° (Michler & Sampaio, B 14, 2168) Large glittering prisms (from water), insol alcohol, v sol hot water — CaA'_2 (dried at 130°) Nodules — BaA'_2 (dried at 130°) Laminæ, v sol hot water — ZnA'_2 , needles

TETRA METHYL-DI AMIDO-DITOLYL

$\text{NMe}_2\text{C}_6\text{H}_4\text{MeC}_6\text{H}_4\text{MeNMe}_2$ Tetra - methyl-toluidine [80°] Formed by oxidising di methyl-o toluidine with MnO_2 and dilute H_2SO_4 Formed also by methylation of di amido ditolyl (Michler & Sampaio, B 14, 2170) White plates, sol ether and hot alcohol, insol water — $\text{B}^+\text{H}_2\text{Cl}_2$ white needles — $\text{B}^+\text{H}_2\text{PtCl}_4$ yellow crystalline pp

Tetra methyl di amido-ditolyl $\text{C}_{18}\text{H}_{22}\text{N}_4$, $\pm \text{NMe}_2\text{C}_6\text{H}_4\text{MeC}_6\text{H}_4\text{MeNMe}_2$ [190°] Formed in small quantity by heating dimethyl o-toluidine with H_2SO_4 (M & S) Long needles, sol hot alcohol, ligroin, and ether, insol water Gives a green colouration with Fe_2Cl_6 and yellow colour with CrO_3

Tetra-methyl-di-amido-ditolyl (?) $\text{C}_{18}\text{H}_{22}\text{N}_4$, $\pm \text{NMe}_2\text{C}_6\text{H}_4\text{MeC}_6\text{H}_4\text{MeNMe}_2$ [57°] Prepared by heating dimethyl p toluidine with H_2SO_4 (Michler & Pattinson, B 14, 2167) White needles, sol alcohol and ether, insol water — $\text{B}^+\text{H}_2\text{Cl}_2\text{PtCl}_4$

DI - METHYL - AMIDO - TOLYL - BUTANE

$\text{C}_{11}\text{H}_{17}\text{N}$, $\pm \text{C}_6\text{H}_5\text{C}_6\text{HMeNMe}_2$ [251°] From amido tolyl butane and MeI (Effront, B 17, 2339) — $\text{B}^+\text{H}_2\text{PtCl}_4$

DI - METHYL AMIDO - TOLYL METHYL

KETONE $\text{C}_{11}\text{H}_{17}\text{NO}$, $\pm \text{C}_6\text{H}_5\text{C}_6\text{H}_2(\text{CH}_3)(\text{NHMe})\text{COCH}_3$ [95°] Flat yellowish prisms, e sol alcohol, ether, and hot water, nearly insol petroleum ether Formed by methylation of amido tolyl methyl ketone (Khngel, B 18, 2699)

METHYL α AMIDO α VALERIC ACID

$\text{C}_6\text{H}_9\text{NO}_2$, $\pm \text{CH}_3\text{CH}_2\text{CH}_2\text{CH}(\text{NHMe})\text{COH}$ Obtained by heating butyric aldehyde with conc aqueous HCl, adding methylamine, and heating again The nitrile then separates as an oil, which is hydrolysed by prolonged boiling with dilute HClAq (Menozzi & Belloni, G 17, 116) Long glistening needles (containing aq), decomposed partially at 110° with formation of a sublimate

Salts — $\text{CuA}'_2\text{aq}$ blue prisms — $\text{HA}'\text{HNO}_3$, prisms, v sol water

Methyl α -amido isovaleric acid $\text{C}_6\text{H}_9\text{NO}_2$, $\pm (\text{CH}_3)_2\text{CHCH}(\text{NHMe})\text{CO}_2\text{H}$ Formed by boiling α bromo isovaleric acid with aqueous methylamine (Duvillier, C R 88, 425, A Ch [5] 21, 434) White crystalline powder, v e sol water, m sol hot alcohol, insol ether Neutral in reaction With cyanamide it forms a creatinin $\text{C}_6\text{H}_9\text{N}_3\text{O}$, crystallising in slender needles, v. sol boiling alcohol (Duvillier, C R 95, 456) — $\text{HA}'\text{HCl}$ Crystallises with difficulty — $\text{H}_2\text{A}'\text{H}_2\text{SO}_4$ prisms, v sol water — $\text{HA}'\text{HAuCl}_4\text{aq}$ short trimetric prisms The copper salt forms an intense blue solution

Di - methyl - α - amido - isovaleric acid. Methyl-o-tolide of the methyl ether.

Fr OH(NMe₂I) CO₂Me. Formed by heating MeI (4 pts) with zinc amido-valerate (1 pt) and ZnO (1 pt) at 110° for 16 hrs (Duvillier, *C R* 110, 640). Yields [Fr OH(NMe₂Cl) CO₂H], PtCl₄aq and Fr.OH(NMe₂Cl) CO₂HauCl.

METHYLAMINE ONH, $\frac{1}{2}$ CH₃NH₂. Mol w 81 (−6°) (Hofmann, *B* 22, 701) S G −103.699 H F p 9,540 H F v 8,880 (Thomsen, *Th*) H C 256,900 (Müller, *Bl* [2] 44, 609) Heat of neutralisation by HCl 25,900 (Müller, *A Ch* [6] 15, 531) S (gas) 1150 at 12.5°, 950 at 25°

Occurrence—1 In bone oil (Anderson, *B* 80, 44)—2 In *Mercurialis annua* and *M. perennis* (Schmidt, *B* 10, 2226, *A* 193, 73, cf Reichardt, *Z* 1868, 734)—3 In herring bone (Bookhisch, *B* 18, 1922)—4 In crude wood spirit (G Williams, *Chem Gas* 1853, 409, Commaile, *J* 1873, 686, Vincent, *C R* 77, 898)—5 In the product of the distillation of beet root molasses (Duvillier a Buisne, *A Ch* [5] 23, 317)—6 In the 'yolk' or 'sunt' of sheep When an aqueous extract of this substance is allowed to stand for some time, and is then boiled, it gives off 4 pts of NH₂Me for 95 pts of NH₃, and 1 pt of NMe₂ (Buisne, *C R* 104, 1292)

Formation—1 Discovered by Wurtz, who obtained it by distilling methyl cyanate (or cyanurate) with potash (*C R* 28, 223, 323, *A Ch* [3] 30, 443)—2 By decomposing methylurea with potash (Wurtz)—3 In small quantity, together with NMe₂, and NMe₃, by heating ammonia with MeI (Hofmann, *A* 79, 19) Formed also by heating the methyl ethers of other inorganic acids with ammonia—4 By heating wood spirit with ammonium chloride or iodide at 300° (Berthelot, *J* 1852, 551, Girard, *Bl* [2] 24, 121, cf Weith, *B* 8, 458) By heating methyl alcohol (3 pts) with NH₄Cl (2 pts) and HCl (1 pt) at 207° for 30 hours Dusart a Bardy converted one third of the NH₃ into NH₂Me (*C R* 74, 188).—5 By heating methyl alcohol with ammoniacal ZnCl₂ (Gasiorski, *B* 17, 639)—6 By heating morphine or codeine with KOH (Wertheim, *A* 73, 210, Anderson, *A* 77, 874)—7 By distilling glycochol with BaO (Cahours, *A* 109, 28)—8 By the action of chlorine on theobromine or caffeine (Rochleder a Hlasiwetz, *J* 1850, 434, 437)—9 In small quantity by reducing HCl with zinc and dilute H₂SO₄ (Mendius, *A* 121, 129)—10 By passing a mixture of HCl and hydrogen over platinum-black at 110° (Debus, *A* 128, 200)—11 By reducing nitro methane with iron filings and dilute HOAc (Freibisch, *J pr* [2] 7, 480)—12 By digesting di methyl urea acid with HClAq for several hours at 170° (Hill a Mabery, *Am* 2, 310)

Preparation—1 By distilling methyl cyanurate with aqueous KOH (Wurtz) The distillate is received in dilute HCl, and the methylamine hydrochloride dried and distilled with quicklime 2 A mixture of acetamide (1 mol) and bromine (1 mol) is treated in the cold with a 10 p c solution of KOH till nearly decolourised The solution of methyl-bromo-amide thus obtained is then run in a slow stream into a 30 p c solution of 8 mols of KOH heated to 60°–70° and digested for 10 or 15 mins till decolourised. The solution is then boiled and the methylamine received in HCl, the yield is 87 p c of the theoretical (Hofmann, *B* 15, 765, 18, 2741).—

3 Methyl nitrate (1 mol) is heated in closed vessels at 100° with a solution of ammonia (1 mol) in wood spirit Besides the nitrate of mono methyl-amine there is formed tetra methyl ammonium nitrate together with small quantities of di- and tri methylamine nitrates At the end of the reaction the product is neutralised with sulphuric acid, heated to evaporate off the alcohol, and the residue decomposed by potash, and the bases received in hydrochloric acid The hydrochlorides are treated with absolute alcohol to remove the ammonium chloride, and again decomposed, the bases being received in sulphuric acid The sulphates are then treated with absolute alcohol in which mono methyl amine sulphate is insoluble To remove the last traces of impurities the insoluble sulphate is again decomposed and converted into di methyl oxamide, which yields on decomposition pure mono-methyl amine (Duvillier a Buisne, *A Ch* [5] 23, 322, cf Juncadella, *C R* 48, 342)—4 By the action of tin and HCl on chloropierin (Wallach, *A* 184, 51, cf Geisse, *A* 109, 282)—5 By heating dry ammonium methyl sulphate at 300° and distilling the product with potash (Milner Morrison, *Pr E* 10, 275)

Properties—Colourless gas with strong ammoniacal odour Not solid at −75° Turns red litmus blue Fumes strongly with HCl Rapidly absorbed by water and by charcoal Of all known gases it is the most soluble in water Burns in air with livid yellowish flame (difference from NH₃) The aqueous solution of methylamine is extremely caustic, and gives off the gas when boiled It ppts metallic salts, for the most part in the same manner as ammonia Zinc hydroxide is, however, soluble in a large excess of methylamine With cupric salts it gives a bluish-white pp dissolving in excess and forming a deep blue solution With salts of Cd, Ni, and Co it forms pps insoluble in excess (difference from NH₃) It ppts lead nitrate but not lead acetate With mercurous nitrate it gives a black pp, with HgCl₂ a white pp, with AgNO₃ it gives a pp of Ag₂O soluble in excess It dissolves AgCl With chloride of gold it gives a brownish yellow pp, soluble in excess Platinic chloride gives a yellow crystalline pp

Reactions—1 Passage through a red-hot tube converts it into hydrogen, CH₄, ammonia, and HCl No acetylene, benzene, or C₂H₂ is formed (Müller, *Bl* [2] 45, 438) When an aqueous solution of methylamine is set on fire, HCl is found in the residue (Tollens, *Z* [2] 2, 516)—2 Heated potassium forms hydrogen and KCy—3 Cyanogen chloride forms methyl cyanamide (Cahours a Cloëz, *C R* 38, 354)—4 Iodine forms MeNI₂—5 CO₂ forms methylamine methyl-carbamate—6 According to Berthelot, heating with saturated HIAq yields ammonia and CH₄—7 COCl₂ yields NHMe COCl (90°)—8 Liquid MeCl under pressure, forms NMe₂Cl and NMe₂H₂Cl (Vincent a Chappuis, *C R* 102, 436)—9 MeBr (1 mol) in MeOH at 100° forms, chiefly, NMe₂Br In like manner MeI forms NMe₂I (Duvillier a Buisne, *C R* 90, 1426)—10 Benzene aldehyde added to aqueous NMe₂H, forms PhOH NMe₂, an oil (c 180°) which yields benzyl-methylamine (185°) on reduction (Zaun-schirm, *A* 245, 281).—11 o-Oxy benzoic aldehyde forms C₆H₅NO (229°) (Dennstedt a Zimmer-

mann, *B* 21, 1553) —12 'Dicyanamidobenzoyl' (cf vol 1 p 155) forms small needles of $C_6H_4N_4O$ (Griss, *B* 18, 2420) —13 *Diazo-benzene chloride* added to a cool 33 p.c. solution of methylamine forms $MeN(N_2Ph)_2$ [118°] which crystallises in yellow needles, v sol. ether, m sol. alcohol. It is reduced by Zn and $HOAc$ to methylamine and phenylhydrazine. Boiling dilute H_2SO_4 gives nitrogen, aniline, $MeOH$, phenol, and some $PhN_2C_6H_5NH_2$ (Goldschmidt & Badl, *B* 22, 934) —14 *o* *Diazo-anisole chloride* forms yellow needles of the compound $MeN(N_2C_6H_4OMe)_2$ [141°] (G & B) —15 *p* *Diazo-anisole chloride* forms $MeN(N_2C_6H_4OMe)_2$ [112°] (G & B) —16 *p* *Diazo-toluene chloride* forms $MeN(N_2C_6H_4Me)_2$ [147°]

Salts — $BHCl$. Large deliquescent laminae (from water), may be sublimed. Sol. alcohol, m sol. $CHCl_3$ — $B^+HAuCl_4^-$ aq. Trimetric crystals (Topsoe, *J* 1883, 618) — $B^+HAuCl_4^-$ Monoclinic (T) — $B^+H_2PtCl_6^-$ Golden hexagonal scales (Ludecke, *J* 1890, 511) S 2 at 14° Insol. alcohol, not decomposed by boiling water (De Coninck, *Bl* [2] 45, 131) — $B^+H_2Cl_2PtBr_2^-$ Scarlet crystals (Maly & Hinterberger, *M* 3, 89) — $B^+PtCl_6^-$ Insoluble green powder — $B^+PtCl_6^-$ V e sol. water — $B^+H_2IrCl_6^-$ Small hexagonal brownish black plates (Vincent, *Bl* [2] 43, 154) — $B^+H_2RhCl_6^-$ (Vincent, *C* R 101, 322) — $B^+H_2HgCl_6^-$ Monoclinic crystals, v sol. water — $B^+HHgCl_6^-$ Rhombohedral crystals (T) — $B^+H_2CuCl_6^-$ Trimetric crystals — $B^+H_2PdCl_6^-$ B^+HBr Large deliquescent plates (from alcohol), v sol. water and alcohol — B^+HI — $B^+H_2I_3BiI_4^-$ Scarlet pp (Kraut, *A* 210, 312) — $B^+H_2I_3BiI_4^-$ Crystalline pp (K) — B^+HNO_3 Elongated trimetric prisms. Deliquescent, v sol. water, sl sol. cold alcohol [100°] (Franchimont, *R T C* 2, 338) — $B^+H_2SO_4$ Deliquescent needles, insol. alcohol — B^+HMeSO_4 Crystals, v e sol. water (Classon & Lundvall, *B* 13, 1701) — B^+HVO_4 (Bailey, *C* J 45, 692) — B^+HVO_4 aq. Colourless acicular crystals (Ditte, *C* R 104, 1844) — $B^+(H_2O)(V_2O_5)_4$ aq. Yellow powder (D) — $B^+(H_2O)(V_2O_5)_3$ aq. (B) — $B^+H_2CO_3$ Formed, together with methyl carbamic acid $NMeHCO_2H$, by decomposing $CaCO_3$ with methylamine hydrochloride. Deposited in crystals from the liquid distillate — $B^+H_2C_2O_4$ Prisms, v sol. water, insol. alcohol —Benzene sulphonate [147°] (Norton & Westenhoff, *Am* 10, 129) —Valerate $NH_2Me_2CO_2MeCO_2H$? [81°] (175°) From NH_2Me and tri methyl-acetic acid at 150° (Franchimont & Klobbie, *R T C* 6, 234)

Acetyl derivative C_2H_5NO s. $NMeAcH$ *Methyl acetamide* [28°]. (206°) From $EtOAc$ and aqueous methylamine at 150° (Hofmann, *B* 14, 2725) — $NMeAcHNH_2$ [58°] Large hygroscopic crystals (Franchimont, *R T C* 2, 341)

Di-acetyl derivative $C_4H_8NO_4$ s. $NMeAc_2$ (192°) A product of the action of Ac_2O on methyl-acetyl urea (H.) Liquid, miscible with water. Split up by HCl into methylamine and acetic acid

Tri-chloro-acetyl derivative $CCl_3CO NMeH$ [106°] From CCl_3CO_2Et and aqueous methylamine (Franchimont & Klobbie, *R T C* 6, 234) White crystals, sl sol. water and ether. Slowly attacked by pure HNO_3 , which gives off N_2O

Valeryl derivative $CMe_3CO NMeH$ [91°] (204°) VD 398 From Me_3CCOCl and NH_2Me Methylamine and Me_3CCO_2Me yield only (Me_3CCO_2H), NH_2Me V sol. water and alcohol (F & K.) Pure HNO_3 gives off N_2O

Heptyl derivative $C_7H_{15}CO NMeH$ [9°] (266°) SG 13 895 Thick liquid (F & K)

Benzoyl derivative $C_6H_5CO NMeH$ [78°] Crystallises from alcohol (Romburg, *R T C* 4, 388)

o-Amido benzoyl derivative $C_6H_4(NH_2)CO NMeH$ [80°] From isatoic acid and methylamine solution (Weddige, *J* pr [2] 36, 150) Thick prisms (from benzene), v e sol. alcohol and ether, sol. hot water. Ac_2O gives $C_6H_4(NHAc)CO NMeH$ [172°] $BzCl$ forms $C_6H_4(NHbz)CO NMeH$ [181°] (Körner, *J* pr [2] 36, 159)

Other alkoyl derivatives are described under the acids from which they are derived

Methyl-di-chloro-amine CH_2NCl_2 (60° uncor.) Pungent yellow liquid (Köhler, *B* 12, 770)

Methyl-di-bromo-amine CH_2NBr_2 Formed by the action of bromine and potash on methylamine hydrochloride (Hofmann, *B* 15, 767) Extremely pungent liquid. Slowly converted into methylamine by HCl

Methyl-di-iodo-amine CH_2NI_2 Formed by the action of iodine on aqueous methylamine (Wurtz, *A Ch* [3] 30, 455) Prepared by adding iodine (1g) and water (50g) to methylamine hydrochloride, and then adding aqueous $NaOH$ (Raschig, *A* 230, 221) Brownish red pp., quickly becoming brick red. Completely soluble in aqueous HCl , soon decomposing into methylamine and ICl . Ammonia forms $NMeH$ and NH_3 . Cold potash dissolves it, forming methylamine, KIO_3 , and KI

Methyl nitro-amine CH_2NHNO_2 [38°] Obtained by treating methyl chloroformate with methylamine, nitrating the resulting methyl methyl-carbamate CH_2NHCO_2Me , and boiling the product with ammonia (Franchimont & Klobbie, *R T C* 7, 354, 8, 297) Formed also by decomposing NO_2 , $NMeCOCO_2NMeNO_2$, with aqueous ammonia. Colourless needles, sol. alcohol. Strongly acid in reaction. Converted by KOH and MeI into crystalline Me_2NNO_2 — $KMeNNO_2$. Slender needles, sol. alcohol and water

Dimethylamine C_2H_5N s. $(CH_3)_2NH$ Mol w 45 (72°) (Hofmann, *B* 22, 702) SG -33 687 SV 152 4 (Ramsay) HFp 12,720 (Thomson), 3,500 (Muller, *Bl* [2] 44, 609) HFv 10,980 (Thomson, *Th*) HC 42,600 (M.) *Heat of neutralisation* (by HCl) 23,980 (Müller, *A Ch* [6] 15, 531)

Occurrence —In herring-brine (Bookisch, *B* 18, 1924)

Formation —1 One of the products of the action of MeI on ammonia (Hofmann) —2 A mixture of mono-, di-, and tri methylamine is obtained by heating methyl alcohol with ammoniacal $ZnCl_2$ at 200°–220° (Menz & Gasiorowski, *B* 17, 639) —3 In small quantity by heating the sulphite of aldehyde ammonia in a sealed tube, or by distilling it with lime (Petersen, *A* 102, 317)

Preparation—1 The mixture of iodides obtained by heating ammonia with MeI is distilled with potash, and the evolved bases dried by KOH and condensed by a freezing mixture, the bases NMe_2 , NMe_2H , and NMe_2H_2 being separated by means of oxalic ether (Hofmann, *Pr* 12, 380)—2 The bases obtained from MeBr and NH_3 are treated with nitrous acid, and the dimethylnitrosamine decomposed by HCl and distilled over sodium (Hofmann)—3 Nitroso-dimethylaniline hydrochloride (2 pts) is boiled with water (90 pts) and aqueous NaOH (10 pts of S G 125) (Baeyer & Caro, *B* 7, 964, 8, 616).

Properties—Highly alkaline liquid or gas. Has not been solidified. If to an alcoholic solution of the base picryl chloride $\text{C}_6\text{H}_2(\text{NO}_2)_3\text{Cl}$ be added, and then H_2SO_4 followed by water, there is formed a characteristic yellow pp of tri nitro-dimethyl-aniline (Van Romburgh, *R T C* 2, 106).

Reactions—1 MeCl forms only NMe_2Cl and NMe_2HCl (Vincent & Chappuis, *C R* 102, 436)—2 SO_2Cl_2 forms only $\text{NMe}_2\text{SO}_2\text{Cl}$ (183° — 187°) and $\text{SO}_2(\text{NMe}_2)_2$ (Behrend, *B* 14, 1810)—3 Cyanamide heated with NHMe_2 at 110° forms di-methyl guanidine (Tatarinoff, *C R* 89, 608) 4 *p*-Diazotoluene chloride forms $\text{NMe}_2\text{N}_2\text{O}_2\text{H}$, [46°] (Goldschmidt & Badl, *B* 22, 935).

Salts— B^+HCl V sol chloroform (differ from NH_4Cl and NMe_2HCl (Behrend, *B* 15, 1611, *A* 222, 119)— $\text{B}^+\text{H}_2\text{AuCl}$, Monoclinic needles— $\text{B}^+\text{H}_2\text{PtCl}_2$, Trimeric needles— $\text{B}^+\text{H}_2\text{IrCl}_4$, Trimeric octahedra, *abc* = 1.969 1.1954 (Vincent, *Bl* [2] 43, 154)— $\text{B}^+\text{Rh}_2\text{Cl}_3$, 8aq Large dark garnet red prisms (Vincent, *Bl* [2] 44, 513, *C R* 101, 322)— $\text{B}^+\text{H}_2\text{HgCl}_2$, Monoclinic crystals (Topsoe, *J* 1883, 618)— $\text{B}^+\text{THg}_2\text{Cl}_2$, Monoclinic crystals (*T*)— $\text{B}^+\text{H}_2\text{Hg}_2\text{Cl}_2$, Triclinic crystals— $\text{B}^+\text{H}_2\text{SnCl}_4$, Trimeric tables (Hjortdahl, *J* 1882, 474)— $\text{B}^+\text{H}_2\text{CuCl}_2$ — $\text{B}^+\text{H}_2\text{CuCl}$, Crystals— $\text{B}^+\text{HClCuCl}$, Monoclinic crystals (*T*)— B^+HBr — $\text{B}^+\text{H}_2\text{PbBr}$, Trimeric needles— $\text{B}^+\text{H}_2\text{I}_2\text{BiI}_4$ — $\text{B}^+\text{H}_2\text{I}_2\text{BiI}_4$ (Kraut, *A* 210, 314)— B^+HNO_2 , [74°] Long hygroscopic needles or prisms V sol alcohol (Franchimont, *R T C* 2, 338, 3, 229) Decomposed by heat, yielding nitrogen, CO_2 , and dimethylamine (Romburgh, *R T C* 5, 246)— B^+HVO , (Bailey, *C J* 45, 693)— $\text{B}^+(\text{H}_2\text{O})_2(\text{V}_2\text{O}_5)_2$, 4aq—Benzene sulphionate [110°] (Norton, *Am* 10, 129)—*p*-Toluene sulphionate [78°] (*N*)

Acetyl derivative NMe_2Ac (166°) S G 22 941 From dimethylamine and AcCl dissolved in ether (Franchimont, *R T C* 2, 121, 342) Colourless liquid Fuming HNO_3 forms in the cold NMe_2NO , [57°]

Tri-chloro-acetyl derivative $\text{CCl}_3\text{CO NMe}_2$, [104°] (*C*) S G 12 1441 (*F* & *K*) From $\text{CCl}_3\text{CO CCl}_3$ and NMe_2 (Closz, *A Ch* [6] 9, 145)

Heptyl derivative $\text{C}_7\text{H}_{15}\text{CO NMe}_2$, (248°) S G 12 894 Solidifies below -10° (Franchimont & Klobbie, *R T C* 6, 249) With HNO_3 it gives di-methyl-nitro-amine

Valeryl derivative $\text{CMe}_2\text{CO NMe}_2$, (185°) S G 12 912 Liquid, v sol water Not solidified at -17° HNO_3 (S G 152) forms NMe_2NO .

Benzoyl derivative $\text{C}_6\text{H}_5\text{CO NMe}_2$, [42°] (256° uncor.) Crystals, v sol water Split up by aqueous HCl at 200° into HOBz and

HNMe_2 (Hallmann, *B* 9, 846) Conc HNO_3 gives $\text{C}_6\text{H}_5(\text{NO}_2)\text{CO NMe}_2$ (Romburgh, *R T C* 4, 385) With COCl_2 it forms deliquescent crystals of $\text{C}_6\text{H}_5\text{CCl}_2\text{NMe}_2$, [36°] decomposed by water into HCl and $\text{C}_6\text{H}_5\text{CO NMe}_2$.

Di-methyl-iodo-amine NMe_2I From dimethylamine, iodine, and NaOH Pale yellow pp, which rapidly decomposes (Raschig, *A* 230, 223)

Dimethylnitrosamine NMe_2NO (148° at 725 mm) Formed by treating an aqueous solution of dimethylamine hydrochloride with potassium nitrite (E Fischer, *B* 8, 1587, Renout, *B* 13, 2169) Yellow oil Volatile with steam Reduced by zinc dust and HOAc to di methyl hydrazine Decomposed by boiling HCl into NMe_2H and nitrous acid— B^+HCl white needles, decomposed by water or alcohol

Di-methyl nitro amine NMe_2NO_2 , [57°] From NMe_2Ac and HNO_3 (Franchimont) Large crystals

Di methyl amine tri-bromide Me_2NBr_3 , 2aq or $\text{Me}_2\text{N} \begin{matrix} \text{Br} \\ \text{H} \end{matrix} \begin{matrix} \text{Br} \\ \text{H} \end{matrix} \begin{matrix} \text{Br} \\ \text{H} \end{matrix} \begin{matrix} \text{OH} \end{matrix}$ Formed, as a yellow pp, on adding an excess of bromine water to a cold solution of dimethylamine— $2\text{Me}_2\text{NH} + 4\text{Br} + 2\text{H}_2\text{O} = \text{C}_2\text{H}_5\text{NBr}_3\text{O}_2 + \text{Me}_2\text{NH}_2\text{Br}$ Yellow powder Sol alcohol and ether, v sl sol water It is very unstable on keeping over night in a closed vessel it liquefies to a brown fluid containing free bromine, but under water it can be kept for several days Alkalis and acids decompose it at once, thus HCl gives dimethylamine hydrochloride according to the equation $\text{C}_2\text{H}_5\text{NBr}_3\text{O}_2 + 2\text{HCl} = \text{Me}_2\text{NH}_2\text{HCl} + 3\text{Br} + \text{Cl} + 2\text{H}_2\text{O}$ (Raschig, *B* 18, 2249)

Trimethylamine $\text{C}_3\text{H}_7\text{N}$ *see* NMe_3 , Mol w 59 (35°) S G 12 662 (Hofmann, *B* 22, 703) H F p 15,870 (Thomsen, *Th*), 14900 (Muller, *Bl*, [2] 44, 609) H F v 13550 H C 577,600 (*M*) Heat of neutralisation (by HCl) 17,900 (Muller, *A Ch* [6] 15, 531)

Occurrence—In herring brine (Hofmann, *C J* 5, 288, cf Wertheim, *J* 1851, 480) In the flowers of *Crataegus oxyacantha* (Wicke, *A* 91, 121), *C monogyna*, *Pyrus aucuparia*, and *P communis* (Wittstein, *J* 1854, 479) In the stinking goosefoot (*Chenopodium vulvaria*) (Dessaigues, *J* 1851, 481), in *Arnica montana* (Hesse, *J* 1864, 458), and in the seeds of the beech (Brandl & Rakowicki, *J* 1864, 607) It also occurs in *Mercurialis annua* (E Schmidt, *B* 10, 2226) In human urine (Dessaigues, *A* 100, 218), and in calves' blood which has stood 12 hours (Dessaigues, *J Ph* [3] 32, 43) In ergot of rye (Walz, *J* 1852, 552, Rithausen, *Rph chim pur* 1863, 420, cf Brieger, *H* 11, 184) In small quantity in guano (Hesse, *J* 1857, 402) In bone oil (Anderson, *A* 80, 51) According to Ludwig (*Z* 4, 96) it occurs in small quantity in several Austrian and Hungarian wines In the product of the destructive distillation of putrid brann (Selmi, *G* 6, 468) and of beet-root molasses (Vincent, *C R* 84, 1139, 85, 667, *J Ph* [4] 80, 132, Roscoe, *C N* 39, 107). In the putrefaction of yeast (*A Müller*, *J* 1857, 402) and of wheat dough (Sullivan, *J* 1858, 231). In most of the cases here mentioned the trimethylamine is probably obtained from betaine,

neurine, or lecithin, either by putrefactive decomposition or in the chemical treatment

Formation—1 Together with NMe_2H and NMe_2H by the action of ammonia on MeI , and separated by treatment with oxalic ether, with which it does not react (Hofmann, *C J* 4, 304) 2 By the distillation of NMe_2OH (Hofmann, *A* 93, 325)—3 By heating narcotine with KOH at 280° (Wertheim, *A* 73, 208)—4 By heating extract of calamus root with alkali (Thoms, *B* 21, 1912)—5 By passing coal gas through heated zinc dust (Williams, *C N* 51, 15)

Preparation—1 The residues in the preparation of sugar from beet root are, after fermentation, subjected to dry distillation. The aqueous portion of the distillate is neutralised by H_2SO_4 , ammonium sulphate separated by crystallisation, and, after evaporation, the trimethylamine expelled by an alkali. Commercial trimethylamine still contains methylamine, ethylamine, dimethylamine, propylamine, and isobutylamine. The proportion in which these bases are present varies greatly in different samples. To the aqueous solution of the bases oxalic ether is added, the primary bases being pptd as di alkyl oxamides. The mother liquor is distilled with addition of KOH and the dry bases dissolved in absolute alcohol. Oxalic ether is then added to the alcoholic solution, when the diamines are converted into di alkyl oxamic ethers, and the trimethylamine can be obtained by distillation (Vincent, *C R* 89, 238, 788, Duvillier a Buisine, *C R* 89, 48, 709, 92, 250, *A Ch* [5] 23, 298, cf. Eisenberg, *B* 13, 1669) 2 Perfectly pure trimethylamine is obtained by distilling NMe_2OH and rectifying over sodium (Hofmann, *B* 22, 699)

Properties—Gas with ammoniacal and fishy odour. Remains liquid at -75° . V e sol water. When an aqueous solution is strongly cooled a hydrate $\text{NMe}_2\cdot 7\text{aq}$ [4 3°] separates

Reactions—1 KMnO_4 oxidises it to CO_2 and oxalic acid (Wallach a Claisen, *B* 8, 1237)—2 At a red heat it is converted into HCy and NH_3 (Willm, *Bl* [2] 41, 449). Passed with hydrogen through a red-hot tube it yields NH_3 , cyanogen, hydrocarbons, and $\text{CH}_2(\text{NMe}_2)_2$ (Bo meny, *B* 11, 835)—3 CS_2 forms $\text{NMe}_2\cdot\text{CS}_2$, crystallising in white needles [125°]. This body is m sol chloroform and dilute alcohol, nearly insol absolute alcohol, CS_2 , and benzene. It splits up into its components, even at ordinary temperatures, but with dilute HCl it forms $\text{NMe}_2\cdot\text{CS}_2\cdot\text{HCl}$ and $(\text{NMe}_2\cdot\text{CS}_2)_2\cdot\text{H}_2\text{Cl}$. Conc HCl aq splits it up into its components. Phosphoric acid forms $(\text{NMe}_2\cdot\text{CS}_2)_2\cdot\text{H}_2\text{PO}_4$ (Bleunard, *C R* 87, 1040)—4 Glycol chlorhydrin forms neurine chloride $\text{NMe}_2\cdot\text{ClCH}_2\cdot\text{CH}_2\cdot\text{OH}$ —5 Chloroacetic acid forms betaine $\text{CH}_2\langle\text{NMe}_2\text{O}\rangle\text{CO}$ —

6 Dichlorhydrin of glycerin forms 'sepine' chloride $\text{C}_3\text{H}_7\text{Cl}(\text{OH})\text{NMe}_2\text{Cl}$ and 'aposepine' chloride $\text{C}_3\text{H}_7(\text{OH})(\text{NMe}_2)_2\text{Cl}$ (Niemi kovitoh, *M* 7, 249)—7. Hexa chloroacetone forms $\text{CCl}_3\cdot\text{CO}\cdot\text{NMe}_2$ [104°] (Closz, *A Ch* [6] 9, 145) 8 When heated with aniline hydrochloride it yields a distillate of methyl aniline

Salts— B^+HCl decomposes at 285° , giving off NMe_2 and MeCl , and leaving mono- and dimethylamine hydrochlorides. At 805° ammonia

and methyl chloride are given off from the residue, and at 325° the whole has sublimed, the sublimate consisting of NH_4Cl and methylamine hydrochloride. This decomposition may be utilised for the manufacture of MeCl (Vincent, *C R* 84, 1139, 85, 666)— B^+HBr decomposed between 230° and 300° into NMe_2 , ammonia, and MeBr — B^+HI white scales, decomposed between 210° and 280° into NMe_2 , ammonia, MeI , and a residue of NMe_2I — $\text{B}^+\text{H}_2\text{PtCl}_4$, orange regular crystals (Ludecke, *J* 1880, 512, Topsoe, *J* 1883, 618) S (alcohol) 0362 More soluble than the di- and still more so than the mono-methylamine platinumchloride (Eisenberg, *A* 205, 139)— $\text{B}^+\text{H}_2\text{IrCl}_4$, reddish brown octahedra (Vincent, *Bl* [2] 43, 155)— B^+HAuCl_4 , yellow monoclinic crystals. V al sol water, sol alcohol [220°] (Hesse, *J pr* 71, 480, Zay, *G* 13, 420)— $\text{B}^+\text{H}_2\text{Rh}_2\text{Cl}_4$, 9aq (Vincent, *C R* 101, 322)— B^+HCdCl_4 , trimetric crystals (Hjortdahl, *J* 1882, 475)— $\text{B}^+\text{H}_2\text{HgCl}_4$, monoclinic crystals (T)— B^+HHgCl_4 , monoclinic crystals (T)— $\text{B}^+\text{HHg}_2\text{Cl}_4$, triclinic crystals (T)— $\text{B}^+\text{HHg}_2\text{Cl}_4$, rhombohedra (T)— B^+HCuCl_4 , 2aq monoclinic crystals— B^+HCdBr_4 , six sided hexagonal prisms— $\text{B}^+\text{H PtBr}_4$ — $\text{B}^+\text{H}_2\text{I}_3\text{BiI}_4$ — $\text{B}^+\text{H}_2\text{I}_2\text{BiI}_4$, six-sided crimson plates (Kraut, *A* 210, 316)— B^+HNO_3 [153°] Long needles or prisms, v sol hot alcohol (Franchimont, *R T C* 2, 338)— $\text{B}^+\text{H}_2\text{SO}_4\cdot\text{Al}_2(\text{SO}_4)_3$, 24aq [100°] Large crystals (Reckenschufs, *A* 83, 343)— $\text{B}^+(\text{H O})_2(\text{V}_2\text{O}_5)_3$, 7aq (Bailey, *C J* 45, 692)— $\text{B}^+\text{H}_2\text{C}_2\text{O}_4$, plates (Loschmidt, *J* 1865, 375)—Benzene sulphate [89°] (Westenhoff, *Am* 10, 129)

Tetra methyl ammonium hydroxide NMe_4OH Heat of neutralisation (by HCl) 27,900 (Muller, *A Ch* [6] 15, 531) Obtained by digesting NMe_2I with moist Ag_2O (Hofmann, *Tr* 1850, 93, *C J* 4, 321) White crystalline deliquescent mass, v e sol water. Absorbs CO_2 with avidity. Its solution is strongly alkaline and caustic. Decomposed by heat into NMe_2 and MeOH (Hofmann, *B* 14, 494) Neutralises acids, forming the following salts

Iodoform NMe_4I SG 11 1827, 105 1831 (Clarke, *Am S* [3] 16, 401) The chief product of the action of MeI on ammonia and on mono, di, and tri methylamine. Best obtained by heating a solution of NH_3 in methyl alcohol with MeI at 100° – 120° (Lawson a Collie, *C J* 53, 624) Dimetric prisms (from water) Sl sol water, almost insol alcohol, insol ether. Decomposes at a dull red heat into NMe_2 and MeI . Unites with iodoform forming red crystals of $\text{NMe}_4\cdot 12\text{CHI}_3$. When heated with liquefied NH_3 , in which potassium has been dissolved, the products are KI , NMe_2 , and ethane (Thompson a Cundall, *C J* 53, 761) Unites with halogens, forming NMe_4I , (Welzien, *A* 91, 41, 99, 1), NMe_4I , [130°], NMe_4I , [110°] (Geuther, *A* 240, 68), NMe_4I , NMe_4ICl , $\text{NMe}_4\text{I}_2\text{Cl}$, (NMe_4I) $_2\text{Cl}$, (W), NMe_4IBr , [190°] (Dobbin a Masson, *C J* 49, 851), and $\text{NMe}_4\text{I}_2\text{Cl}$, [216° – 220°] The compound NMe_4I is converted by ammonia into $\text{NMe}_4\text{I}\cdot\text{NH}_3$, a dark-coloured explosive body (Stahlschmidt, *P* 119, 421)— NMe_4HgI_2 , small light-yellow prisms, m sol alcohol— $(\text{NMe}_4)_2\cdot 3\text{HgI}_2$, lemon-yellow scales (Risse, *A* 107, 223)— $(\text{NMe}_4)_2\cdot 2\text{BiI}_3$, amorphous scarlet pp (Kraut, *A* 210, 316)— $\text{NMe}_4\cdot 12\text{HgCy}$, white crystals. On heating for a

long time at 200° it is converted into the isomeric $\text{NMe}_2\text{CyHgICy}$ (Claus & Merck, *B* 16, 2738)

Bromide NMe_2Br *S* 55 26 at 15° Obtained by neutralising the hydroxide by HBr Very deliquescent needles Dissociates at 360° into NMe_2 and MeBr (Lawson & Collier, *C J* 53, 625) Forms crystalline compounds with the halogens (Dobbin & Masson, *C J* 49, 848) — (NMe_2Br) $\cdot\text{PtBr}_2$ regular octahedra (Topsoe)

Chloride NMe_2Cl *H.F.* 27,500 (Müller, *BI* [2] 44, 192) Deliquescent crystals Decomposed above 360° into NMe_2 and, doubtless, MeCl (*L a C*) — (NMe_2Cl) $\cdot\text{HgCl}_2$ trimetric crystals (*T*) — (NMe_2Cl) $\cdot\text{HgCl}_2$ rhombohedra — (NMe_2Cl) $\cdot\text{CuCl}_2$ trimetric crystals — $\text{NMe}_2\text{AuCl}_2$ dimetric crystals

Fluoride NMe_2F From the hydroxide and HF Radiating crystals Decomposes at 180° into NMe_2 and MeF (*L a C*)

Nitrate NMe_2NO_3 *S* (94 p.c. alcohol) 3 at 11° Obtained, together with NMe_2HNO_2 , $\text{NMe}_2\text{H}_2\text{NO}_2$, and NMe_2HNO , by heating methyl nitrate with NH_3 dissolved in MeOH (Duvillier & Buisson, *C R* 90, 872) The di and tri methylamine are formed in very small quantity (Duvillier & Malhot, *C R* 100, 177) Formed also by treating NMe_2I with AgNO_3 (Lawson & Collier, *C J* 53, 628) Needles or plates, v. sol water, sl. sol. cold alcohol Not attacked by boiling KOH Above 300° it yields NMe_2 , formic acid, MeNO_2 , and NO (*L a C*)

Nitrite NMe_2NO_2 From NMe_2I and AgNO_2 Deliquescent crystals Decomposed above 300° into NMe_2 , MeNO_2 , Me_2O , NO , and oxygen (*L a C*)

Sulphate (NMe_2) $\cdot\text{SO}_4$ [280°] Deliquescent crystals Decomposed above 280° into NMe_2 and $\text{NMe}_2\text{SO}_4\text{Me}$, which then undergoes further decomposition (*L a C*)

Chromates (NMe_2) $\cdot\text{CrO}_4$ yellow trimetric crystals, v. sol water (Hjortdahl, *J* 1882, 475) — (NMe_2) $\cdot\text{Cr}_2\text{O}_7$ orange trimetric tables, v. sol water

Sulphite $\text{NMe}_2\text{SO}_3\text{H}$ aq. [180°] Decomposes above 300°, yielding NMe_2 , SO_2 , MeOH , and other products (*L a C*)

Sulphydrate NMe_2SH Very deliquescent Decomposes above 200° into NMe_2 and MeSH

Phosphate From NMe_2I and Ag_3PO_4 Forms a strongly alkaline solution Decomposed at a high temperature into NMe_2 , methyl alcohol, and HPO_4 (*L a C*)

Vanadate NMe_2VO_4 (Bailey, *C J* 45, 693)

Carbonates $\text{NMe}_2\text{CO}_2\text{H}$ *H.F.* 20,570 (Müller) Obtained by saturating a solution of the base with CO_2 (*L a C*) Deliquescent crystals, decomposing above 180° into NMe_2 , methyl alcohol, and CO_2 — (NMe_2) $\cdot\text{CO}_2$ *H.F.* 19,100 Obtained by acting on NMe_2I (2 mols) with Ag_2CO_3 , $\frac{1}{2}\text{Ag}_2\text{O}$ (Müller, *BI* [2] 44, 191)

Oxalate (NMe_2) $\cdot\text{C}_2\text{O}_4$ Formed from NMe_2OH and oxalic acid, or from NMe_2I and silver oxalate Deliquescent crystals, decomposed above 360° into NMe_2 and methyl oxalate, the latter being further resolved into Me_2O , CO , and CO_2

Cyanide NMe_2Cy Prisms Sublimes at 226° V. sol water, m. sol alcohol, insol ether and chloroform (Claus & Merck, *B* 16, 2738) — $\text{NMe}_2\text{CyHgCy}$, [275° uncor.] Yellow crystals — $\text{NMe}_2\text{CyAgCy}$, [212° uncor.] From AgCy

and NMe_2I or NMe_2Cy (Thompson, *B* 16, 2338) Long colourless prisms or slender needles, v. sol water and alcohol, insol ether On dry distillation it yields NMe_2 , acetonitrile, and methyl carbamine

Ferrocyanides (NMe_2) $\cdot\text{FeCy}_2$ 13aq Laminar granular mass (Barth, *B* 8, 1484) — (NMe_2) $\cdot\text{H}_2\text{FeCy}_2$ 2aq (E. Fischer, *A* 190, 184)

Ferricyanide (NMe_2) $\cdot\text{FeCy}_3$ 3aq From NMe_2I and silver ferricyanide (Bernheimer, *B* 12, 408) Unstable hygroscopic prisms, sol water, insol alcohol

Cobaltcyanide (NMe_2) $\cdot\text{CoCy}_2$ 13aq Yellow tables (*C a M*)

Acetate NMe_2OAc [*c* 70°] From NMe_2OH and HOAc Deliquescent needles, decomposed at 200° into NMe_2 and MeOAc (*L a C*)

Benzoate NMe_2OBz [220°–230°] Long deliquescent needles Decomposed above 230° into NMe_2 and MeOBz (*L a C*)

Cyanurate $\text{NMe}_2\text{OC}_3\text{N}_3(\text{OH})_2$ aq (Claus, *J pr* [2] 38, 225)

Picrate [313°] (Lossen, *A* 181, 374)

Trimethylamine iodo-methyle iodide NMe_2CHI_2 From NMe_2 and methyleneiodide (Hofmann) Needles Not attacked by NH_3 Moist silver oxide gives $\text{NMe}_2(\text{OH})\text{CHI}_2$ and $\text{NMe}(\text{OH})\text{CH}_2\text{OH}$ It yields the platinum salt $\text{PtCl}_4(\text{NMe}_2\text{ClCHI}_2)_2$

Trimethylamine ethyle-iodide $\text{C}_2\text{H}_5\text{NI}$ *z* NMe_2EtI From NMe_2 and EtI (Müller, *A* 108, 1) Gives the following derivatives (Topsoe, *J* 1883, 620) — (NMe_2EtCl) $\cdot\text{HgCl}_2$ trimetric crystals — ($\text{NMe}_2\text{EtClHgCl}_2$) monoclinic crystals — ($\text{NMe}_2\text{EtCl}(\text{HgCl}_2)_2$) trimetric crystals — (NMe_2EtCl) $\cdot\text{CuCl}_2$ trimetric crystals — Aurochloride $\text{NMe}_2\text{EtAuCl}_2$ dimetric crystals — (NMe_2EtCl) $\cdot\text{PtCl}_2$ regular crystals — Picrate [300°] (Lossen, *A* 181, 374) The ethyle hydroxide is decomposed on distillation into NMe_2 , ethylene, and water NMe_2EtCl on distillation yields MeCl , NMe_2Et , and NMe_2

Trimethylamine ethyle tri-iodide NMe_2EtI_3 [64°] Regular crystals (Ludecke, *A* 240, 85)

Tri-methylamine ethyle penta iodide NMe_2EtI_5 [26°], dark green tables (Geuther, *A* 240, 66)

Trimethylamine ethyle enne-iodide

NMe_2EtI_9 [38°], black green crystals

Trimethylamine bromo-ethyle bromide

$\text{C}_2\text{H}_5\text{Br NMe}_2\text{Br}$ From NMe_2 and ethylene bromide at 45° (Hofmann, *C R* 47, 558) Needles, v. sol hot alcohol Ammonia as well as moist Ag_2O converts it into $\text{C}_2\text{H}_5\text{NMe}_2\text{OH}$ — ($\text{C}_2\text{H}_5\text{Br NMe}_2\text{Cl}$) $\cdot\text{PtCl}_4$ Octahedral crystals — $\text{C}_2\text{H}_5\text{Br NMe}_2\text{I}$ (Baeyer, *A* 140, 312)

Trimethylamine iodo ethyle iodide

$\text{C}_2\text{H}_5\text{I NMe}_2\text{I}$ From neurine, HI , and phosphorus (Baeyer, *A* 140, 309, 142, 324) Crystalline, sl. sol cold water Moist Ag_2O yields $\text{C}_2\text{H}_5\text{NMe}_2\text{OH}$ — ($\text{C}_2\text{H}_5\text{I NMe}_2\text{Cl}$) $\cdot\text{PtCl}_4$ octahedra

Trimethylamine allylo bromide $\text{C}_3\text{H}_5\text{NBr}$ *z* $\text{NMe}_2\text{C}_3\text{H}_5\text{Br}$ Forms a dibromide $\text{NMe}_2\text{C}_3\text{H}_5\text{Br}_2$ [175°] of which the gold salt melts at 148° (Partheuil, *B* 22, 3317)

Trimethylamine bromo-allylo-bromide

$\text{NMe}_2\text{Br C}_3\text{H}_5\text{CH OCHBr}$ [165°] Formed from $\text{NMe}_2\text{O}, \text{C}_3\text{H}_5\text{Br}$, and alcoholic KOH Colourless prisms v. sol water and alcohol, insol ether,

gives a perbromide $\text{NMe}_2\text{BrCH}_2\text{CHBrCHBr}_2$, crystallising in scales [156°]—Platinochloride ($\text{C}_2\text{H}_5\text{BrNMe}_2\text{Cl}$), PtCl_2 , [220°]—Aurochloride $\text{C}_2\text{H}_5\text{BrNMe}_2\text{AuCl}$, [181°]

Trimethylamine propylo-iodide NMe_2PrI [190°] (Langeli, *G* 16, 385)

Trimethylamine iodo-propylo-iodide $\text{NMe}_2\text{C}_2\text{H}_4\text{I}$ [151°] From the allylo iodide and HI at 100° (Partheil, *B* 22, 3320) Colourless needles, sol water and alcohol, insol ether Alcoholic KOH regenerates the allylo-compound—($\text{C}_2\text{H}_5\text{I NMe}_2\text{Cl}$), PtCl_2 , [237°]—Aurochloride ($\text{C}_2\text{H}_5\text{I NMe}_2\text{Cl}$) AuCl , [135°]

Trimethylamine trimethenyl bromide

$\text{NMe}_2\text{BrCH} \begin{array}{c} \text{CH} \\ \parallel \\ \text{CH} \end{array}$ From the bromo allylo-bromide and alcoholic KOH —Aurochloride ($\text{NMe}_2\text{Cl C}_2\text{H}_5\text{AuCl}$)

Dibromide $\text{NMe}_2\text{BrCH} \begin{array}{c} \text{CHBr} \\ \parallel \\ \text{CHBr} \end{array}$ [187°]

From the preceding and Br Colourless, somewhat hygroscopic crystals Forms a platinochloride [232°] and an aurochloride [193°] both crystallising in tables

Trimethylamine bromo-pentenyl bromide $\text{C}_2\text{H}_5\text{BrNMe}_2\text{Br}$ Formed from trimethylamine and di bromo amylene (valerylene bromide) (Ladenburg, *B* 14, 231, 1342) With HI it forms $\text{C}_2\text{H}_5\text{BrNMe}_2\text{I}$ Silver chloride forms $\text{C}_2\text{H}_5\text{BrNMe}_2\text{Cl}$, whence ($\text{C}_2\text{H}_5\text{BrNMe}_2\text{Cl}$), PtCl_2 , and $\text{C}_2\text{H}_5\text{BrNMe}_2\text{ClAuCl}$, both crystalline

Trimethylamine isoamyl triiodide $\text{NMe}_2\text{C}_2\text{H}_4\text{I}_3$ [80°] From NMe_2 and $\text{C}_2\text{H}_5\text{I}$, the product being treated with iodine Dark-brown prisms, nearly insol water, v sol alcohol

TRIMETHYLAmmELIDE v vol u p 325

DI-METHYL-AMMELINE v vol u p 321

METHYL-AMYL-ACETAL v ALDEHYDE

METHYL-ISOAMYL-ANILINE $\text{C}_2\text{H}_5\text{N}$ v $\text{C}_2\text{H}_5\text{NMe C}_2\text{H}_4\text{N}$ (257°) $\text{SG} \approx 906$ Obtained together with C_2H_5 and water by distilling $\text{C}_2\text{H}_5\text{NMeEtC}_2\text{H}_4\text{OH}$ (Hofmann, *A* 79, 15) Formed also by heating dimethylaniline with isoamyl bromide (Claus & Rautenberg, *B* 14, 622)— $\text{B}^+\text{H}_2\text{PtCl}_2$ crystalline pp—(B^+HI), 3BiI_3

METHYL AMYL ARSINE v Organic compounds of ARSENIC

METHYL AMYL BENZENE v AMYL TOLUENE

Di methyl isoamyl benzene v AMYL XYLENE
METHYL-ISOAMYL-BENZENE SULPHONIC ACID $\text{CH}_2\text{C}_2\text{H}_4(\text{C}_2\text{H}_5)_2\text{SO}_3\text{H}$ From p isoamyl toluene by sulphonation (Fitting & Bigot, *A* 141, 166) — KA^+ The Ba salt is a deliquescent gummy mass

Di methyl isoamyl benzene sulphonic acid $(\text{OH})_2\text{C}_2\text{H}_4(\text{C}_2\text{H}_5)_2\text{SO}_3\text{H}$ Formed by sulphonating isoamyl-xylene (F & B) The K and Ba salts were not obtained in crystals

METHYL AMYL CARBINOL v HEPTYL ALCOHOL

DI METHYL AMYLENE DIKETONE $\text{C}_2\text{H}_5\text{O}_2$ v $\text{CEt}_2(\text{COCH}_2)_2$ Di ethyl-acetyl acetone (c 203°) From $(\text{CH}_3\text{CO})_2\text{CEtNa}$ and EtI at 180° (Combes, *A Ch.* [6] 12, 250) Liquid Split up by KOH into $\text{CH}_3\text{CO}_2\text{K}$ and $\text{CH}_3\text{COCH}_2\text{Et}$

Di methyl amylenone diketone $\text{C}_2\text{H}_5\text{O}_2$ v $(\text{CH}_3\text{COCH}_2\text{CH}_2)_2\text{CH}_2$ Di acetyl-pentane

[49°] (212°–215° at 300 mm) From the carboxylic ether and a dilute solution of KOH in MeOH (Kipping & Perkin, *C J* 55, 337) v sol cold ether, alcohol, acetone, chloroform, and light petroleum Cold conc HNO_3 dissolves it without decomposition Conc H_2SO_4 forms a colourless oil $\text{C}_2\text{H}_5\text{O}$

Di oxim $(\text{CH}_3\text{C}(\text{NOH})\text{CH}_2\text{CH}_2)_2$ [85°] Colourless moss like crystals, v sol cold water

DI METHYL AMYLENE DIKETONE CARBOXYLIC ACID

$\text{CO}_2\text{HCHAcCH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{COCH}_2$ From its ether and KOH dissolved in MeOH Liquid, miscible with water Decomposed by heat into CO_2 and the diketone

Ethyl ether EtA^+ (240° at 200 mm) Prepared from sodium aceto acetic ether and $\text{CH}_3\text{COCH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{Br}$ (Kipping & Perkin, *C J* 55, 333) Thick oil Alcoholic NH_3 converts it into $\text{AcCH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{C} \begin{array}{c} \text{CMe} \\ \parallel \\ \text{CO} \end{array} \text{NH}$

[c 235°], which forms monoclinic crystals, $a b c = 7487.1 \ 3997.7 \ 79^\circ 11'$ This 'dehydro amide' yields an acetyl derivative $\text{C}_{12}\text{H}_{17}\text{AcNO}_2$, an oil, converted by boiling water into the amide $\text{CH}_3\text{COCH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{CONH}_2$, which melts between 200° and 228°

METHYL AMYL ETHER v METHYL AMYL OXIDE

METHYL AMYL KETONE $\text{C}_2\text{H}_5\text{O}$ v $\text{CH}_3\text{COCHCH}_2\text{CH}_2\text{CHCH}_2\text{CH}_2$ (151°) $\text{SG} \approx 837$ Formed by oxidising sec heptyl alcohol obtained from *n* heptane (Schorlemmer, *A* 161, 279, 217, 149) Fragrant liquid Unites with NaHSO_4 Yields on oxidation acetic and *n* valeric acids Formed also by dissolving heptinene C_2H_5 in conc H_2SO_4 , and distilling the product with water (Behal, *A Ch.* [6] 15, 270)

Methyl isoamyl ketone $\text{C}_2\text{H}_5\text{O}$ v $\text{CH}_3\text{COCH}_2\text{CH}_2\text{CH}_2\text{CH}(\text{CH}_3)_2$ (144°) $\text{SG} \approx 828$, μ 818 (Rohn, *A* 190, 308), η 821 (Wagner, *J R* 16, 705)

Formation—1 By heating a mixture of calcium hexoate with calcium acetate (Schmidt, *B* 5, 604)—2 By the action of zinc isoamyl on acetyl chloride (Popoff, *A* 145, 283)—3 By oxidising the corresponding heptyl alcohol (Grimshaw *A* 166, 169)—4 By boiling isobutyl aceto acetic acid with aqueous KOH the yield being 70 p c (Purdie, *C J* 39, 467)

Properties—Oil Reduced by sodium to sec heptyl alcohol and di isopropyl pinacone CrO_2 oxidises it to acetic, isovaleric, and isohexonic acids It combines with NaHSO_4

Methyl amyl ketone Me CO CHMePr (142°–147°) From methyl propyl aceto acetic ether (E J Jones, *A* 226, 293) Oil, smelling of peppermint

Methyl amyl ketone $\text{CH}_3\text{COCHMePr}$ (135°) $\text{SG} \approx 815$ One of the products of the saponification of methyl isopropyl aceto acetic ether (Van Romburgh, *R T C* 5, 235) Does not combine with NaHSO_4 Does not give a solid phenyl hydrazide

Methyl amyl ketone $\text{CH}_3\text{COCH}_2\text{CMe}_2$ (125°–130°) Obtained by oxidising the alcohol $\text{CMe}_2\text{CH}_2\text{CMe}_2\text{OH}$ (Butlerow, *A* 189, 78) Oxidised by CrO_2 to acetic acid and $\text{CMe}_2\text{CO}_2\text{H}$ Does not combine with NaHSO_4

Methyl amyl ketone CH_3COCHEt (138°), $\text{SG} \approx 817$ Obtained by boiling di ethyl aceto-

acetic ether with baryta water (Frankland & Duppe, *A* 138, 212) Forms an oily compound with NaHSO_3 .

Methyl amyl ketone $\text{CH}_3\text{CO CMe}_2\text{Et}$ *Methyl-amyl-pinacolin* (132°) SG 2 842, 21 825 Obtained by the action of ZnMe_2 on $\text{CMe}_2\text{Et COCl}$ (Wyschnegradsky, *A* 178, 108) CrO_3 oxidises it to HOAc and $\text{CMe}_2\text{Et CO}_2\text{H}$

Methyl amyl ketone $\text{CH}_3\text{CO C}_4\text{H}_9$ (142°–146°) From *sec*-heptyl alcohol derived from petroleum heptane (90°) Gives acetic acid on oxidation (Schorlemmer, *A* 166, 172)

Methyl isoamyl diketone $\text{CH}_3\text{CO CO C}_4\text{H}_9$ *Methyl isoamyl-glyoxal* (163°) SG 2 8814 From methyl nitrosohexyl ketone (Otto & Pechmann, *B* 22, 2123) Oil, solidified by cold

Phenyl-hydrazide [100°] White needles (from benzene ligroin)

Di-phenyl-di hydrazide. [114°] Needles (from dilute alcohol)

Oxam [173°] Needles

Oxam-phenyl-hydrazide [132°] White needles (from benzene ligroin)

METHYL AMYL KETONE CARBOXYLIC ACID $\text{CH}_3\text{CO CH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{CO}_2\text{H}$ [30°] A product of the action of a boiling solution of KOH in MeOH upon dimethyl-amylene diketone carboxylic ether (Kipping & Perkin, *jun*, *C J* 55, 338) Also from sodium malonic ether and $\text{CH}_3\text{CO CH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{Br}$, the resulting dicarboxylic acid being distilled Crystalline plates, v sol water— Ag^+ colourless plates

METHYL ISO AMYL OXIDE $\text{C}_7\text{H}_{14}\text{O}$ *ze* $\text{CH}_3\text{O C}_4\text{H}_9$ Mol w 102 (92°) VD 3 74 SV 148 1 (Schiiff) From MeONa and isoamyl iodide (Williamson, *C J* 4, 233)

METHYL-AMYL-PIPERIDINE $\text{C}_{11}\text{H}_{21}\text{N}$ *ze* $\text{C}_4\text{H}_9\text{N}(\text{C}_2\text{H}_5)(\text{CH}_3)$ (190°–193°) Prepared by dry distillation of the alkaline hydrate obtained by the action of moist Ag_2O on amyl piperidine methylo iodide (Schotten, *B* 15, 422) Colourless fluid Sl sol water With MeI it forms a crystalline methylo iodide— B^+HCl hygroscopic salt— $(\text{B}^+\text{HCl})_2\text{PtCl}_6$ sparingly soluble pp, melts at [140°]

METHYL AMYL SULPHIDE $\text{CH}_3\text{S C}_4\text{H}_9$ (137°) VD 58 6 From NaSC_2H_5 and MeI (Obermeyer, *B* 20, 2924)

METHYL AMYL DI-THIO-CARBONATE $\text{*CO}(\text{SMe})(\text{SC}_2\text{H}_5)$ (c 140°) From $\text{Cl CO SC}_2\text{H}_5$ and NaSMe (Schöne, *J pr* [2] 82, 244) Liquid, smelling like CS_2 With ammonia it gives HSC_2H_5 , urea, and HSMe Alcoholic potash forms K_2CO_3 , methyl mercaptan, and HSC_2H_5

METHYL-ANHYDRO-ACETONE-BENZYL *v*. vol 1 p 463

METHYL-ANILINE $\text{C}_6\text{H}_5\text{NHMe}$ Mol w 107 (193° uncor) (Friswell & Green, *B* 19, 2035) SG 15 976 HF –5500 H C v 973,000 (Petit, *C R* 107, 266) *Heat of neutralisation* (by HCl) 6,910 (Vignon, *C R* 106, 1722)

Formation—1 Together with dimethylaniline from aniline and MeI or MeBr (Hofmann, *A* 74, 150, *B* 10, 591, cf Kern, *B* 10, 135)—2 Together with dimethylaniline by heating methyl alcohol with aniline and HCl at 200° under pressure (Girard, *BI* [2] 24, 120, cf Poirner & Chappat, *J* 1866, 903)—3 By heating aniline hydrobromide (or hydriodide) with 10 p.c. more than the calculated quantity of methyl

alcohol to 150° for 8 hours, the yield is 34 p.c. of the theoretical (Reinhardt & Staedel, *B* 16, 29, cf Kramer & Grodzky, *B* 18, 1006)—4 By heating acetanilide with alcoholic sodium ethylate at 170°–200° under pressure (Seifert, *B* 18, 1355)—5 By adding sodium to a solution of acetanilide in xylene, treating the resulting sodium-acetanilide with MeI and boiling the product ($\text{C}_6\text{H}_5\text{NMeAc}$) with alcoholic potash (Hepp, *B* 10, 327)—6 From its formyl derivative (Pictet & Crépieux, *B* 21, 1108)—7 By boiling diazobenzene methyl anilide (100 g) with conc HCl (200 cc), making alkaline with NaOH , and distilling with steam, yield 40 g (Friswell & Green, *B* 19, 2035)

Purification—By the action of MeOH and HCl on aniline a mixture of bases is obtained On adding dilute H_2SO_4 aniline sulphate separates, and the bases liberated from the filtrate may then be heated with AcCl On pouring the product into water the acetyl derivative of methyl aniline separates in long needles, while dimethyl aniline hydrochloride remains in solution The acetyl derivative may be quickly saponified by boiling with conc HClAq (Hofmann, *B* 7, 523) The mixture of bases may also be treated with nitrous acid, whereby a diazo benzene salt, phenyl-methyl-nitrosamine, and nitroso dimethylaniline are formed The nitrosamine, being insoluble in water and acids, separates as a yellow oil, and may be recovered by tin and HCl into methyl aniline (Noelting & Boasson, *BI* [2] 28, 2)

Properties—Oil Its aqueous solution is not coloured by bleaching powder With NaOBr it gives a yellow pp (Deniges, *C R* 107, 662) With CuSO_4 it forms a compound $\text{B}(\text{CuSO}_4)_4\text{CuO}$ (Lachovitch, *M* 9, 514)

Estimation—1 It is treated with a mixture of Ac_2O (1 vol) and di methyl aniline (10 vols), water is added after the reaction, and the solution titrated with phenol phthalein The di methyl aniline does not interfere with the reaction, and the mixture of it with the acetic anhydride keeps well (Girard, *BI* [3] 2, 142)—2 In a mixture of aniline, methylaniline, and dimethyl aniline the aniline is first determined by diazotisation and ppn by (8) naphthol disulphonic acid (*R*) and NaCl Another portion of the mixed bases (2 g) is mixed with Ac_2O (4 g), and, after 30 minutes water (50 cc) is added The liquid is boiled to decompose the excess of Ac_2O , and the acid solution titrated with Na_2CO_3 using phenol phthalein as indicator The amount of Ac_2O found plus that required by the aniline previously determined is then subtracted from the amount taken, and gives a measure of the methyl-aniline present (Reverdin & De la Harpe, *B* 22, 1004)—3 In a mixture of mono- and di methyl aniline the amount of the former present may be known by observing the rise of temperature on mixing with an equal volume of Ac_2O

Reactions—1 HNO_3 gives off red fumes on tanning CO_2 , and forms tetra-nitro-methyl aniline (Van Romburgh, *R T C* 2, 81)—2 NaNO_2 added to a solution of its hydrochloride forms phenyl-methyl-nitrosamine $\text{C}_6\text{H}_5\text{NMeNO}$, a yellow oil solidifying when cooled as needles [12°–15°] It gives no reaction with gallic acid (Reverdin & De la Harpe, *B* 22, 1006)—3 Boiling sulphur forms methenyl amido-phenyl mercap-

tan and crystalline C_6H_5NS , [89°] (360°) (Möhlau a Krohn, *B* 21, 59) —4 A solution of methyl-aniline sulphate saturated with SO_2 forms with an aqueous solution of alloxan small yellowish prisms of $CO \begin{smallmatrix} \text{NH} & \text{CO} \\ \diagdown & / \\ \text{NH} & \text{CO} \end{smallmatrix} C(OH) SO_2 NH_2 PhMe 2aq$ (Pelizzari, *A* 248, 148) —5 An aqueous solution of alloxan forms $(C_6H_5N_2O_2)(NHPhMe)$, a compound crystallising in white scales, with a hydrochloride crystallising in prisms —6 *Diazobenzene chloride* and aqueous $NaOAc$ form $PhNMe N NPh$, a yellowish oil, decomposed by dilute H_2SO_4 into diazobenzene sulphate (or phenol) and methyl-aniline, and by $SnCl_2$ and HCl into phenyl hydrazine and methyl aniline (Nöbling a Binder, *B* 20, 3017) —7 *p-Nitro-diazobenzene chloride* and $NaOAc$ form $C_6H_5(NO_2) N N C_6H_5NHMe$ crystallising in red needles [134°], v sol hot alcohol (N a B) —8 *Bromo-acetophenone* forms $C_6H_5 CO CH_2 NPhMe$ in the cold, and on boiling it gives phenyl indole and phenyl methyl-indole (Cullmann, *B* 21, 2595)

Salts — $B^+H^+PtCl_6^-$ Yellow scales Decomposed by boiling water (De Coninck, *Bl* [2] 45, 131) — $B^+H^+CdBr_4^-$ Trimetric crystals (Hjortdahl, *J* 1882, 522) — $B^+H^+SnBr_6^-$ Monoclinic crystals — $B^+HSO_4^-$ Needles (Classens a Lundvall, *B* 13, 1703)

Formyl derivative $C_6H_5NMeCHO$ [12 5°] (N a L) (250°) (N a L), (256°) (Pictet a Crépieux, *B* 21, 1108) $SG \frac{1}{2}$ 1097 From sodium formamide and MeI (Norton a Livermore, *B* 20, 2273) Formed also from methyl-aniline and the hydrochloride of formimide ether (Pinner, *B* 16, 1652) Oil, could not be solidified by Pictet When boiled with 124 p c nitric acid it yields dimethyl-aniline

Acetyl derivative C_6H_5NMeAc 'Ex-algin' [101 5°] (Kamensky, *A* 214, 236) (256°) (P a C) Formed from sodium acetanilide and MeI (Hepp, *B* 10, 328, Hofmann, *B* 10, 599) Formed also by the action of acetyl bromide on dimethyl-aniline (Staedel, *B* 19, 1947) White needles and tablets, sl sol cold water, v sol dilute alcohol As a medicine it has an energetic action on the cerebro spinal system (Dujardin Beaumetz, *C R* 108, 571, cf Giraud, *C R* 108, 749) It is a good antiseptic Boiling nitric acid forms (4, 2, 1)-di nitro methyl aniline

Thio acetyl derivative $CH_3CS NPhMe$ [59°] (290°) From the acetyl derivative and sulphide of phosphorus (Wallach, *B* 13, 528) Monoclinic plates (from $CHCl_3$) Insol. water and alkalis, sol alcohol and ether

Propionyl derivative $CH_3CH_2CO NPhMe$ [58 5°] From sodium propionamide and MeI (Norton a Allen, *B* 18, 1998) With boiling HNO_3 (100 pts of SG 1029) it yields (4, 2, 1) di-nitro methyl aniline

Oxalyl derivative $(CONPhMe)_2$ (250°) Forms oxalic acid, alcohol, and methyl-aniline (Norton a Livermore, *B* 20, 2273) Boiling nitric acid converts it into di nitro-methyl-aniline [177°]

Benzoyl derivative C_6H_5NMeBz [68°] From methyl-aniline and $BaCl_2$ (Hepp, *B* 10, 329) Formed also by heating di methyl aniline with benzoyl chloride at 180° (Hess, *B* 18, 685) Large monoclinic crystals, insol water, but v

sol other solvents Gives a mono nitro- derivative [136°]

Nitrosamine $PhMeNNO$ [12°-15°] From methyl-aniline hydrochloride and aqueous $NaNO$, (Hepp, *B* 10, 329, Fischer, *A* 190, 151, Reverdin a De la Harpe, *B* 22, 1006) Yellow oil, solidified by cold Reduced by tin and HCl to methyl-aniline Gaseous HCl passed into its solution in alcohol ether forms the isomeric [4 1] $NO C_6H_5NHMe$ [118°] which crystallises from water in prisms, v sol alcohol, and is converted by heating with aqueous $NaOH$ into *p* nitrosophenol and methylamine (Fischer a Hepp, *B* 19, 2991)

Di-methyl-aniline $C_6H_5NMe_2$ Mol w 121 [2°-2 5°] (Friswell a Green, *private communication*) (193°) $SG \frac{2}{3}$ 9575 $\mu_n = 1.559$ (Bruhl, *A* 235, 14) SH (9°-82°) 443 (Schiff, *G* 17, 286) Heat of neutralisation (by HCl) 6,810 (Vignon, *C R* 106, 1722) Formed by heating aniline with MeI or $MeCl$

Preparation —1 Aniline hydrobromide (or hydroiodide) is heated with (2 mols + 10 p c excess of) methyl alcohol to 150° for 8 hours, the yield is 95 p c of the theoretical (Reinhardt a Staedel, *B* 16, 29, cf Lauth, *Bl* 7, 448) —2 Aniline (18 pts) saturated with HCl is mixed with a further quantity (75 pts) of aniline and methyl alcohol (75 pts) The mixture is heated at 230° in closed vessels until the internal pressure falls The use of a comparatively small quantity of HCl avoids the formation of toluidine and allows of the use of iron vessels The HCl is sometimes replaced by H_2SO_4 (Schoop, *Chem Zeit* 11, 253)

Purification —Dimethyl-aniline can be separated from methyl aniline by the methods described under methyl aniline It can also be purified by freezing (Hubner, *A* 224, 347)

Reactions —1 When its vapour is passed through a red hot tube it forms benzonitrile (25 p c), carbazole, NH_3 , benzene, and HCy (Nietzki, *B* 10, 474) —2 When heated in a current of HCl at 180° the products are $MeCl$ and aniline (Lauth, *B* 6, 677) —3 With sodium hypobromite it gives a greenish yellow pp in the cold and a red pp on heating (Denigès, *C R* 107, 662) —4 Bromine (1 mol) at 115° forms methyl violet and naphthalene (Brunner a Brandenburg, *B* 11, 697) —5 By nitration with a mixture of equal volumes of ordinary conc HNO_3 and water at 0° di nitro di-methyl aniline $C_6H_5(NO_2)_2NMe$ [4 2 1] is formed (yield 116 p c) If the mixture is allowed to get warm another di nitro di methyl aniline [probably 5 3 1] is formed (yield 15 p c) By further nitration of the first isomeride by boiling it with fuming nitric acid tri nitro phenyl methyl nitramide $C_6H_5(NO_2)_3NMe(NO_2)$ [6 4 2 1] is produced By the same treatment the second isomeride is converted into di nitro phenyl methyl nitramide $C_6H_5(NO_2)_2NMe(NO_2)$ [5 3 1 7] (Mertens, *B* 19, 2123, cf Bomburgh, *R T O* 2, 31) When nitrated in presence of a large excess of H_2SO_4 (20 pts) it yields as chief product the *m* nitro-derivative whilst the *p* nitro derivative is formed in smaller quantity (Groll, *B* 19, 198, Nöbling, *B* 19, 545) —6 Combines with aluminum chloride with great evolution of heat forming long prisms [88°] (H Giraud, *Bl* [3] 1, 691).

Heated in sealed tube with 5 pts of aluminium chloride for 10 hours at 250° no change is effected. Heated in air with excess of aluminium chloride it yields a tetra methyl benzidine $(\text{CH}_3)_4\text{N}_2\text{C}_6\text{H}_4\text{N}(\text{CH}_3)_2$, small needles [195°]. This base is split at 180° by action of HCl gas into methyl chloride and benzidine [118°]—7 *Nitric oxide* passed for 12 days into a solution of dimethylaniline (500 g) in absolute alcohol (510 g) forms $\text{NMe}_2\text{C}_6\text{H}_4\text{N}(\text{CH}_3)_2$. After 3 or 4 weeks' passage of the gas there are formed a base $\text{C}_6\text{H}_4\text{N}_2$ [173°] and a violet colouring matter $\text{C}_6\text{H}_4\text{N}_2\text{O}$ (?) (Lippmann & Lange, B 13, 2136). 8 *Nitrous acid* forms nitroso di methyl aniline (q v)—9 By heating with sulphur there is formed a compound $\text{C}_6\text{H}_4\text{NS}$ which is converted

by HNO_3 into a base $\text{C}_6\text{H}_4\text{N}(\text{CH}_3)_2\text{S}$ of which

the nitrate $\text{B}'\text{HNO}_3$ crystallises in colourless needles and the platinumchloride $\text{B}'_2\text{H}_2\text{PtCl}_6$ in plates (Mohlau & Krohn, B 21, 65). The compound $\text{C}_6\text{H}_4\text{NS}$ is neither acid nor basic and

appears to be $\text{C}_6\text{H}_4\text{N}(\text{CH}_3)_2\text{S}$. It melts at

89° and forms prisms, insol water, and volatile with steam. On heating with sulphur it is converted into methenyl amido phenyl mercaptan. 10 *Chloride of sulphur* forms tetra methyl di amido di phenyl disulphide $\text{S}_2(\text{C}_6\text{H}_4\text{NMe}_2)_2$ (Hannmann, B 10, 403)—11 Heated with *persulphocyanic acid* it gives $\text{S}(\text{C}_6\text{H}_4\text{NMe}_2)_2$ with simultaneous formation of thiocyanic acid, CS_2 , H_2S , and NH_3 (Tursini, B 17, 586)—12 Mixed with CS_2 and then treated with zinc dust and HCl it gives tetra-methyl di amido di phenyl methane [90°] and thioformic paraldehyde [212°] (Wiernik, B 21, 3204)—13 *Oxidising agents* give rise to penta methyl tri amido tri-phenyl carbinol (methyl violet) (O & E Fischer, B 11, 2099)—14 COCl_2 forms $\text{CO}(\text{C}_6\text{H}_4\text{NMe}_2)_2$ and, at 120°, $\text{NMe}_2\text{C}_6\text{H}_4(\text{CO C}_6\text{H}_4\text{NMe}_2)_2$ —15 CCl_4 gives at 180°C $(\text{C}_6\text{H}_4\text{NMe}_2)_4$ (Hannmann, B 10, 1689). 16 *Chloroform* at 230° yields $\text{CH}(\text{C}_6\text{H}_4\text{NMe}_2)_3$ (Hannmann, B 10, 1235)—17 *Chloral hydrate* in presence of ZnCl_2 forms $\text{CCl}_3\text{CH}(\text{OH})\text{C}_6\text{H}_4\text{NMe}_2$ (Knöfler & Boessneck, B 20, 3195)—18 *Furfuraldehyde* in presence of zinc chloride yields $\text{C}_6\text{H}_4\text{OCH}(\text{C}_6\text{H}_4\text{NMe}_2)_2$ which crystallises in pale yellow needles [83°] and is a strong base (O Fischer, A 206, 141)—19 With *heptol aldehyde* and ZnCl_2 it also forms a condensation-product (Auger, Bl [2] 47, 42)—20 *Benzoin aldehyde* and ZnCl_2 forms $\text{C}_6\text{H}_4\text{CH}(\text{C}_6\text{H}_4\text{NMe}_2)_2$ (O Fischer, B 10, 1624)—21 When treated in the cold with *acetyl bromide* it becomes hot and yields acetyl mono methyl aniline (Staedel, B 19, 1947)—22 *Heptyl chloride* in presence of ZnCl_2 forms a base [72.5°] (278° at 15 mm) (Krafft, B 19, 2987)—23 By heating with *benzoyl chloride* at 190°, benzoyl methyl aniline is formed with splitting off of MeCl (Hess, B 18, 685, cf Michler, B 9, 1899)—24 *Benzoin acid* (1 mol) and P_2O_5 at 200° forms di-methyl amido-benzophenone [83°] (O Fischer, A 206, 88)—25 *Phthalic anhydride* in presence of ZnCl_2 forms di-methyl-aniline phthalein—26 *Benzotrichloride* forms 'malachite green'

$(\text{NMe}_2\text{C}_6\text{H}_4)_2\text{CCl}_3\text{C}_6\text{H}_5$. The same body is formed by the action of Bz_2O or BzCl in presence of P_2O_5 —27 *Orthoformic ether* and ZnCl_2 form $\text{CH}(\text{O}_2\text{H}_2\text{NMe}_2)_2$ —28 *Vanillin* and ZnCl_2 form $\text{C}_6\text{H}_4\text{N}_2\text{O}$ [136°] (Fischer & Schmidt, B 17, 1895)—29 A blue compound is obtained by the oxidation of dimethylaniline with *chloranil* or other chlorinated quinone. It appears to have the same constitution whichever chloro quinone is used, as on reduction it always gives a leuco base of the formula $\text{C}_{12}\text{H}_{10}\text{N}_2$, and melting point [173°]. The latter forms glistening plates, sl sol cold alcohol, v sol hot alcohol, benzene, and ether. It forms the salts $\text{B}'\text{H}_2\text{Cl}_2$, colourless crystals— $\text{B}'\text{H}_2\text{Cl}_2\text{PtCl}_6$, fine needles— $\text{B}'\text{Me}_2\text{Cl}_2$, crystals, sol water, sl sol alcohol, with wet Ag_2O it gives an alkaline fluid (Meister, Lucius, & Bruning, B 13, 212, 2100, Wöhlerhaus, B 14, 1952)—30 *Acetone* saturated with SO_2 forms $(\text{C}_6\text{H}_5\text{O})(\text{SO}_2)(\text{NMe}_2\text{Ph})$ crystals in lustrous plates, v sol water and alcohol, insol acetone (Boessneck, B 21, 1906). Acetone and ZnCl_2 at 150° forms $\text{CMe}_2(\text{C}_6\text{H}_4\text{NMe}_2)_2$ [83°] (Doebner, B 12, 810)—31 Di-methylaniline dissolved in a saturated solution of SO_2 and mixed with an aqueous solution of *alloxan* forms $(\text{C}_6\text{H}_4\text{N}_2\text{O})(\text{NMe}_2\text{Ph})\text{H}_2\text{SO}_4$ 4aq crystallising in monoclinic plates, $a, b, c = 1.3399, 1.3394, \beta = 93^\circ 23'$ (Pellizzari, A 218, 148, G 18, 329)—32 A saturated aqueous solution of *alloxan* forms a compound $\text{C}_{12}\text{H}_{10}\text{N}_2\text{O}_4$ aq crystallising in colourless needles, sl sol water, decomposing at 250° (Pellizzari, G 17, 409). The compound forms a crystalline hydrochloride, nitrate, and oxalate, and yields a silver salt $\text{C}_{12}\text{H}_{12}\text{AgN}_2\text{O}_4$. It is decomposed by alkalis forming a compound $\text{C}_{11}\text{H}_{12}\text{N}_2\text{O}_5$, decomposing at 281°—33 *s-Trinitrobenzene* forms a compound $\text{NMe}_2\text{PhC}_6\text{H}_4(\text{NO}_2)_3$ [108°] which crystallises in long dark violet needles, sl sol alcohol (Hepp, A 215, 368)—34 *Tri nitro toluene* forms a corresponding body (H)—35 *Tri nitro aniline* forms $\text{NMe}_2\text{PhC}_6\text{H}_4(\text{NO}_2)_3(\text{NH}_2)$ [141°] (Hepp).

Salts—Dimethylaniline is apparently less basic than aniline, for if a mixture of the bases is treated with an insufficient quantity of cold aqueous HCl the base left uncombined is chiefly dimethylaniline (Morley, C J 51, 580)— $\text{B}'\text{H}_2\text{SO}_4$ [80°] Sol water, insol ether and benzene. HF 8400. The normal sulphate could not be formed. At 190° the acid sulphate splits up into CO_2 and the sulphonic acid (Vignon, C R 107, 263)— $\text{B}'_2\text{H}_2\text{HgCl}_2$ [149°] Large tables or prisms, m sol hot water and hot alcohol. Prepared by adding aqueous HgCl_2 to an alcoholic solution of dimethylaniline (Klein, B 11, 1741)— $\text{B}'_2\text{Hg}_2\text{OCl}_2$ Formed together with the preceding. Small needles or glistening leaflets, m sol hot water and hot alcohol, insol cold water, v sol benzene. On heating to 100° a blue colouring matter is formed— $\text{B}'_2\text{HgCl}_2$ (Leeds, J 1882, 504)— $\text{B}'_2\text{H}_2\text{SnCl}_4$ Large monoclinic crystals (Hjortdahl, J 1882, 523)— $\text{B}'_2\text{H}_2\text{PtCl}_6$ Tables— $\text{B}'_2\text{H}_2\text{PtCl}_6$ 2aq (Reinhardt & Staedel, B 16, 29)— $\text{B}'_2\text{H}_2\text{PtBr}_6$ Brownish-red monoclinic needles— $\text{Bi}_2\text{B}'\text{HI}$ (Kraut, A 210, 324)— $(\text{Bi}_2)_2\text{B}'\text{HI}$ — $(\text{Bi}_2)_2\text{B}'\text{HI}$ — $(\text{Bi}_2)_2\text{B}'\text{HI}$ — $\text{Bi}_2\text{B}'\text{HI}$ — $\text{B}'_2\text{H}_2\text{FeCl}_2$ Lamine, sl. sol cold water, being less soluble than the methylaniline

and aniline ferrocyanides (Fischer, *A* 190, 184) — $B^{\cdot}H_2FeCy_2$ 2aq (Eisenberg, *A* 205, 266)

Methylo-iodide $C_6H_5NMe_2I$ *Phenyl trimethyl-ammonium iodide* S (alcohol) 2 2 at 8° The combination of $PhNMe_2$ and MeI takes place rapidly (Lauth, *Bl* 7, 448) It may also be obtained by shaking a mixture of aniline (1 mol) with MeI (3 mols) and KOH (2 mols) dissolved in water (Pawlinoff, *J R* 13, 448, *Bl* [2] 37, 493) Plates (from alcohol) Converted by moist Ag_2O into $C_6H_5NMe_2OH$ a deliquescent, crystalline, caustic base When the methylo iodide is distilled it splits up into NMe_2Ph and MeI , but these recombine in the receiver If, however, a current of HCl be passed through the retort NMe_2PhHCl condenses in the receiver, while MeI escapes and may be condensed in a second colder receiver (Merill, *J pr* [2] 17, 286) When treated in a sealed tube with a solution of potassium in anhydrous liquefied ammonia it yields KI , NMe_2 , and possibly benzene (Thompson a Cundall, *C J* 53, 761) Boiling conc KOH slowly decomposes $PhNMe_2I$ into di methylaniline, KI , and $MeOH$ (Claus a Rautenberg, *B* 14, 621) The methylo iodide gives rise to the following derivatives — $PhNMe_2I_2$ [115°] Brown trimetric plates (Geuther, *A* 240, 69) — $PhNMe_2I_3$ [87°] (Dafert, *M* 4, 500) Lustrous green monoclinic needles, $a b c = 1.794 : 1.1251$, $\beta = 69^\circ 43'$ — $PhNMe_2I_4$ [65°] Violet black plates — $PhNMe_2IZnI_2$ Trimetric crystals (Hjortdahl) — $PhNMe_2ClHgCl_2$ [188°] S 36 at 62° Needles (Hubner, *A* 224, 352) — $(PhNMe_2Cl)_2PtCl_2$ Orange needles (from water) S 33 at 74° — $(PhNMe_2)_2Cr_2O_7$ Prisms S 5 — $(PhNMe_2)_2Cr_2O_{10}$ Monoclinic crystals (Hjortdahl)

Ethyl iodide $B^{\cdot}EtI$ [125°] Identical with methyl ethyl aniline methylo iodide, KOH splits off dimethylaniline (Claus a Howitz, *B* 17, 1325) — $PhNMe_2EtI_2$ [81°] Dark violet hexagonal rhombohedra (Geuther) — $PhNMe_2EtI_3$ [50°] Lustrous bluish-green plates — $PhNMe_2EtI_4$ [45°] Violet black plates

References — Bromo, Bromo di-nitro, Chloro, Chloro nitro, Iodo di-, Nitro, and Nitroso di methylaniline

DIMETHYLANILINE - AZYLINE v *Di-methyl-amido benzene* *azo-dimethylaniline*

DI-METHYL-ANILINE-PHTHALEIN v **TETRA-METHYL-DI-AMIDO-DI-PHENYL-PHTHALIDE**

METHYLANILINE SULPHONIC ACID v. **METHYLAMIDOBENZENE SULPHONIC ACID**

DIMETHYLANISIDINE v *o* **DIMETHYL-AMIDO-PHENOL**

METHYL-ANISOL is the methyl ether of CRESOL

(*B* 1) **METHYL-ANTHRACENE** $C_{14}H_{12}$, t.e.
 $C_6H_5 \begin{array}{c} \diagup CH \\ | \\ \diagdown CH \end{array} C_6H_5Me \left[\begin{array}{c} 6 \\ 5 \end{array} \right]$ Mol w 192 [203°]

(Börnstein), [200°] (Burukoff) Formed by distilling *erythro-oxo* methyl anthraquinone with zinc dust (Burukoff, *B* 20, 2070) Formed also by boiling *iso* methylanthraquinone with zinc dust and ammonia, and splitting off water from the resulting methyl hydroxanthranol by boiling with xylene (Börnstein, *B* 15, 1821) White plates — $B^{\cdot}C_6H_5(NO_2)_2OH$ red needles Gives on oxidation methyl-anthraquinone [167°] and

the corresponding anthraquinone carboxylic acid. Forms a di bromo methyl anthracene [148°]

Hexahydride $C_{14}H_{14}$ [α 66°] Formed by reducing the dilactone of benzophenone dicarboxylic acid with phosphorus and hydric iodide (Graebe a Juillard, *A* 242, 256) Plates V sol alcohol, ether, and chloroform Passed through a red hot tube it yields a hydrocarbon [195°] On oxidation it yields methylanthraquinone [154°]

(*B* 2) **Methyl-anthracene**

$C_6H_5 \begin{array}{c} \diagup CH \\ | \\ \diagdown CH \end{array} C_6H_5Me \left[\begin{array}{c} 6 \\ 5 \end{array} \right] 2$ [200°] Occurs in

coal tar oil, and hence is present in crude anthracene and phenanthrene (Japp a Schultz, *B* 10, 1049)

Formation — 1 By passing di-tolyl methane through a red hot tube (Weiler, *B* 7, 1181) — 2 By passing di tolyl ethane through a red hot tube (O Fischer, *B* 7, 1191, 8, 675) — 3 Apparently occurs among the products obtained by passing vapour of oil of turpentine through a red-hot tube (Schultz, *B* 10, 84) — 4 By the action of zinc dust at high temperatures on chrysophanic acid $C_{14}H_8O_2(OH)_2$, on emodin $C_{14}H_8O_4(OH)_2$ (Liebermann, *B* 8, 970, *A* 183, 163), on aloin (in small quantity) (E Schmidt, *B* 8, 1275, *Ar Ph* [3] 8, 496), on the acid [4 1] $CH_3C_6H_4CO_2C_6H_4CO_2H$ (Gresly, *A* 234, 238), on abietic acid (colophony), on gum benzoin (Ciamician, *B* 11, 273), on chrysarobin (Liebermann a Seidler, *A* 212, 34), and on methyl quinizarin (Nietzki, *B* 10, 2013) — 5 By boiling phenyl xylol ketone $C_6H_5CO_2C_6H_4Me$ [1 2 4] for a long time (Elbs, *J pr* [2] 35, 472) The isomeric $C_6H_5CO_2C_6H_4Me$ [1 2 5] condenses on boiling to the extent of 10 to 20 p.c. to (*B* 2) methyl anthracene, dehydrating agents either stop the reaction or split off benzoic acid

Properties — Yellowish plates (from alcohol) Sublimes in large white plates, exhibiting blue fluorescence Sl sol alcohol, ether, and $HOAc$, v sol $CHCl_3$, benzene, and CS_2 Chromic acid in $HOAc$ oxidises it to anthraquinone carboxylic acid [282°] Conc HNO_3 added to its alcoholic solution forms methyl anthraquinone Bromine in CS_2 forms a di bromo derivative [156°] (Fischer), which yields, on further bromination, a tetra bromo derivative crystallising from toluene in needles The picric acid compound melts at [93°] (Gresly)

Di-methyl-anthracene $C_{14}H_{14}$, t.e.

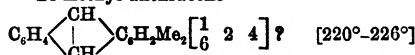
$C_6H_5 \begin{array}{c} \diagup CH \\ | \\ \diagdown CH \end{array} C_6H_5Me_2$ [71°] (above 360°) VD

7 19 One of the products obtained by passing benzyl mesitylene through a red hot tube Separated from the di methyl anthracene [218°] which accompanies it in smaller quantity, by crystallisation from toluene, in which the compound, melting at 71°, is extremely soluble Further purified by means of its picric acid compound (Louise, *Bl* [2] 44, 180, *A Ch* [6] 6, 191) White needles, v e sol cold benzene and toluene, m sol $HOAc$ and alcohol On oxidation with CrO_3 in $HOAc$ it yields di methyl-anthraquinone [158°] The picric acid compound forms long red needles. Forms a grey compound with nitro anthracene.

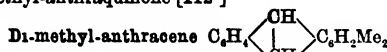


[219°] One of the products of the passage of benzyl-mesitylenethrough a red hot tube (Louise, *Bl* [2] 44, 178) Rhomboidal plates, insol cold alcohol, sl sol ether, ligroin, and acetic acid, v sol hot benzene and toluene. Forms small colourless leaflets when sublimed. Br in CS_2 forms a crystallised bromo-derivative. The picric acid compound crystallises in red needles. CrO_3 in HOAc oxidises it to a di-methyl anthraquinone [170°]. Forms a green compound with nitro-anthracene.

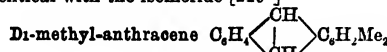
Di-methyl-anthracene



Obtained by distilling with zinc dust the tri-ox-di-methyl-anthraquinone derived from *m* xylene carboxylic acid, gallic acid, and H_2SO_4 (Birukoff, *B* 20, 871) Plates. Oxidised by CrO_3 to a di-methyl-anthraquinone [112°]

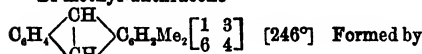


[218°] Obtained by treating a mixture of toluene (3 pts), CS_2 (3 pts), and chloroform (1 pt) with $AlCl_3$ (2 pts) (Elbs a Wittich, *B* 18, 348). With chromic acid it gives a quinone [162°]. Probably identical with the isomeride [219°]



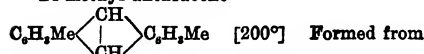
[203°] From [2 4 1] $C_6H_4Me_2$, CO , C_6H_4 , CO_2H [1 2] by heating with red hot zinc-dust (Gresly, *A* 234, 238) Plates. Possibly identical with the isomeride [220°-226°]

Di-methyl-anthracene



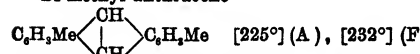
reduction of di-methyl-anthraquinone [183°] with zinc-dust and NH_3 . White plates, with bluish-green fluorescence. Its picrate forms garnet-red needles, decomposed by alcohol (Elbs a Eurich, *B* 20, 1863)

Di-methyl-anthracene



coal tar xylene by chlorination at boiling temperature, and subsequently heating the resulting [3 1] $C_6H_4(CH_3)(CH_2Cl)$ with water at 210° (Van Dorp, *B* 5, 674) White fluorescent plates. Yields on oxidation with chromic acid a quinone [158°]. Br gives a di-bromo di-methyl anthracene [154°]. Perhaps identical with the following isomeride

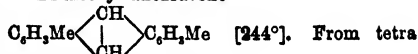
Di-methyl-anthracene



a C) Occurs in coal-tar (Zincke a Wachen-dorff, *B* 10, 1481) Formed from toluene, $AlCl_3$, and acetylene tetra-bromide (Anschütz, *A* 235, 172) Prepared by acting on toluene with methylene chloride in presence of $AlCl_3$ (Friedel a Crafts, *Bl* [2] 41, 323, *A Ch* [6] 11, 266). Scales. With CrO_3 in HOAc it gives dimethyl-anthraquinone [160°], which forms an orange

solution in H_2SO_4 . The picric acid compound crystallises in slender needles

Di-methyl-anthracene



methyl anthracene dihydride,

$C_6H_4Me \begin{array}{c} \diagup CHMe \\ \diagdown CHMe \end{array} C_6H_4Me$ by distillation over red hot zinc dust (Anschutz, *A* 235, 320), Greenish-yellow laminae (from benzene), m sol benzene, sl sol alcohol. Gives on oxidation di-methyl-anthraquinone [238°]

s Di-methyl-anthracene dihydride $C_{16}H_{16}$, i.e.

$C_6H_4 \begin{array}{c} \diagup CMeH \\ \diagdown CMeH \end{array} C_6H_4$ [181°] From ethylidene bromide, benzene, and $AlCl_3$ (Anschutz, *A* 235, 305) Yellow laminae (from alcohol) Sublimes in yellow needles. V sol benzene, CS_2 , and ether, sl sol cold alcohol and glacial HOAc. Distillation over red hot zinc dust gives anthracene. Oxidation gives anthraquinone. Bromine in HOAc gives $C_6H_4 \begin{array}{c} \diagup CMeBr \\ \diagdown CMeBr \end{array} C_6H_4$, which crystallises from toluene in needles

Picrate $C_{16}H_{16} \cdot C_6H_3(NO_2)_3OH$ [174°]

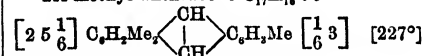
u-Di-methyl-anthracene dihydride

$C_6H_4 \begin{array}{c} \diagup CH \\ \diagdown CMe_2 \end{array} C_6H_4$ [56°] Formed by the action of HI and red phosphorus on dimethyl-anthron $C_6H_4 \begin{array}{c} \diagup CO \\ \diagdown CMe \end{array} C_6H_4$ at 150° (Hallgarten, *B* 21, 2508) White crystals, sol ether, benzene, and HOAc



[1 2 3 5] [236°] (W), [243°] (G) Obtained by distilling 'ψ cumene phthaloylo' acid $C_6H_4Me_2$, CO , C_6H_4 , CO_2H with zinc dust (Gresly, *A* 234, 239) Formed also by distilling tri-ox-di-methyl anthraquinone [244°] over zinc dust (Wende, *B* 20, 868) Exhibits green fluorescence

Tri-methyl-anthracene $C_{17}H_{14}$, i.e.



Formed by boiling di-*p* xylol ketone $C_6H_4Me_2$, CO , $C_6H_4Me_2$ for six hours, H O being eliminated. Colourless plates, with bluish green fluorescence. Sublimes below 100°, slightly volatile with alcohol. V sl sol cold alcohol, v sol ether. By CrO_3 and acetic acid it is oxidised to tri-methyl anthraquinone [184°] (Elbs a Olberg, *B* 19, 409, *J pr* [2] 35, 483)

Tetra-methyl-anthracene $C_{18}H_{14}$, [c 280°] A small quantity (3 g) is formed from *m* xylene (100 g), Al_2Cl_6 , and acetylene tetrabromide (Anschutz, *A* 235, 173) CrO_3 gives a substance (tetra-methyl anthraquinone?) which forms needles [c 300°]

Tetra-methyl-anthracene (?) [280°] From *o*-xylene, Al_2Cl_6 , and acetylene tetrabromide (Anschutz, *A* 235, 175) Fluorescent needles

Tetra-methyl-anthracene (?) [280°] Formed similarly from *p* xylene (A)

Tetra-methyl-anthracene $C_{18}H_{14}Me_2$, [163°]. Formed by the action of methylene chloride in presence of $AlCl_3$ on *m*-xylene, and in smaller

quantity on ψ -cumene (Friedel & Crafts, *A Ch* [6] 11, 268) Crystallises from benzene Gives a dark red crystalline compound with picric acid Conc H_2SO_4 gives a yellow solution CrO_3 in HOAc oxidises it to tetra methyl anthraquinone [206°]

Tetra-methyl-anthracene dibromide

$C_6H_4Me_4 \begin{smallmatrix} <CBrMe> \\ <CBrMe> \end{smallmatrix} C_6H_4Me_4$ From the corresponding tetra methyl anthracene dihydride by bromination (Anschutz, *A* 235, 321) Yellow needles, decomposes when heated

Tetra methyl-anthracene dihydride

$MeC_6H_4 \begin{smallmatrix} <CMeH> \\ <CMeH> \end{smallmatrix} C_6H_4Me_4$ [171°] Obtained by the action of ethylidene chloride on toluene in presence of $AlCl_3$ (Anschutz, *A* 235, 317) Pale yellow trimetric laminae, $a, b, c = 675, 1, 924$ (from alcohol and HOAc), ν sol benzene, sl sol HOAc, ν sl sol alcohol Distillation over red hot zinc dust gives di methyl anthracene [244°] Oxidation gives di-methyl-anthraquinone [236°]

Picrate $C_{18}H_{12}O_6N_2$ (OH) [165°] Red glistening needles

Hexa-methyl-anthracene $C_{18}H_{12}Me_6$ [c 220°]

One of the products of the action of methylene chloride on ψ -cumene in presence of $AlCl_3$ (Friedel & Crafts, *A Ch* [6] 11, 272) Not volatile at 410° The alcoholic solution gives with picric acid a brownish black pp [203°] Conc H_2SO_4 forms a red solution, becoming colourless after absorbing moisture

References — DI BROMO METHYL-ANTHRACENE AND AMIDO METHYL ANTHRACENE DIHYDRIDE AND DIBROMIDE

DI-METHYL-ANTHRACHRYSONE ν Tetra

oxy di methyl-anthraquinone

DI METHYL ANTHRACYLAMINE ν Di-methyl anthramine

DI-METHYL ANTHRAFLAVIC ACID ν Di-oxy di methyl anthraquinone

METHYL ANTHRAGALLOLS ν (1 2 3) Tri-oxy di methyl anthraquinones

DI METHYL ANTHRAMINE $C_{18}H_{12}NMe_2$. *Di methyl anthracylamine* [155°] Formed by heating the methyl hydrate Thin golden plates Soluble in alcohol with a green fluorescence

Salts — $BHCl$ colourless plates, decomposed by water — $B'HCl, PtCl_4$ yellow pp

Methylo iodide $B'MeI$ [215° uncor]

Formed by heating anthramine with methyl iodide at 100° Flat needles, sol hot water, sl sol cold, nearly insol alcohol

Methylo chloride platinum salt

$B'_2Me Cl.PtCl_4$ yellow crystalline pp

Methylo hydrate $B'Me(OH)$ strongly alkaline Formed by the action of Ag_2O on the iodide, on boiling the aqueous solution it decomposes into di methyl anthramine and methyl alcohol (Bollert, *B* 16, 1636)

METHYL-ANTHRANILIC ACID ν AMIDO-TOLUIC ACID

(B 1) METHYL-ANTHRAQUINONE

$C_{18}H_{10}O_2$ ν $C_6H_4 \begin{smallmatrix} <CO> \\ <CO> \end{smallmatrix} C_6H_4Me$ [5 1] Mol w 222 [154°] (Graebe), [167°] (Birukoff), [176°] (Börnstein) Formed by oxidising (B 1)-methyl-anthracene with CrO_3 and HOAc (Birukoff, *B* 20, 2070) Formed also by oxidising

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(B 1) methyl anthracene hexahydride (Graebe *A* 242, 256) The same, or the following, methyl anthraquinone is a by product in the preparation of anthraquinone (Wachendorff & Zinke, *B* 10, 1485, Börnstein, *B* 15, 1820) Small needles (from dilute HOAc) ν e sol alcohol and benzene

(B 2) Methyl-anthraquinone

$C_6H_4 \begin{smallmatrix} <CO> \\ <CO> \end{smallmatrix} C_6H_4Me$ [6 2] [163°] (F), [172°] (E), [177°] (Römer & Link, *B* 16, 695)

Formation — 1 By warming an alcoholic solution of (B 2) methyl anthracene with nitric acid, ppg with water, and subliming (O Fischler, *B* 8, 675) — 2 In small quantity by boiling phenyl m -xylyl ketone (Elbs, *J pr* [2] 35, 471) 3 By warming [4 1] $CH_3, C_6H_4, CO, C_6H_4, CO, H$ with H_2SO_4 at 170° for ten minutes (Gresly, *A* 234, 239)

Properties — Yellow needles Sublimes in almost colourless needles ν sol alcohol, benzene, and HOAc (R & L), according to Fischer, however, it is sl sol these solvents Conc H_2SO_4 forms a blood red solution which becomes violet on heating When heated with zinc-dust it yields methyl anthracene [203°] With fuming H_2SO_4 it yields a disulphonic acid, which on fusion with potash forms di oxy methyl anthraquinone (methyl alizarin) [252°] (Fischer)

Isomeride of Methyl anthraquinone ν METHANTHRENE

Di methyl-anthraquinone $C_{18}H_{12}O_2$ ν $C_6H_4 \begin{smallmatrix} <CO> \\ <CO> \end{smallmatrix} C_6H_4Me_2$ [1 2 4] ν [153°] Obtained

by oxidising the corresponding di methyl anthracene [71°] Formed also by heating benzoyl mesitylenic acid $C_6H_3CO, C_6H_4Me, CO, H$ with P_2O_5 and subliming the product (Louise, *A Ch* [6] 6, 193, 228, *Bl* [2] 44, 181) Yellow acicular prisms (from chloroform acetone) or needles (by sublimation) Insol water, sl sol alcohol, sol chloroform and acetone With zinc-dust and KOH it gives an intense red colour This di methyl anthraquinone ought theoretically to be identical with those melting at 180 and 112° (ν infra)

Di methyl anthraquinone $C_{18}H_{12}O_2$ [170°] Obtained by oxidising di methyl anthracene [219°] with CrO_3 in HOAc (Louise, *A Ch* [6] 6, 189, *Bl* [2] 44, 180) Yellow needles (from alcohol) With zinc dust and KOH it gives a characteristic red tint, which disappears on heating

Di methyl anthraquinone

$C_6H_4 \begin{smallmatrix} <CO> \\ <CO> \end{smallmatrix} C_6H_4Me$ [1 3 4] [183°] Obtained by heating o -xylyl phenyl ketone carboxylic acid (o -xylyl- o -benzoic acid) [3 4 1] $C_6H_4Me, CO, C_6H_4, CO, H$ [2] with conc H_2SO_4 By HNO_3 (S G 1 2) at 220° it is oxidised to anthraquinone di carboxylic acid [340°] (Elbs & Eulich, *B* 20, 1361).

Di-methyl-anthraquinone

$C_6H_4 \begin{smallmatrix} <CO> \\ <CO> \end{smallmatrix} C_6H_4Me_2$ [1 2 4] [180°] Formed by heating m -xylyl phenyl-ketone- o -carboxylic acid (m -xylyl- o -benzoic acid) [2 4 1] $C_6H_4Me, CO, C_6H_4, CO, H$ [1 2] with conc H_2SO_4 , the yield is 60 to 70 p c of the theoretical (Gresly, *A* 234, 240, Elbs & Gunther, *B* 20, 1364) Small needles, sl sol benzene

U

and alcohol By dil HNO_3 it is oxidised to anthraquinone *m* di carboxylic acid [above 330°] Reduced by zinc dust and ammonia to a hydrocarbon [85°] which forms with picric acid reddish-brown scales [136°]

Di methyl anthraquinone

$\text{C}_6\text{H}_4 \begin{smallmatrix} \text{CO} \\ \diagup \quad \diagdown \\ \text{CO} \end{smallmatrix} \text{C}_6\text{H}_2\text{Me}_2 \left[\begin{smallmatrix} 1 \\ 6 \end{smallmatrix} \begin{smallmatrix} 2 \\ 5 \end{smallmatrix} \right]$ [118°] Formed by warming $\left[4 \frac{1}{2} \right] \text{C}_6\text{H}_2\text{Me}_2 \cdot \text{CO} \cdot \text{C}_6\text{H}_4 \cdot \text{CO}_2\text{H} \left[2 \frac{1}{2} \right]$ with conc H_2SO_4 at 120° (Gresly, *A* 234, 240)

Di methyl anthraquinone

$\text{C}_6\text{H}_4 \begin{smallmatrix} \text{CO} \\ \diagup \quad \diagdown \\ \text{CO} \end{smallmatrix} \text{C}_6\text{H}_2\text{Me}_2 \left[\begin{smallmatrix} 1 \\ 6 \end{smallmatrix} \begin{smallmatrix} 2 \\ 4 \end{smallmatrix} \right]$? [112°] Obtained by oxidising the di methyl anthracene [220°–226°] (Birukoff, *B* 20, 871)

Di methyl anthraquinone

$\text{C}_6\text{H}_4 \begin{smallmatrix} \text{CO} \\ \diagup \quad \diagdown \\ \text{CO} \end{smallmatrix} \text{C}_6\text{H}_2\text{Me}_2$ [162°] Obtained by oxidising di methyl anthracene [216°] (Elbs & Wittich, *B* 18, 348)

Di-methyl anthraquinone

$\text{C}_6\text{H}_2\text{Me} \begin{smallmatrix} \text{CO} \\ \diagup \quad \diagdown \\ \text{CO} \end{smallmatrix} \text{C}_6\text{H}_2\text{Me}$ [236°] Formed by oxidation of di methyl anthracene [244°] Pale yellow needles, sl sol HOAc , v sl sol alcohol (Anschutz, *A* 235, 321)

Di-methyl-anthraquinone

$\text{C}_6\text{H}_2\text{Me} \begin{smallmatrix} \text{CO} \\ \diagup \quad \diagdown \\ \text{CO} \end{smallmatrix} \text{C}_6\text{H}_2\text{Me}$ [155°] Obtained by oxidising the di methyl anthracene occurring in coal tar xylene (Wachendorff & Zincke, *B* 10, 1482) Small light yellow needles (from dilute alcohol) May be sublimed M sol alcohol, ether, and HOAc

Di-methyl-anthraquinone

$\text{C}_6\text{H}_2\text{Me} \begin{smallmatrix} \text{CO} \\ \diagup \quad \diagdown \\ \text{CO} \end{smallmatrix} \text{C}_6\text{H}_2\text{Me}$ [160°] Obtained by oxidising the di-methyl anthracene formed from toluene, methylene chloride, and AlCl_3 (Friedel & Crafts, *A Ch* [6] 11, 266) Forms an orange solution in H_2SO_4 Probably identical with the preceding

Tri-methyl anthraquinone

$\text{C}_6\text{H}_4 \begin{smallmatrix} \text{CO} \\ \diagup \quad \diagdown \\ \text{CO} \end{smallmatrix} \text{C}_6\text{HMe}_2$ [Me Me Me = 1 2 4] [161°] Formed by warming ψ cuminoyl benzoic acid C_6HMe , $\text{CO} \cdot \text{C}_6\text{H}_4 \cdot \text{CO}_2\text{H}$ for a short time with conc H_2SO_4 (Gresly, *A* 234, 240) Needles

Tri-methyl-anthraquinone

$\left[2 \frac{5}{6} \frac{1}{3} \right] \text{C}_6\text{H}_2\text{Me}_2 \begin{smallmatrix} \text{CO} \\ \diagup \quad \diagdown \\ \text{CO} \end{smallmatrix} \text{C}_6\text{H}_2\text{Me} \left[\begin{smallmatrix} 1 \\ 6 \end{smallmatrix} \begin{smallmatrix} 3 \end{smallmatrix} \right]$ [184°] Formed by oxidation of the tri methyl anthracene obtained from di *p* xylil ketone (Elbs & Olberg, *B* 19, 409)

References — AMIDO-, NITRO, and OXY-METHYL ANTHRAQUINONES

METHYL ANTHRAQUINONE CARBOXY-

LIC ACID $\text{C}_6\text{H}_4\text{O}_4 \cdot \text{C}_6\text{H}_2\text{Me} \begin{smallmatrix} \text{CO} \\ \diagup \quad \diagdown \\ \text{CO} \end{smallmatrix} \text{C}_6\text{H}_4 \cdot \text{CO}_2\text{H}$ [246°] Occurs among the products of the oxidation of coal tar di-methyl anthracene by CrO_3 in HOAc (Wachendorff & Zincke, *B* 10, 1483) Small needles (by sublimation); v. sol. hot alcohol

Di-methyl-anthraquinone carboxylic acid

$\text{C}_6\text{H}_4\text{O}_4 \cdot \text{C}_6\text{H}_2 \begin{smallmatrix} \text{CO} \\ \diagup \quad \diagdown \\ \text{CO} \end{smallmatrix} \text{C}_6\text{H}_2\text{Me}_2 \cdot \text{CO}_2\text{H}$ [240°] Formed by heating ψ -cuminoyl-benzoic acid $\text{C}_6\text{H}_2\text{Me}$, $\text{CO} \cdot \text{C}_6\text{H}_4 \cdot \text{CO}_2\text{H} \left[2 \frac{1}{2} \right]$ with fuming H_2SO_4 (Gresly, *A*. 234, 241). Small needles. V. sol.

alkalis, al. sol alcohol and benzene May be sublimed

DI-METHYL-ANTHRARUFIN v DI OXY DI-METHYL ANTHRAQUINONE

DI-METHYL-ANTHRONE $\text{C}_{16}\text{H}_{14}\text{O}$ s.s.

$\text{C}_6\text{H}_4 \begin{smallmatrix} \text{CO} \\ \diagup \quad \diagdown \\ \text{CMe}_2 \end{smallmatrix} \text{C}_6\text{H}_4$ [94°] Formed by the action of MeI and KOH on anthranol (Hallgarten, *B* 21, 2508) Transparent crystals, v sol benzene, ether, sl sol petroleum ether Converted by the action of HI and red phosphorus into di methyl anthracene dihydride

METHYL-ARBUTIN v ARBUTIN

METHYL ARSENATE Me_2AsO_3 (214°) SG 1.559 Formed from Ag_3AsO_4 and MeI (Crafts, *Bl* [2] 14, 99)

METHYL ARSENITE Me_2AsO_2 (129°) SG 1.428 VD 6.01. From AsCl_3 and NaOMe (Crafts, *Bl* [2] 14, 104) When arsenious acid is dissolved in aqueous NaOH and treated with alcoholic MeI the product is not mono methyl arsenite but methane arsonic acid $\text{CH}_3\text{AsO}(\text{OH})_2$ (Klinger, *A* 249, 149, cf vol 1 p 317)

METHYL ARSINE v Organic ARSENIC COMPOUNDS

METHYL-ATROLACTIC ACID v OXY TOLYL-PROPIONIC ACID

METHYL-ATROPIC ACID v PHENYL CROTONIC ACID

METHYL-AURIN $\text{C}_{20}\text{H}_{12}\text{O}_4$ aq A by product in the preparation of aurin (Zulkowsky, *A* 194, 131, 202, 210, M 3, 476) Brick red crystals with green lustre (from 60 p c alcohol) Its alcoholic solution is yellowish red, and becomes crimson on addition of alkalis It loses aq at 100°, but does not melt below 200° HCl added to its dilute alcoholic solution ppts $\text{C}_{20}\text{H}_{12}\text{ClO}_4$ as red crystals with blue reflex On heating with water in sealed tubes at 245° methyl aurin is split up into *p* cresol and di oxy benzo phenone Alkaline KMnO_4 oxidises methyl aurin to $\text{C}_{16}\text{H}_{10}\text{O}_4$ Potash fusion forms *p* oxy-benzoic acid Zinc dust and HOAc reduce it to methyl leucaurin $\text{C}_{20}\text{H}_{14}\text{O}_4$ which crystallises in long colourless needles H_2SO_4 forms red crystals of $(\text{C}_{20}\text{H}_{14}\text{O}_4)_2 \cdot \text{H}_2\text{SO}_4$ which are blue by reflected light Bromine in HOAc forms crystals of $\text{C}_{20}\text{H}_{12}\text{Br}_2\text{O}_4 \cdot \text{HBr}$ aq, which have a steel blue reflex

METHYL-AZELAIC ACID v OCTANE DI-CARBOXYLIC ACID

METHYL DIAZO-COMPOUNDS v Di Azo COMPOUNDS

TETRA-METHYL TETRAZONE $\text{C}_4\text{H}_2\text{N}_4$ s.s. $(\text{CH}_3)_2\text{N} \cdot \text{N} \cdot \text{N} \cdot \text{N} \cdot (\text{CH}_3)_2$ (130°) Prepared by the oxidation of dimethyl hydrazine in ethereal solution with HgO (Renouf, *B* 13, 2173) Only fluid Explodes with violence if heated above 130° Alkaline in reaction Reduces AgNO_3 to a silver mirror It is decomposed by boiling aqueous acids into dimethylamine, methylamine, formic acid, and nitrogen

Salts — The picrate $\text{B} \cdot \text{C}_6\text{H}_3(\text{NO}_2)_3\text{OH}$ forms yellow prisms V sol water, sl sol alcohol. The other salts are also easily soluble in water

METHYL- α -BARBITURIC ACID $\text{C}_4\text{H}_7\text{N}_3\text{O}_4$ Formed by treating nitro methyl-uracil

$\text{CO} \begin{smallmatrix} \text{NMe} \text{CH} \\ \text{NH} \text{CO} \end{smallmatrix} \text{CNO}$, with tin and HCl (Lehmann, *A* 253, 80) Needles, v sol cold water.

Di-methyl barbituric acid v *Di methyl derivative of BARBITURIC ACID*, vol 1 p 439

METHYL BENZAMIDES v *Benzoyl derivatives of METHYLAMINES*

METHYL-BENZENE v **TOLUENE**

Di-methyl-benzene v **XYLENE**

Tri-methyl-benzene v ψ **CUMENE**, **MESITYLENE**, and **HEMIMELLITHENE**

Tetra-methyl benzene v **DURENE**

Penta-methyl benzene C_7H_8 , i.e. C_6HMe_5 , Mol w 148 [52°] (231° i V) V D 5.27 (calc 5.12) H F 31,900 H C 1,554,100 (Stohmann, Kleber, a Langbein, *J pr* [2] 40, 82) One of the products of the action of MeCl on benzene or toluene in presence of AlCl_3 (Friedel a Crafts, *A Ch* [6] 1, 472, Ador a Rilliet, *B* 12, 332) Formed also in like manner by the action of MeCl and AlCl_3 on tri methyl benzenes at 100°–110°, the fraction (220°–235°) on crystallisation from alcohol deposits C_6Me_5 first, and the penta-methyl benzene remaining in the mother liquor may then be purified by means of its sulphamide (Jacobsen, *B* 20, 896) Penta methyl benzene is a by product in the formation of tetramethyl-benzene by the action of MeI and AlCl_3 on ψ cumene (Claus, *J pr* [2] 38, 231)

Properties—Flat prisms, v e sol alcohol

Reactions—1 **Bromine** in CHCl_3 forms $\text{C}_6\text{Me}_5\text{Br}$ [163°] (292° i V) (F a C, Jacobsen, *B* 20, 898)—2 AgNO_3 and vapour of Br forms di bromo c durene [202°] (Gottschalk, *B* 20, 3288)—3 ClSO_3H forms the sulphone and sulphochloride (Jacobsen)—4 Cold conc H_2SO_4 does not form the corresponding sulphonic acid, but yields c tetra methyl benzene sulphonic acid and hexa methyl benzene (J)—5 KMnO_4 forms benzene penta carboxylic acid (F a C)—6 Fuming nitric acid forms di nitro c tetra methyl benzene [178°] (Gottschalk, *B* 20, 3287) Dilute nitric acid acting on its solution in benzene produces tetra methyl benzoic acid [165°]—7 Heating with AlCl_3 forms isodurene, C_6Me_4 , and other hydrocarbons (Jacobsen, *B* 18, 340)—8 ClCONH_2 and AlCl_3 convert C_6HMe_5 dissolved in CS, into the amide of penta-methyl benzoic acid (Jacobsen, *B* 22, 1219)

Picric acid compound

$\text{C}_6\text{HMe}_5\text{C}_6\text{H}_3(\text{NO}_3)_3\text{OH}$ [131°] Golden yellow prisms

Hexa methyl-benzene $\text{C}_{12}\text{H}_{12}$, i.e. C_6Me_6 , Mol w 162 [164°] (F a C), [166°] (J) (264°) S (95 p c alcohol) 2 at 0° V D 5.73 (calc 5.61) (F a C), 5.58 (H) H C v 1,709,600 H C p 1,712,200 H F 36,800 (Stohmann, Kleber, a Langbein, *J pr* [2] 40, 84)

Formation—1 The final product of the action of MeCl on benzene or toluene in the presence of AlCl_3 (Friedel a Crafts, *A Ch* [6] 1, 467, *C R* 91, 257, Ador a Rilliet, *B* 12, 332)—2 From o-di chloro benzene, MeCl, and AlCl_3 at 100° (Friedel a Crafts, *A Ch* [6] 10, 411)—3 A by product in the formation of (1,2,3,4)-tetra methyl benzene by the action of MeI and AlCl_3 on ψ cumene (Claus, *J pr* [2] 38, 231)—4 By heating dimethylaniline methylo-iodide at 330° (Hofmann, *B* 5, 721)—5 Said to be a product of the action of fused ZnCl_2 on sugar (*O. J.* 38, 863, *D P. J* 237, 146)—6 Formed as a

by product in the preparation of cumidine by heating xylidine hydrochloride with methyl alcohol, also in small quantity by the action of methyl alcohol at a high temperature on aniline hydrochloride (Hofmann, *B* 13, 1729)—7 By dropping methyl alcohol upon strongly-heated ZnCl_2 (Le Bel a Greene, *Am* 2, 21)—8 By pouring acetone upon fused ZnCl_2 (H Greene, *C R* 87, 931)—9 By shaking crotonylene CMeCMe with H_2SO_4 (3 pts) diluted with water (1 pt) (Favorsky, *J pr* [2] 37, 384, Lwoff a Almédingen, *Bl* [2] 37, 493)—10 Together with other products by the prolonged action of conc H_2SO_4 upon durene (Jacobsen, *B* 19, 1211)—11 A product of the action of cold conc H_2SO_4 on penta methyl benzene (Jacobsen, *B* 20, 901)

Properties—Colourless plates (from alcohol) or prisms (from benzene), insol water, sl sol cold, v sol hot, alcohol, v sol benzene Forms iridescent plates on sublimation Does not dissolve in conc H_2SO_4

Reactions—1 **Bromine** at 100° yields $\text{C}_{12}\text{H}_2\text{Br}_2$ [255°] (F a C), [227°] (H)—2 KMnO_4 oxidises it at a low temperature to mellitic acid—3 Dilute nitric acid yields c tetramethyl benzene dicarboxylic acid (Jacobsen, *B* 22, 1216)—4 When heated with AlCl_3 at 200° a gas is given off and penta methyl benzene, durene (Friedel a Crafts, *C R* 100, 692), isodurene, tri methyl benzenes, xylenes, toluene, and benzene are formed (Jacobsen, *B* 18, 339)—5 PCl_5 forms hexa chloro hexa methyl-benzene (*q v*)

Picric acid compound

$\text{C}_6\text{Me}_6\text{C}_6\text{H}_3(\text{NO}_3)_3\text{OH}$ [169°] Golden plates

METHYL BENZENE CARBOXYLIC ACID v

TOLUIC ACID

Di methyl-benzene carboxylic acid v **MESITYLENIC** and **DI METHYL BENZOIC ACIDS**

Tri-methyl-benzene carboxylic acid v ψ **CUMINIC ACID**

Methyl-benzene dicarboxylic acid v **UVITIC ACID**, **METHYL ISOPHTHALIC ACID**, **METHYL TEREPHTHALIC ACID**, and **TOLUENE DICARBOXYLIC ACID**

Di-methyl-benzene dicarboxylic acid v **DI-METHYL-ISOPHTHALIC** and **DI-METHYL-TEREPHTHALIC ACIDS**

Tri-methyl-benzene dicarboxylic acid $\text{C}_6\text{HMe}_3(\text{CO}_2\text{H})_2$ Obtained by oxidising $\text{C}_6\text{HMe}_3\text{CO}_2\text{C}_2\text{H}_5$ with KMnO_4 in alkaline solution (Ador a Meyer, *J* 1879, 562) Needles (from water)— $\text{BaA}'\text{aq}$ Slender needles, almost insol water

Tetra-methyl-benzene dicarboxylic acid

$\text{C}_6\text{Me}_4(\text{CO}_2\text{H})_2$ [6 5 4 3 2 1] [249°] Formed by boiling C_6Me_4 with dilute HNO_3 (Jacobsen, *B* 22, 1215) Small needles (from hot water) or prisms (from alcohol) Yields c durene when distilled with lime— $\text{BaA}'\text{, 2aq}$

DI-METHYL-BENZENE GLYCOL v **DI-XYLYLENE**

METHYL-BENZENE SULPHINIC ACID v **TOLUENE SULPHINIC ACID**

Tri-methyl-benzene sulphinic acid

$\text{C}_6\text{HMe}_3(\text{SO}_2\text{H})_2$ [1.3 4.6] [98°] Long needles (from water)— NaA' — BaA' S 5 at 7°. Thin plates— AgA' (Radloff, *B* 11, 82)

METHYL-BENZENE SULPHONIC ACID v **TOLUENE SULPHONIC ACID**

Di methyl-benzene sulphonic acid v XYLENE SULPHONIC ACID

c-Tri-methyl benzene sulphonic acid
 $C_6H_3Me_3(SO_3H)$ [1 2 3 5] *Hemimellitene sulphonic acid* Formed by sulphonation of hemimellitene (Jacobsen, *B* 15, 1858, 19, 2617) Crystallises from dilute H_2SO_4 in six sided plates or tables (containing aq) — NaA' aq tables, v sol hot, m sol cold, water — $*BaA'$, brittle plates, v sl sol water

Amide $C_6H_3Me_3SO_2NH_2$ [196°] Prisms or needles Less soluble in alcohol than the amide of ψ -cumene sulphonic acid

Isomerides v ψ -CUMENE SULPHONIC ACID and MESITYLENE SULPHONIC ACID

Tetra methyl-benzene sulphonic acid v. DURENE SULPHONIC ACID

Penta-methyl-benzene sulphonic acid
 $C_5Me_5SO_3H$ Obtained by treating penta methyl-benzene with $ClSO_3H$ and treating the product with NaOH to convert the $C_5Me_5SO_3Cl$ first formed into $C_5Me_5SO_3Na$ (Jacobsen, *B* 20, 899) In the action of $ClSO_3H$ on C_5Me_5H a sulphone [98 5°], crystallising from petroleum ether in needles, is also formed The free acid, liberated by shaking the sodium salt with light petroleum and H_2SO_4 , at once undergoes hydrolysis, C_5H_5Me , dissolving in the petroleum

Salts — $A'Na$ tables (from neutral aqueous solution) or plates (from hot, dilute, NaOH), sl sol cold water, m sol hot water, insol cold, dilute NaOH — $A'K$ plates (from hot water) — $A'Ba$ plates, v sl sol hot water — $A'Ca$ plates — $A'Ag$ plates, v sl sol cold, m sol hot, water — $A'Cu$ greenish white tables, sl sol water

Chloride [82°] Plates (from EtOH), or prisms (from Et₂O), v sol EtOH and Et₂O

Amide [186°] Prisms, v e sol hot, m sol cold, EtOH, insol water $KMnO_4$ produces a sulphamic acid [c 265°]

METHYL - BENZGLYCOCYAMIDINE v BENZOREATININE**METHYL-BENZGLYCOCYAMINE v BENZ-UREATIN****TETRA METHYL BENZIDINE v. TETRA-METHYL-DI-*p* AMIDO DIPHENYL****METHYL-BENZYL v PHENYL TOLYL DIKETOONE****METHYL-BENZOIC ACID v TOLUIC ACID**

Di-methyl-benzoic acid $C_6H_4O_2$ *o,s*
 $C_6H_4Me_2CO_2H$ [6 2 1] *m-Xylene o-carboxylic acid* [97°-99°] Formed by fusing potassium *m*-xylene *c*-sulphonate with sodium formate (Jacobsen, *B* 11, 21) Short needles (from water) Forms *m*-xylene when distilled with lime

Di-methyl-benzoic acid
 $C_6H_4Me_2CO_2H$ [4 2 1] *m Xylene o carboxylic acid Xylylic acid Xyloic acid* [126°]. (267°) at 727 mm

Formation.—1 By the oxidation of ψ -cumene by dilute nitric acid, being separated from the accompanying $C_6H_4Me_2CO_2H$ [5 2 1] through the greater solubility of its Ca salt (Fittig & Laubinger, *A* 151, 269) —2 By treating bromo-*m*-xylene with sodium and CO_2 (Kekulé, *A* 187, 186) —3 By warming its nitrile with H_2SO_4 (85 p c) and heating the resulting amide with conc HCl aq at 170° (Birukoff, *B* 20, 871, *A* 240, 246) —4 By heating its diphenylamide

with HCl aq (*v infra*) —5 By decomposing its chloride (*v infra*) with water

Properties — Slender needles (from water) or monoclinic prisms (from alcohol) May be sublimed Almost insol cold water, sl sol hot water, v sol hot alcohol Gives *m*-xylene when heated with lime Chromic acid mixture oxidises it to $C_6H_4Me_2(CO_2H)_2$ [3 1 4]

Salts — CaA' , 2aq monoclinic prisms, v sol water — BaA' , 8aq ? — AgA'

Chloride $C_6H_4Me_2COCl$ [25 5°] (235°) Formed by passing $COCl_2$ into *m*-xylene containing $AlCl_3$, and heating to 100° (Ador & Meyer, *B* 12, 1968)

Amide $C_6H_4Me_2CONH_2$ [180°] Formed by warming the nitrile with H_2SO_4 (85 p c) Formed also by the action of NH_3 , $COCl$ on *m*-xylene in presence of $AlCl_3$ (Gattermann, *A* 244, 58) Formed also by passing gaseous $HCNO$ and HCl gas into *m*-xylene containing $AlCl_3$ on the water bath (Gattermann & Rossolymov, *B* 23, 1196) Long shining needles (from water) Almost insol cold water, v sol alcohol

Anilide $C_6H_4Me_2CONHPh$ [138 5°] Crystals, v sl sol hot water

Diphenylamide $C_6H_4Me_2CONPh_2$ [142°] Formed from *m*-xylene and NPh_2 , $COCl$ in presence of $AlCl_3$ (Lellmann & Bonhöffer, *B* 20, 2120) Monoclinic crystals, v e sol hot alcohol, HOAc, and chloroform, sl sol hot ether, v sl sol hot petroleum ether

Nitrile $C_6H_4Me_2CN$ [25°] (221°) SG 12 9871 (Hinrichsen, *B* 21, 3082) VD 4 64.

Formation.—1 By heating the formyl derivative of *m*-xylydine with zinc-dust, the yield being about 12 p c (Gassioovsky & Merz, *B* 18, 1012) —2 By distilling the corresponding $PO(OC_2H_5)_2$, with $K_2Cr_2O_7$, the yield being 15 p c (Kreysler, *B* 18, 1713) —3 From *m*-xylydine by Sandmeyer's reaction (Bruckoff *B* 20, 871) **Properties** — Triclinic crystals (from dilute alcohol) Volatile with steam Yields benzyl di methyl amine on reduction

Amidoxim $C_6H_4Me_2C(NH)NOH$ [178°] *Xylenylamidoxim* Formed by the action of hydroxylamine on the nitrile at 85° (E Oppenheimer, *B* 22, 2443) White needles, v sol alcohol, ether, chloroform, and hot water, sl sol cold water **Reactions**.—1 EtI acting on its Na derivative forms $C_6H_4Me_2C(NH)NOEt$ which crystallises in white needles, [172°], v sol alcohol, ether, chloroform, benzene, and boiling water, sl sol cold water Excess of HCl converts the ethyl derivative into $C_6H_4Me_2CClNOEt$, a yellow aromatic oil —2 *Chloral* forms a compound $CCl_2CHO(C_6H_4Me_2C(NH)NOH)$ [112°] crystallising in white scales, sl alcohol and ether but decomposed by water and dilute acids 3 $ClCO_2Et$ acting on the amidoxim in chloroform forms $C_6H_4Me_2C(NH)NOCO_2Et$ [143°] which crystallises in white needles, v sol alcohol, ether, and chloroform, sl sol ligroin It is converted by heat into $C_6H_4Me_2C \begin{smallmatrix} N & O \\ \diagdown & / \\ & NH \end{smallmatrix} CO$ which crystallises in needles [182°], v sol alcohol and ether —4 Potassium cyanate acting on the hydrochloride of the amidoxim forms $C_6H_4Me_2C(NOH)NHCO_2NH_4$ [153°] which crystallises from dilute alcohol in white scales, and forms an unstable platinumchloride —5. *Phenyl cyanate* at 100° reacts forming

$C_6H_5Me_2C(OH)NHCO NHPH$ [188°] which crystallises from alcohol in light yellow scales, sol acids, alcohol, ether, chloroform, benzene, and hot water—6 *Phenyl thio-carbimide* at 100° forms $C_6H_5Me_2C(OH)NHOS NHPH$ [150°], sol alcohol, ether, benzene, boiling water, and acids—7 Heating with As_2O_3 forms the azoxim $C_6H_5Me_2C \begin{smallmatrix} \text{N} \text{O} \\ \diagup \quad \diagdown \end{smallmatrix} > CCH_3$ [89°]—

8 *Succinic anhydride* forms the azoxim $C_6H_5Me_2C \begin{smallmatrix} \text{N} \text{O} \\ \diagup \quad \diagdown \end{smallmatrix} > CCH_2CH_2CO_2H$ which forms long white needles, [112°], v sol alcohol, ether, benzene, $CHCl_3$, and hot water, and yields crystalline salts

Acetyl derivative of the *amidoxim* $C_6H_5Me_2C(NOAc)NH_2$ [189°] Small white needles, v sol alcohol and chloroform, sl sol ether (Oppenheimer, B 22, 2445)

Benzoyl derivative of the *amidoxim* $C_6H_5Me_2C(NO_2)NH_2$ [158°] White crystals, v sol alcohol, ether, and chloroform, sl sol water and ligroin Converted by heat into $C_6H_5Me_2C \begin{smallmatrix} \text{N} \text{O} \\ \diagup \quad \diagdown \end{smallmatrix} > CCH_3$, which crystallises in faintly yellow scales, v sol alcohol, ether, and chloroform, volatile with steam (Oppenheimer, B 22, 2444)

Di-methyl benzoic acid $C_6H_5Me_2CO_2H$ [5 2 1] *p* *Xylene carboxylic acid* [132°] (268° ± V)

Formation—1 From bromo-*p* xylene by treatment with $ClCO_2Et$ and sodium amalgam, the resulting ether being then saponified (Jacobsen, B 14, 2111)—2 By oxidation of methyl xylyl ketone $CH_3COCH_2Me_2$ [1 2 5] by dilute HNO_3 or by $KMnO_4$ (Claus, B 18, 1858, 19, 3183)—3 From its amide

Properties—Large needles (from alcohol), v sl sol hot water, v e sol alcohol Volatile with steam Yields $C_6H_5Me(CO_2H)_2$ [2 5 1] on oxidation

Salts— CaA' , 2aq crusts, m sol water— BaA' , 4aq small needles

Amide $C_6H_5Me_2CONH_2$ [186°] Formed by the action of $ClCO_2NH_2$ (or of $HNCO$ and HCl) on *p* xylene in presence of $AlCl_3$ (Gattermann, A 244, 54, B 23, 1199) Colourless needles (from water), m sol hot water, v e sol alcohol

Di-methyl-benzoic acid $C_6H_5Me_2CO_2H$ [3 2 1] *o* *Xylene c* *carboxylic acid* *Hemimellitethylic acid* [144°] Formed by the oxidation of hemimellitene $C_6H_5Me_2$ [3 2 1] by dilute nitric acid (Jacobsen, B 19, 2518) Prisms (from hot alcohol) or plates (from dilute alcohol) Volatile with steam, v sl sol hot water, insol cold water Yields *o*-xylene when distilled with lime CaA' , aq Long prisms, m sol cold water

Di-methyl benzoic acid $C_6H_5Me_2CO_2H$ [4 3 1] *o* *Xylene t* *carboxylic acid* [165°] Obtained by boiling its amide for 2 days with caustic potash solution Formed also, together with the isomeric *m*-xylene *t* *carboxylic acid* [126°] (*u supra*), by oxidising ψ cumene with dilute HNO_3 (Fitting a Laubinger, A 151, 275) Its ether is formed by the action of sodium-amalgam on a mixture of bromo-*o* xylene and $ClCO_2Et$ (Jacobsen, B 17, 2874) Prisms (from alcohol), almost insol cold water, v sl sol boiling water, v e sol alcohol Gives *o* xylene when distilled with lime Gives $C_6H_5Me(CO_2H)_2$

[3 4 1] on oxidation with dilute nitric acid— CaA' , 3aq Needles— BaA' , 4aq Needles, v sol water

Amide $C_6H_5Me_2CONH_2$ [131°] Formed by the action of NH_2COCl on *o* xylene in presence of $AlCl_3$ (Gattermann, A 244, 52) Long shining needles (from water) The amide obtained from *o* xylene, gaseous cyanic acid, HCl , and $AlCl_3$, melts at 165° (Gattermann a Rossolymo, B 22, 1199)

Diphenylamide $C_6H_5Me_2CONPh_2$ [136°] From NPh_2COCl , *o* xylene, and $AlCl_3$ (Lellmann a Bonhoffer, B 20, 2119) Small prisms v sol hot alcohol, sl sol ether and ligroin

Nitrile $C_6H_5Me_2CN$ (232°) VD 4 61 Formed by fusing potassium *o* xylene sulphonate with KCy (Jacobsen, B 11, 23) Formed also by heating the corresponding xylenyl phosphate $PO(OC_6H_5Me_2)_3$ with KCy , the yield being 20 p c (Kreysler, B 18, 1711) Liquid, miscible with alcohol and ether

Di-methyl-benzoic acid $C_6H_5Me_2CO_2H$ [5 3 1] is described as *MESITYLENIC ACID*

Di-methyl-benzoic acid $C_6H_5Me_2CO_2H$? *Lauroxylic acid* [155°] Produced by the action of dilute nitric acid on laurene $C_{11}H_{18}$ (Fitting, Köbrich, a Jilke, A 145, 151) Hard nodules (from alcohol) Nearly insol cold, sl sol boiling, water, v sol alcohol Yields acetic acid on oxidation with chromic acid mixture— BaA' , 4aq Concentrically grouped needles, v sol water— CaA' , 4aq— AgA' White pp, may be crystallised from water

Tri-methyl-benzoic acid ψ *CUMINIC ACID* Another tri methyl-benzoic acid, *Hemimellitene carboxylic acid*, is described as *c-CUMINIC ACID*

Tetra-methyl-benzoic acid $C_6HMe_4CO_2H$ [5 4 3 2 1] [165°] (G), [150°] (C) (c 270°) Formed by oxidising penta methyl benzene, dissolved in benzene, with dilute nitric acid (Gottschalk, B 20, 3286) Formed also by oxidising tetra methyl-phenyl methyl ketone or tetra methyl phenyl acetic acid with $KMnO_4$ (Claus, J pr [2] 38, 234) Needles (G), m sol hot water, v e sol alcohol, ether, benzene, CS_2 , and chloroform (C) Its alkaline salts are very soluble The Ag salt is v sl sol water The cupric salt forms light-green plates— BaA' , 2aq Plates or nodules, v sol water and alcohol (G)— BaA' , 6aq (C)— CaA' , 3aq (G)— NaA' 3aq

Tetra-methyl-benzoic acid $C_6HMe_4CO_2H$ [6 4 3 2 1] Formed by oxidising the corresponding $C_6HMe_4CO_2CH_3$ with $KMnO_4$ (Claus a Foecking, B 20, 8103) Yellow liquid, not solidified at 0°

Tetra-methyl-benzoic acid $C_6HMe_4CO_2H$ [6 5 3 2 1] [109°] (C a F), [112°] (G) Formed by oxidising the corresponding $C_6HMe_4CO_2CH_3$ with $KMnO_4$ (Claus a Foecking, B 20, 8103) Formed also by boiling its amide for several days with alcoholic potash (G) Plates

Amide $C_6HMe_4CONH_2$ [173°] From durenene, $ClCONH_2$, and $AlCl_3$ (Gattermann, A 244, 55) Shining plates (from dilute alcohol)

Tetra-methyl-benzoic acid $C_6HMe_4CO_2H$ [179°] From durenene $COCl_2$, and $AlCl_3$, the product being decomposed by water (Jacobsen, B 22, 1223) Also from $C_6HMe_4CO_2C_2H_5$ and $KMnO_4$ (Meyer a Ador, J 1879, 562) Must be identical with one of the three preceding acids,

probably with that of Gottschalk Large plates (from dilute alcohol), v sl sol cold water Volatile with steam May be distilled Conc HClAq at 210° yields durenene and CO₂—CaA', Short prisms, sl sol hot water—BaA', 4aq Small plates (from hot water)

Methyl ether C₆HMe, CO₂Me [59°] (269° v) Plates (Jacobsen, B 22, 1223) Saponified by alcoholic potash at 210°

Nitrile C₆HMe, CN [77°] Formed by distilling tetra methyl benzoic acid with PbC₂ (Jacobsen, B 22, 1224) Needles, v e sol alcoh. Split up by HCl at 215° into durenene and CO₂. A crystalline nitrile C₆H(CH₃), CN [69°], (260°), probably identical with the last is formed by the isomeric change of the carbamine derived from duridine (q v) by distillation It is very stable towards HCl, by which it is not saponified to the acid, but at 250° it decomposes with formation of tetra methyl benzene (Hofmann, B 17, 1914)

Amide C₆HMe, CONH₂ [173°] Formed by passing gaseous cyanic acid and HCl into durenene containing AlCl₃ at 100° (Gattermann a Rossolymo, B 23, 1199) It is probably the amide of the acid [179°], but has not been fully examined

Penta-methyl-benzoic acid C₆Me₅CO₂H [210 5°] Formed from penta methyl benzene, COCl₂, and AlCl₃ at 0°, after a fortnight the product is exposed to moist air and then treated with water (Jacobsen, B 22, 1220) Needles (from water) or prisms (from 70 p.c alcohol), v sl sol cold, sl sol hot, water, v e sol hot alcohol Volatile with steam May be distilled With H₂SO₄ it yields C₆Me₅ and c durenene (prehnitene) HNO₃ forms di nitro c durenene Fuming HClAq at 200° yields CO₂ and penta-methyl-benzene—CaA', Prisms, m sol water—BaA', 2aq Plates, sol hot water

Methyl ether MeA' [67 5°] (300° v) Plates, v sol alcohol

Amide C₆Me₅CONH₂ [206°] From C₆HMe₅, chloro formamide, and AlCl₃ Plates, sl sol hot, v sl sol cold, water

Nitrile C₆(CH₃), CN [170°] (J), [168°] (H) (292°) (H), (295°) (J) Formed from the carbamine C₆Me₅NC by intra-molecular trans formation by heating it a few degrees above its melting point (Hofmann, B 18, 1825) Large white needles, sol alcohol and ether, insol water It is remarkably stable, and could not be saponified by treatment with acids or alkalis. By heating with HI at 220°–230° it yields penta-methyl benzene, NH₃, and CO₂. Conc HClAq at 215° also yields C₆HMe₅.

METHYL-BENZOIC ALDEHYDE v TOLUIC ALDEHYDE

Di-methyl-benzoic aldehyde C₆H₂Me₂CHO [4 2 1] m *Xylobenzaldehyde* *Xylylic aldehyde*. [–8°] (216°) Formed by oxidising di methyl-benzyl alcohol with K₂Cr₂O₇ and H₂SO₄ (Hinrichsen, B 21, 3085, 22, 121) Separated by K₂CO₃. Colourless oil, volatile with steam HNO₃ oxidises it to di-methyl benzoic acid [126°]

Di-methyl-benzoic aldehyde C₆H₂Me₂CHO [5 3 1] (221°) Formed from mesitylene by treatment in CS₂ with CrO₃Cl₂, followed by water (Etard, C. R 97, 909) The yield is almost the theoretical quantity Yields mesitylene acid on oxidation.

Tri-methyl-benzoic aldehyde *Phenyl hydrazide* Me₃C₆H₂CH N₂HPH [129°] Formed by the action of tri methyl benzoic aldehyde on a solution of phenyl hydrazine in dilute HOAc (Rudolph, A 248, 100) Colourless needles, very sensitive to light, v sol ether, hot alcohol, and petroleum ether

DI-METHYL-BENZOIN C₆H₂O₂ v e [4 1] C₆H₂Me CO CH(OH) C₆H₂Me [4] [89°] p *Toluoin* Prepared by boiling 10 pts of p toluic aldehyde (from p xylene) with 2 pts of KCy and 30 pts of 50 p.c alcohol, and shaking the mass, after cooling, till the yellow product separates (Sterlin, B 22, 380, cf Grimaux a Lauth, B. 7, 233) Pale yellow prisms (from dilute alcohol), sl sol hot water, v sol alcohol, ether, and benzene Fuming H₂SO₄ gives a beautiful green colour

Acetyl derivative C₆H₂AcO₂ [100°] White crystals, v sol alcohol and ether
Benzoyl derivative C₆H₂BzO₂ [119°] White crystals, v sol alcohol and ether

METHYL BENZONITRILE v *Nitrile* of TOLUIC ACID

Di-methyl-benzonitrile v *Nitrile* of Di-METHYL BENZOIC ACID

METHYL-BENZOPHENONE v PHENYL TOLYL KETONE

Di-methyl-benzophenone v PHENYL XYLIL KETONE and Di TOLYL KETONE

TRI-METHYL-BENZOQUINONE CARBOXYLIC ACID v ψ-CUMOQUINONE CARBOXYLIC ACID

METHYL-BENZOYL-ACETIC ACID v. a BENZOYL PROPIONIC ACID

Di-methyl-benzoyl-acetic acid [5 2 1] C₆H₂Me₂CO CH₂CO₂H *Xylyl methyl ketone carboxylic acid* [132°] Formed, together with di methyl benzoic acid, by oxidising xylyl ethyl ketone with very dilute KMnO₄ (Claus a Fickert, B 19, 3183) Needles, sl sol water, v sol alcohol and ether—NaA' aq crusts, v sol water—CaA', 2½aq needles, sl sol cold water—BaA', 4aq prisms, sl sol cold water—AgA'

TETRA-METHYL-BENZOYL-BENZOIC ACID C₆HMe₄CO₂C₆H₂CO₂H [2 1] *Duroyl-benzoic acid* [c 260°] Formed by heating phthalic anhydride with durenene in presence of AlCl₃ (Friedel a Crafts, A Ch [6] 14, 454, C R 92, 833) Insol water, v sol alcohol, ether, and benzene May be crystallised from HOAc The Pb, Cu, and Ag salts are insol water—KA' minute needles, v sol cold water—NaA', v sl sol alcohol—NH₄A' needles—BaA', aq groups of needles, v sl sol water—CaA', aq needles, v sl sol water

METHYL-BENZOYL-ETHYL-HYDROXYLAMINE v HYDROXYLAMINE DERIVATIVES

TRI-METHYL-BENZOYL-PROPIONIC ACID v ψ CUMYL ETHYL KETONE-CARBOXYLIC ACID

METHYL-DIBENZYL v PHENYL TOLYL-ETHANE

METHYL-BENZYL-ACETIC ACID v PHENYL-250 BUTYRIC ACID

DI-METHYL-BENZYL ALCOHOL C₆H₂O v e [4 2 1] C₆H₂Me₂CH₂OH *Xylyl alcohol* [22°] (232°) Formed by adding KNO₃ to a solution of dimethyl benzyl amine sulphate (Hinrichsen, B 21, 3085) Colourless liquid, with aromatic odour, volatile with steam Oxidised by chromic acid mixture to di-methyl-benzoic aldehyde

Acetyl derivative $C_6H_5MeCH_2OAc$ (230°–234°).

Benzoyl derivative $C_6H_5MeCH_2OBz$ (333°)

Yellow oil (Hinrichsen, *B* 22, 123)

Penta-methyl-benzyl alcohol $C_6H_5MeCH_2OH$ *Mesityl alcohol* [160.5°] Obtained by saponifying its acetyl derivative with alcoholic potash (Jacobsen, *B* 22, 1217) Dimeric prisms, insol water, v sol alcohol

Acetyl derivative $C_6H_5MeCH_2OAc$ [85°] (310°) Obtained by the action of KOAc and HOAc on the chloride $C_6H_5MeCH_2Cl$, which is got by heating hexamethyl benzene with PCl_5 at 140° (Jacobsen) Plates or prisms (from alcohol), v sol ether, m sol alcohol at 0°, insol water

METHYL-BENZYL-AMINE

[2 1] $C_6H_5MeCH_2NH_2$ *o-Tolyl methyl-amine Xylylamine* (202°) Formed by heating methyl-benzyl phthalamide (derived from ω bromo-o xylene) with conc $HClAq$ in sealed tubes at 200° (Strassmann, *B* 21, 577) Liquid, which absorbs moisture and CO from the air— $B_2H_3PtCl_6$, yellow needles— $BHCl$ needles (from alcohol)—Picrate $B'C_6H_5N_2O_6$, long yellow needles, decomposing above 170°

Acetyl derivative $C_6H_5MeCH_2NHAc$ [69°] Formed by heating the hydrochloride with NaOAc and Ac_2O Crystallises from alcohol

m Methyl-benzyl-amine

[3 1] $C_6H_5MeCH_2NH_2$ (202°) Formed by heating ω bromo m xylene with potassium phthalamide, and decomposing the product with conc $HClAq$ at 190° (Bromme, *B* 21, 2700) Colourless liquid, miscible with alcohol and ether Absorbs CO_2 from the air— $BHCl$ needles $B_2H_3PtCl_6$, [212°] Golden plates—Sulphate [248°]— $B_2H_3C_2O_4$, [172°] Plates, v sol water—Picrate [156°]

Acetyl derivative $C_6H_5MeCH_2NHAc$ (235°–240°) Oil

Benzoyl derivative $C_6H_5MeCH_2NHBz$ [69°] White plates (from alcohol), v sol chloroform, HOAc, and benzene

p Methyl-benzyl-amine

[4 1] $C_6H_5MeCH_2NH_2$ Formed by treating an alcoholic solution of the amide of thio p toluo acid with zinc and $HClAq$ (Paterno a Spica, *B* 8, 441)

m (?) Methyl-benzyl-amine $C_6H_5MeCH_2NH_2$ *Xylylamine* (196°) Formed, together with di methyl di benzyl amine and tri methyl tri benzyl amine by heating ω chloro xylene with alcoholic NH_3 at 116° (Pieper, *A* 151, 129) Oil, smelling like herring brine Is either the m-compound or a mixture Lighter than water Absorbs CO_2 from the air— $BHCl$ [185°] Needles, v sol water and alcohol— $B_2H_3PtCl_6$

Di m (?) methyl-di-benzyl-amine $C_6H_5MeCH_2NH_2$ Formed as above (Pieper) Oil smelling like herring brine Lighter than water Decomposes above 210°— $BHCl$ [198°] Needles, sl sol cold water, v sol hot water and alcohol— B_2H_3Br [196°]

Tri m (?) methyl-tri-benzyl-amine $C_6H_5MeCH_2NH_2$ Formed as above (Pieper) Oil, sl sol alcohol Cannot be distilled With bromine water it yields $(C_6H_5MeCH_2)_3NH$ and C_6H_5MeCHO — $BHCl$ [212°] (P), [204°] (Jannasch, *A* 142, 803) Needles, sl sol cold alcohol, insol water and ether Heated in a

current of dry HCl it yields $(C_6H_5MeCH_2)_3NH$ and $C_6H_5MeCH_2Cl$ — $B'NO_3$, [122°] (J)

Di methyl-benzyl-amine $C_6H_5MeCH_2NH_2$ [4 2 1] $C_6H_5MeCH_2NH_2$ (219°) *Xylyl-methyl-amine Xylobenzyl amine* Formed by reducing with sodium the nitrile of the corresponding di methyl benzoic acid dissolved in alcohol (Hinrichsen, *B* 21, 3083, 22, 122) Liquid, sl sol water, v sol alcohol and ether Readily absorbs CO_2 and moisture from the air Nitrous acid converts it into the corresponding alcohol— $BHCl$ [210°] Needles or plates— $B_2H_3PtCl_6$, [228°]— $B_2H_3HgCl_2$, [205°]—Nitrate [158°]—Sulphate [254°]—Picrate [223°]— $B'CdI_2$ — $B'H_2CdI_2$

Benzoyl derivative $C_6H_5MeCH_2NHBz$ [98°] Needles, v sl sol water, v e sol alcohol and benzene

Di-methyl-benzyl amine v. BENZYL DI METHYL AMINE

METHYL-BENZYL ANILINE

$C_6H_5NMeCH_2Ph$ (306°) (N), (above 360°) (S) From methyl aniline and benzyl chloride (Noelting, *J* 1883, 702, Stebbins, *A C J* 7, 42). Pale yellow oil

p METHYL-BENZYL-BENZOIC ACID

$C_6H_5MeCH_2CO_2H$ [4 1] $C_6H_5MeCH_2CO_2H$, CO_2H [134°] Formed from $C_6H_5MeCO_2C_6H_5$, CO_2H by reducing with zinc dust and ammonia, filtering, diluting with water, adding HCl , dissolving the pp in NH_4Cl , and re ppg with acid (Gresly, *A* 234, 235) Long needles (from alcohol), v sol alcohol, HOAc, and benzene, almost insol water— BaA_2 plates, v sol water

Di-methyl benzyl benzoic acid $C_6H_5MeCH_2CO_2H$ [4 2 1] $C_6H_5MeCH_2CO_2H$, CO_2H [121°] [158°] Obtained by reducing $C_6H_5MeCO_2C_6H_5$, CO_2H with zinc dust and ammonia (Gresly, *A* 234, 237) Small needles (from alcohol)— BaA_2 , aq Plates (from dilute alcohol), sl sol water

METHYL-BENZYL BROMIDE v. o BROMO

XYLENE

DI METHYL-BENZYL CARBAMIC ETHER

$C_6H_5MeCH_2NHCO_2Et$ [6 4 2 1] $C_6H_5MeCH_2NHCO_2Et$ *Mesityl carbamic ether* [62°] From mesidine and $ClCO_2Et$ (Eisenberg, *B* 15, 1016) Long needles (from water) Volatile with steam

METHYL-BENZYL-CARBINOL v. PHENYL-

iso PROPYL ALCOHOL

PENTA-METHYL-BENZYL CHLORIDE

$C_6H_5MeCH_2Cl$ [99°] (c 285°) Formed by heating C_6H_5Me , (40 g) with PCl_5 (50 g) at 140° (Jacobsen, *B* 22, 1217) Plates (from alcohol ether), v e sol ether, sl sol alcohol

TRI METHYL BENZYL CYANATE

[6 4 2 1] $C_6H_5MeCH_2CO$ (219°) From tri methyl-benzyl carbamic ether and P_2O_5 (Eisenberg, *B* 15, 1017) Pungent liquid

METHYL-BENZYL GLYOXIM v. BENZYL-

METHYL GLYOXIM

DI- ω METHYL DI-BENZYLIDENE-ETHYL-

ENE DIAMINE $C_6H_5MeCH_2N_2$ + $C_6H_5MeCH_2N_2$ [c 104°] Formed by heating ethylene diamine (1 mol) with acetophenone (2 mols) to 120° White needles V sol alcohol and benzene, sl sol ether It is readily decomposed into its constituents, especially by acids (Mason, *B* 20, 273)

METHYL BENZYL KETONE v. BENZYL-METHYL-KETONE.

Methyl benzyl diketone $C_{11}H_{10}O_2$ *ze* $CH_3COCOCH_2C_6H_5$ (176°) SG $\frac{1}{2}$ 10721 This ketone is formed by distilling its monoxim $CH_3COC(NO)CH_2C_6H_5$ with $FeCl_3$ and dilute HCl (H Müller a Pechmann, B 22, 2133) Thick yellow oil, smelling like honey.

Di-phenyl-dihydrazide $CH_3C(N,HPH)CH_2C_6H_5$ [173°] Almost colourless flat silky needles

Mono oxim $CH_3COC(NO)CH_2C_6H_5$, **Isonitroso benzyl acetone** [81°] Formed by the action of nitrous acid on barium benzyl-acetoacetate (Ceresole, B 15, 1876, 3072, 16, 886) Needles, insol ligroin May be sublimed Dissolves in aqueous alkalis forming yellow solutions With $NaOEt$ and benzyl chloride it gives $CH_3COC(NOCH_2Ph)CH_2C_6H_5$, a thick yellow oil, volatile with steam

Di-oxim $CH_3C(NO)C(NO)CH_2C_6H_5$, **Methylbenzylacetoximic acid** [181°] Formed by adding an alcoholic solution of hydroxylamine hydrochloride to an alcoholic solution of the mono-oxim (Schramm, B 16, 181, 2185) Small needles With Ac_2O it yields a di acetyl derivative $CH_3C(NOAc)C(NOAc)CH_2C_6H_5$ [80°]

METHYL-BENZYL MALONIC ACID

BENZYL-METHYL-MALONIC ACID

o METHYL-BENZYL-PHTHALAMIC ACID

$C_{11}H_{11}NO_3$ *ze*

[1 2] $C_6H_4MeCH_2NHCOCH_2CO_2H$ [2 1] *o* **Xylyl phthalamic acid** [156°] Formed by boiling *o* methyl-benzyl phthalimide with aqueous $NaOH$ and ppg the solution with HCl (Strassmann, B 21, 576) Slender needles — AgA'

m-Methyl-benzyl-phthalamic acid

[1 3] $C_6H_4MeCH_2NHCOCH_2CO_2H$ [2 1] [131°] Formed in like manner from *m* methyl benzyl-phthalimide (Bromme, B 21, 2700) Slender needles (from hot alcohol) — AgA' *v* sol boiling water

o METHYL-BENZYL-PHTHALIMIDE

[1 2] $C_6H_4MeCH_2N<\begin{smallmatrix} CO \\ CO \end{smallmatrix}>C_6H_4$ [149°] Obtained by heating *o* bromo-*o*-xylene with potassium phthalimide at 200°, the yield being 76 p.c. of the theoretical amount (Strassmann, B 21, 576) White hexagonal crystals (from alcohol), insol water With conc $HClAq$ at 200° it forms methylbenzyl amine (202°) Aqueous $NaOH$ converts it into *o* methyl benzyl-phthalamic acid

m Methyl-benzyl-phthalimide

[1 3] $C_6H_4MeCH_2N<\begin{smallmatrix} CO \\ CO \end{smallmatrix}>C_6H_4$ *m*-Xylyl-phthalimide [118°] Formed in like manner by heating *o* bromo-*m*-xylene with potassium phthalimide at 190° (Brömme, B 21, 2700) Slender hexagonal needles (from hot alcohol), *v* sol ether, chloroform, and alkalis

METHYL-BENZYL-PIPERIDINE v BENZYL-METHYL-PIPERIDINE

METHYL BENZYL SULPHIDE $C_{11}H_{10}S$ *ze* $CH_3SCH_2C_6H_5$ (195°-198°) Formed by heating benzyl chloride with $Pb(SMe)_2$ at 100° (Obermeyer, B 20, 2926)

o Methyl-benzyl thiocarbimide C_8H_9NS *ze*, [2 1] $C_6H_4MeCH_2NCS$ *o* **Xylyl thiocarbimide** (256°) Formed by boiling an ethereal solution of *o*-methylbenzyl-amine with CS_2 and $HgCl_2$

(Strassmann, B 21, 578) Oil, smelling strongly like horse radish

Di-methyl-benzyl thiocarbimide $C_{10}H_{11}NS$ *ze* [4 2 1] $C_6H_4Me_2CH_2NCS$ **Xylobenzyl mustard oil** Formed by heating the corresponding urea ($C_6H_4Me_2CH_2NH_2$)CO with syrupy phosphoric acid (Hinrichsen, B 22, 123)

METHYL-BENZYL-THIO-UREA

$CH_3NHCSNHCH_2C_6H_5$ [74°] Formed by exposing an alcoholic solution of benzylamine mixed with methyl thiocarbimide for some days to the air (Dixon, C J 55, 619) White octahedra, *v* sol hot benzene and hot alcohol, *m* sol ether Gives a black pp with ammoniacal $AgNO_3$, but the aqueous or alcoholic solution is not desulphurised by prolonged boiling with alkaline lead tartrate

Methyl di-benzyl ψ thio-urea v Di BENZYL

METHYL- ψ -THIO UREA

o-Methyl-benzyl-thio-urea C_8H_9NS *ze*

[2 1] $C_6H_4MeCH_2NHCSNH_2$ *o* **Xylyl thio urea** [167°] Formed by evaporating a solution of potassium sulphocyanide and *o* methyl benzyl amine hydrochloride, and heating the residue to 140° (Strassmann, B 21, 578) White needles (from water), *m* sol hot water Turns red in air

m Methyl-benzyl-thio-urea

[3 1] $C_6H_4MeCH_2NHCSNH_2$ [112°] Prepared in the same way as the preceding body, using [3 1] $C_6H_4MeCH_2NH_2$ (Bromme, B 21, 2700)

Di m methyl-di-benzyl-thio-urea $C_{11}H_{12}N_2S$ *ze* ($C_6H_4MeCH_2NH_2$)CS [97°] Prepared by boiling *m* methylbenzyl amine with alcoholic CS_2 (B) Needles, insol water, acids, and alkalis

Tetra methyl-di-benzyl thio-urea $C_{15}H_{18}N_2S$ *ze* ($C_6H_4Me_2CH_2NH_2$)CS **Dizylobenzylsulphourea** [177°] From (4, 2, 1) di methylbenzyl amine and alcoholic CS_2 (Hinrichsen, B 22, 123) Glittering needles (from hot alcohol)

o METHYL-BENZYL-UREA $C_8H_9N_2O$ *ze*

[2 1] $C_6H_4MeCH_2NHCONH_2$ *o*-Xylyl urea [173°] Formed from *o* methylbenzyl amine sulphate and potassium cyanate (Strassmann, B 21, 578) Radiating moss like crystals (from alcohol), insol water

m Methyl-benzyl-urea

[3 1] $C_6H_4MeCH_2NHCONH_2$ [148°] Long needles from alcohol (Brömme, B 21, 2700)

Di-m-methyl-di-benzyl-urea

($C_6H_4MeCH_2NH_2$)CO [137°] Formed from *m* methylbenzyl-amine and $COCl_2$ in ethereal solution (B) Slender needles (from alcohol or ether)

Di-methyl-benzyl-urea $C_{11}H_{12}N_2O$ *ze*

[4 2 1] $C_6H_4Me_2CH_2NHCONH_2$ *m* **Xylobenzyl-urea** [184 5°] Formed by mixing concentrated solutions of (4, 2, 1) di methylbenzyl amine hydrochloride and potassium cyanate (Hinrichsen, B 22, 122) Slender needles (from alcohol), *m* sol water from which it separates as a flocculent pp

METHYL-BISMUTHINE v Bismuth methide under BISMUTH, ORGANIC DERIVATIVES

METHYL BORATES

Tri-methyl-borate Me_3BO_3 (72°) (E a B), (65°) (S) SG $\frac{1}{2}$ 955 (E a B), 940 (S) VD 3 66 (E a B) Prepared by heating B_2O_3 with methyl alcohol at 100° in sealed tubes, and distilling The distillate is freed from $MeOH$ by shaking with conc H_2SO_4 , and the upper

layer is then rectified (Schiff, *Bl* [2] 5, 372, 6, 36) Formed also by the action of BCl_3 on dry methyl alcohol, the upper layer of the product being rectified (Ebelmen a Bouquet, *A Ch* [3] 17, 59, A 60, 251) Colourless mobile liquid, sol alcohol and ether Burns with a green flame (greener than that of Et_2BO) Decomposed by water into boric acid and MeOH

Methyl metaborate MeBO A thick liquid, formed by heating Me_2BO_3 with B_2O_3 When heated, it begins to decompose at 160° giving off Me_2BO_3 , and at 250° there remains a mass of the composition MeB_2O_5 , which becomes vitreous on cooling

An acid methyl borate $\text{Me B}_2\text{O}_5$ was described by Ebelmen (*A Ch* [3] 16, 17) as obtained by treating B_2O_3 with dry MeOH , and as being a vitreous mass, readily decomposed by water into boric acid and MeOH It was probably a mixture of Me_2BO_3 and MeBO

METHYL-BORNYL UREA v **BORNYL-METHYL UREA**

TRI-METHYL-BRAZILIN $\text{C}_6\text{H}_{11}\text{Me}_3\text{O}_3 \cdot \frac{3}{2}\text{aq}$ Formed in the preparation of tetra methyl brazilin (v *infra*) Crystalline, sol dilute alcohol Its solution in aqueous NaOH is colourless and gives a brown pp with FeCl_3

Acetyl derivative $\text{C}_6\text{H}_{10}\text{AcMe}_2\text{O}_3$ [97°]

Tetra-methyl-brazilin $\text{C}_6\text{H}_{10}\text{Me}_4\text{O}_4$ [139°] Formed by the action of NaOEt and MeI on brazilin (Schall a Dralle, *B* 20, 3365, 21, 3009) Snow white crystals Bromine in HOAc forms $\text{C}_6\text{H}_8\text{Br}_2\text{Me}_2\text{O}_4$ [181°] and crystalline di bromo tetra methyl brazilin dibromide $\text{C}_6\text{H}_8\text{Br}_2\text{Me}_2\text{O}_4$

METHYL-BROMACETOL v *aa* **DI BROMO-IRAPANE**

METHYL BROMIDE CH_3Br *Bromo methane* Mol w 95 (45°) SG $\frac{4}{8}$ 1.7331, $\frac{12}{23}$ 1.7235 (Perkin), $\frac{8}{8}$ 1.732 (Merrill) VD ($H=1$) 95 MM 4.644 at 15° (Perkin, *C J* 45, 454) HFp 14,790 HFv 14,210 (Thomsen) SV 55.7 (Lossen, *A* 254, 68)

Preparation—Methyl alcohol (800 grms) and amorphous phosphorus (133 grms) are put into a retort Bromine (800 grms) is slowly run in After some hours the retort is heated and the product condensed in a receiver surrounded by a freezing mixture It is washed with dilute potash, dried over calcic chloride and distilled (Merrill, *J pr* 126, 296, cf Pierre, *J Ph* [3] 13, 156, Bunsen, *A* 46, 44)

Properties—Colourless, mobile liquid, with burning taste and pleasant smell resembling chloroform Burns with greenish brown smoky flame Miscible with alcohol, ether, chloroform, and CS_2 Poured into cold water a white ice like mass is formed It is approximately CH_3Br , 20aq At 5° it begins to split up with evolution of methyl bromide gas

METHYL-BROMO-ACETOACETIC ETHER v **BROMO ACETO-ACETIC ACID**

METHYL-BROMO-ACETOL v **DI BROMO PROPANE**

METHYL α -BROMO-ALLYLOXIDE $\text{C}_3\text{H}_5\text{BrO}$ v $\text{CH}_2\text{OCH}_2\text{CBrCH}_3$ (116°) SG $\frac{12}{13}$ 1.35 Formed by the action of NaOH on $\text{CH}_3\text{OCH}_2\text{CHBrCH}_3$ (Henry, *B* 5, 455)

METHYL BROMO-ALLYL SULPHIDE $\text{C}_3\text{H}_5\text{BrS}$ v $\text{CH}_3\text{SCH}_2\text{CH}_2\text{CHBr}$ Formed by heating $\text{Pb}(\text{SMe})_2$ with *s*-tri-bromo propane in

ethereal solution at 150° (Obermeyer, *B* 20, 2925) Decomposes at 120° – 130° Combines with Br (1 mol)

METHYL-BROMO-AMINE v **METHYLAMINE**

METHYL-DI-BROMO-ANTHRACENE v **LI-BROMO METHYL ANTHRACENE**

METHYL-DI-BROMO ATROLACTIC ACID v **DI BROMO OXY-TOLYL PROPIONIC ACID**

METHYL BROMO-BUTYL KETONE v **ACETYL BUTYL BROMIDE**

TRI METHYL-BROMO-ETHYL AMMONIUM BROMIDE v **TRIMETHYLAMINE bromo-ethyl bromide**

METHYL-BROMO ISATIN v *Methyl derivative of Bromo ISATIN*, p 71

METHYL BROMO ISATOYD v *Bromo methyl-isatoyd*, p 71

METHYL BROMOPROPYL KETONE

$\text{CH}_3\text{COCH}_2\text{CH}_2\text{CH}_2\text{Br}$ *Acetyl propyl bromide*. (118°) at 90 mm (P), (106°) at 60 mm (L)

Formed by treating acetyl propyl alcohol (q v) with a saturated aqueous solution of HBr (Colman a Perkin, jun, *C J* 55, 357, Lipp, *B* 22, 1196) Colourless mobile liquid with penetrating odour Turns brown in light Sl sol cold water, quickly decomposed by hot water, dissolving as acetyl propyl alcohol Forms a crystalline compound with NaHSO_4 , NaOEt and KOH act on it, forming a light ethereal oil $\text{C}_6\text{H}_8\text{O}$ (113°), probably methylene furfuran tetrahydride

METHYL BROMO-STYRENE v **BROMO TOLYLETHYLENE**

METHYL BRUCINE v *Methylo-compounds of BRUCINE*, vol 1 p 637

METHYL BUTENYL TRICARBOXYLIC ACID v **PENTANE TRICARBOXYLIC ACID**

METHYL BUTENYL KETONE v **ALLYL-ACETONE**

METHYL ISOBUTYL ACETAL v **ALDEHYDE**

METHYL-BUTYL-ACETIC ACID v **HEPTOIC ACID**

Methyl-di-butyl acetic acid v **HENDECIOIC ACID**

METHYL-BUTYL-ACETYLENE v **HEPTENF**

METHYL-ISOBUTYL ANILINE $\text{C}_6\text{H}_5\text{N} \cdot \text{C}_4\text{H}_8\text{NMeCH}_2\text{Pr}$ (235°) (Noelting, *J* 1883, 702)

METHYL ISOBUTYL-BENZENE v **ISOBUTYL TOLUENE**

METHYL-BUTYL-CARBINOL v **HEXYL ALCOHOL**

Di methyl butyl-carbinol v **HEPTYL ALCOHOL**

METHYL ISOBUTYL CARBONATE $\text{C}_4\text{H}_9\text{O}_2$ v $\text{CH}_3\text{O CO OC}_4\text{H}_9$ (143.6° cor) SG $\frac{12}{13}$ 0.95 (Rose, *A* 205, 230)

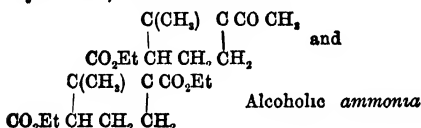
DI-METHYL-BUTYLENE-DIKETONE $\text{C}_6\text{H}_8\text{O}_2$ v $\text{CH}_3\text{COCH}_2\text{CH}_2\text{CH}_2\text{COCH}_3$

Di-acetyl butane (44°) Obtained by heating its dicarboxylic ether with NaOMe in MeOH (Marshall a Perkin, jun, *C J* 57, 241) Crystalline mass, sl sol water, v sol other menstrua Combines with NaHSO_4 Reacts with phenylhydrazine and with hydroxylamine Boiling alcoholic potash condenses it forming methylpenta methenyl trihydride methyl ketone $\text{CH}_3\text{C} \begin{smallmatrix} \diagup \text{C}(\text{COCH}_3) \\ \diagdown \end{smallmatrix} \text{CH}_3$

DI-METHYL-BUTYLENE-DIKETONE CARBOXYLIC ETHER $\text{C}_6\text{H}_8\text{O}_4$ v $\text{CH}_3\text{COCH}_2\text{CH}_2\text{CH}_2\text{CH}(\text{CO}_2\text{Et})\text{COCH}_3$

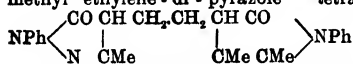
di-acetyl n valeric ether (195°–200°) at 100 mm. Obtained by distilling the corresponding di-carboxylic ether (di-acetyl adipic ether) with potash, neutralising the residue with H_2SO_4 , extracting with ether, washing the ether with water, drying and evaporating (Perkin, jun, *C J* 57, 229). Colourless oil, v sol ether and alcohol, sl sol water. Its alcoholic solution is coloured violet by $FeCl_3$. On hydrolysis it yields acetyl-valeric acid and acetic acid.

Di-methyl-butylene-diketone dicarboxylic ether $C_8H_{14}O_6$ *ie* $CH_3COCH(CO_2Et)CH_2CH_2CH(CO_2Et)COCH_3$, *Ethylene diaceto acetic ether* *Di acetyl adipic ether*. Formed by the action of sodium (46 g) on aceto acetic ether (260 g) and ethylene bromide (190 g), dissolved in alcohol (Perkin, jun, *C J* 57, 215). It may be purified by conversion into the yellow flocculent di sodium compound ($AcCNa(CO_2Et)CH_2$)₂, decomposing this with very dilute HCl , and extracting with ether. Thick oil, miscible with alcohol and ether, sl sol dilute aqueous KOH . $FeCl_3$ colours its alcoholic solution intense violet red. Combines with great difficulty with $NaHSO_4$. Decomposed by heat into di-methyl butylene diketone carboxylic ether, and the two ethers



converts the dicarboxylic ether into the diimide ($CH_2C(NH)CH(CO_2Et)CH_2$)₂ [174°] which is reconverted by warming with dilute HCl into the original ketonic ether.

Phenyl hydrazide $(CH_2C(N_2HPh)CH(CO_2Et)CH_2)_2$ [α 145°]. Formed by heating di-methyl butylene diketone carboxylic ether with phenyl hydrazine on the water bath (Perkin, jun, a Obremsky, *B* 19, 2049; Perkin, jun, *C J* 57, 221). Plates or needles (from $MeOH$), v sol conc $HClAq$, insol alkalis, v sl sol ether. At 200° it gives off $EtOH$ (2 mols), forming di-oxy di-phenyl dimethyl ethylene-di-pyrazole tetrahydride



METHYL-ISOBUTYL-GLYOXALINE *Methylo iodide* $C_8H_{14}N_2I_2$ *ie* $N_2C_2H_4MeC_2H_4MeI$ [170°]. Formed by treating isobutyl glyoxaline (glyoxal-isocamylne) with MeI in $MeOH$ (Radziszewsky, a Szul, *B* 17, 1294). Trimetric prisms (from alcohol).

METHYL-n-BUTYL KETONE $CH_3COCH_2CH_2CH_2CH_2COCH_3$ (127°) *SG* 2°830. Formed by the oxidation of sec hexyl alcohol (Erlenmeyer, a Wanklyn, *A* 135, 144; Schorlemmer, *A* 161, 263). Combines with $NaHSO_4$. Gives on oxidation acetic, n-butyric, and valeric acids (Wagner, *B* 18, 2267).

Methyl isobutyl ketone $CH_3COCH_2CH_2CH_2CH_2COCH_3$ (115°) *SG* 2°8195, 2°8034 (Wagner, *J R* 16, 703). A product of the distillation of potassium isovalerate with $NaOAc$ (Williamson, *A* 81, 86). Formed by the action on valeryl chloride of $ZnMe$, followed by water. Formed also by boiling isopropyl-aceto acetic ether with baryta water (Frankland, a Duppes, *A* 145, 82).

Combines with $NaHSO_4$. Yields on oxidation acetic, isobutyric, and isovaleric acids.

Methyl sec-butyl ketone $CH_3COCHMeCH_2CH_2CH_3$ (118°) *V* *SG* 14°5 8181. Formed by boiling methyl ethyl acetoacetic ether with dilute KOH or baryta water (Walsleben, *A* 219, 307; Wagner, *J R* 16, 711). Oil, smelling of peppermint. Oxidised by chromic acid mixture to methyl ethyl ketone and $HOAc$. Sodium reduces it to the corresponding hexyl alcohol and methyl sec butyl pinacone [249°].

Methyl tert-butyl ketone $CH_3COCHMe_2$, *Pinacolone* (105°) (Schiff, *B* 19, 562) *SG* 2°8265, 1°800.

Formation—1 By distilling pinacone with dilute sulphuric acid (Fittig, *A* 114, 56).—2 By the action of CMe_2COCl on $ZnMe_2$, followed by water (Butlerow, *A* 174, 125).—3 By the dry distillation of calcium isobutyrate (Barbaglia, a Gucci, *B* 13, 1572).

Properties—Liquid, smelling of peppermint, nearly insol water. Sodium amalgam forms the corresponding hexyl alcohol. Combines with $NaHSO_4$. Chlorine forms pungent $C_8H_{14}ClO$ crystallising in needles [51°], (178°).

Oxim $CH_3C(NOH)CMe_2$, [75°]. Slender needles, very volatile with steam (Janny, *B* 15, 2780). May be sublimed. Sl sol cold, v sol hot, water. Has a camphor like smell and taste. Not decomposed by acids.

Methyl isobutyl diketone $C_8H_{14}O_2$ *ie* $CH_3COCH_2CH_2CH_2CH_2COCH_3$, *Acetyl isovaleryl Diketoheptane* (138°) *SG* 2°908. Formed by distilling its mono oxim with dilute H_2SO_4 (Otte, a Pechmann, *B* 22, 2122). Yellow oil with irritating smell, becoming fruity when diluted. Sl sol water.

Mono oxim $CH_3COCH(NOH)CH_2CH_2CH_2CH_2COCH_3$, *Acetyl isobutyl acetone* [42°]. Formed by adding $NaNO_2$ (10 g) to a solution of isobutyl aceto acetic ether (27 g) in water (300 cc) and KOH (8 g), and extracting with ether (Treadwell, a Westenberger, *B* 15, 2786). White plates, v sol alcohol and ether, sl sol cold water, dissolves in alkalis.

Dioxim $CH_3C(NOH)C(NOH)CH_2CH_2CH_2CH_2COCH_3$ [c 172°]. Small white shining plates (from dilute alcohol) (O a P).

Oxim phenylhydrazide $C_{13}H_{18}N_2O$ *ie* $CH_3C(N_2HPh)C(NOH)CH_2CH_2CH_2CH_2COCH_3$ [151°]. Almost colourless needles.

Phenylhydrazide $C_{13}H_{18}N_2O$ [98°]. Almost colourless needles.

Diphenylhydrazide $C_{19}H_{24}N_4$ *ie* $CH_3C(N_2HPh)C(N_2HPh)CH_2CH_2CH_2CH_2COCH_3$ [116°]. Almost colourless needles.

METHYL n-BUTYL KETONE CARBOXYLIC $CH_3C_2H_4O_2$ *ie*

$CH_3COCH_2CH_2CH_2CH_2CO_2H$ *ω*-Acetyl valeric acid [42°]. A product of the hydrolysis of di-methyl butylene diketone carboxylic ether by a solution of KOH in methyl alcohol (Perkin, jun, *C J* 57, 231). Colourless crystals, v sol water, alcohol, and ether. Its ammonium salt is very soluble— AgA' leafy masses (from hot water). The cupric salt forms minute spherules.

METHYL-ISOBUTYL-KETONE SULFONIC ACID $CH_3COCH_2CH_2CH_2CH_2SO_3H$. The sodium salt of this acid is slowly formed by the action of a

saturated solution of NaHSO_4 on mesityl oxide. It is easily soluble in water and alcohol, melts at 95° , is not attacked by Na_2CO_3 , but by NaOH it gives mesityl oxide (Pinner, *B* 15, 592).

METHYL BUTYL KETOXIM *v.* Oxim of METHYL BUTYL KETONE

METHYL *n* BUTYL OXIDE $\text{C}_6\text{H}_{12}\text{O}$ *ie* $\text{CH}_3\text{O C}_4\text{H}_9$ (70.3°) SG 7635 SV 127.2 CE (0° - 10°) 00125 (Dobrenner, *A* 243, 3).

Methyl isobutyl oxide $\text{CH}_3\text{O C}_4\text{H}_9$ (60°). One of the products of the action of NaOCH_2Pr on methylene iodide or iodoform (Gorboff, *J pr* [2] 41, 238, 254). Oil. Conc HIAq yields MeI and isobutyl iodide.

METHYL-ISOBUTYL-PHENOL $\text{C}_{11}\text{H}_{14}\text{O}$ *ie* $\text{C}_6\text{H}_5(\text{CH}_2)(\text{C}_2\text{H}_5)\text{OH}$ [136] (236°). *Isobutyl o cresol*. Formed by the action of nitrous acid on methyl isobutyl phenyl amine (Effront, *B* 17, 2324). Oil. V sol alcohol and ether, nearly insol water.

Isomeride *v.* Methyl ether of ISOBUTYL-PHENOL

METHYL-ISOBUTYL PHENYL AMINE

$\text{C}_6\text{H}_5(\text{CH}_2)(\text{C}_2\text{H}_5)\text{NH}_2$ [136] *Isobutyl tolyl amine or toluisobutylamine* (243°). Formed by heating *o* toluidine hydrochloride with isobutyl alcohol at 200° - 300° (Effront, *B* 17, 2317). Colourless liquid. Volatile with steam. Miscible with alcohol and ether, nearly insol water.

Salts— B^+HCl . Long thin needles, sl sol cold water, v sol hot water— B^+HBr long soluble needles— $\text{B}^+\text{H}_2\text{SO}_4$ needles, sl sol cold water— $\text{B}^+\text{H}_2\text{C}_2\text{O}_4$ silvery needles, insol ether.

Formyl derivative $\text{C}_{11}\text{H}_{15}\text{NH(COH)}$ [106°]. Colourless tables, v sol alcohol and ether, nearly insol cold water.

Acetyl derivative $\text{C}_{11}\text{H}_{15}\text{NHAc}$ [162°]. Colourless plates, sparingly sol hot water, easily in alcohol.

Benzoyl derivative $\text{C}_{11}\text{H}_{15}\text{NHBz}$ [168°]. Small white needles, insol cold water, v sl sol hot.

METHYL-ISOBUTYL PHENYL-AMINE

$\text{C}_6\text{H}_5(\text{CH}_2)(\text{C}_2\text{H}_5)\text{NH}$ [156] (244°). Formed by heating *o* toluidine with isobutyl alcohol and ZnCl_2 at 270° - 280° (Effront, *B* 17, 2339). Colourless liquid. Volatile with steam.

Salts— B^+HCl . Plates— $\text{B}^+\text{H}_2\text{SO}_4$ plates— $\text{B}^+\text{H}_2\text{C}_2\text{O}_4$ easily sol ether.

Formyl derivative $\text{C}_{11}\text{H}_{15}\text{NH(CHO)}$ [105°]. White plates, v sol alcohol and ether, nearly insol water.

Acetyl derivative $\text{C}_{11}\text{H}_{15}\text{NHAc}$ [141°]. Long silky needles.

Benzoyl derivative $\text{C}_{11}\text{H}_{15}\text{NHBz}$ [142°]. Silvery plates.

METHYL - ISOBUTYL - PHENYL - DI -

METHYL-AMINE $\text{C}_6\text{H}_5(\text{CH}_2)(\text{C}_2\text{H}_5)\text{NMe}$ [136] *Dimethyl toluisobutylamine* (250°). Colourless liquid. Formed by methylation of methyl isobutyl phenyl amine— $\text{B}^+\text{H}_2\text{Cl}_2\text{PtCl}_4$ (Effront, *B* 17, 2339).

DI METHYL-DI-BUTYL-DI PHENYL-THIO-UREA *v.* DI BUTYL DI-TOLYL THIOUREA

METHYL BUTYL PINACONE *v.* DI-OXY-DODECANE

DI METHYL DI ISOBUTYL-PYRAZINE

$\text{C}_{11}\text{H}_{20}\text{N}_2$ *ie* $\text{N} \langle \text{CMe C(C}_2\text{H}_5) \rangle \text{N}$ *Di isobutyl-ketone* (243°). Formed by reducing the oxim of methyl isobutyl diketone $\text{Me CO C(NO}_2\text{H) C}_2\text{H}_5$

with tin and HCl (Lang, *B* 18, 1364, Oeconomidis, *B* 19, 2526, Wolff, *B* 20, 433). Yellow oil— $\text{B}^+\text{H}_2\text{PtCl}_4$, orange needles.

DI METHYL ISOBUTYL PYRIDINE $\text{C}_{11}\text{H}_{17}\text{N}$ *ie* $\text{N} \langle \text{CMe CH} \rangle \text{C}_2\text{H}_5$, *Isobutyl lutidine*,

(210° - 213°) SG 8961. Formed by distilling its carboxylic acid with lime (Engelmann, *A* 231, 65). Oil, smelling of violets. Tastes bitter. More soluble in cold water than in hot water— $\text{B}^+\text{H}_2\text{PtCl}_4$, [209°]. Yellow crystalline pp— $\text{B}^+\text{H}_2\text{CrO}_4$, Yellow plates, v sol boiling water—Picrate [115°]. Yellow needles.

Hexahydrate $\text{C}_{11}\text{H}_{21}\text{N}$ *ie*

$\text{NH} \langle \text{CHMe CH} \rangle \text{CH C}_2\text{H}_5$, *s Isobutyl lupetidine*. *Di methyl isobutyl piperidine* (198°) at 720 mm. Formed by reducing di methyl isobutyl pyridine in alcoholic solution with sodium (Jaekle, *A* 246, 47). Oil. With NaNO_2 and HCl it yields a nitroso compound— B^+HCl [184°]. Prisms, v sol water and alcohol— B^+HBr . Prisms. The platinochloride and chromate could not be crystallised.

DI METHYL-ISOBUTYL PYRIDINE DI-CARBOXYLIC ACID $\text{C}_{11}\text{H}_{17}\text{NO}_4$ *ie*

$\text{N} \langle \text{CMe C(CO}_2\text{H)} \rangle \text{C}_2\text{H}_5$, [273°]. Formed by boiling its diethyl ether with alcoholic, and afterwards with aqueous, potash (Engelmann, *A* 231, 57). Long monoclinic prisms (containing 2aq). Decomposed by fusion. M sol cold, v sol hot, alcohol— CaA'' 3aq. Small four-sided pyramids, v sol water— BaA'' 5aq— $\text{H}_2\text{A}''\text{HCl}$. M sol cold water.

Mono ethyl ether HEtA'' [185°]. Formed by boiling the di ethyl ether with alcoholic potash. Long prisms, v sol water and alcohol— $\text{Ba(EtA}'')$ 3aq— $\text{HEtA}''\text{HCl}$ 2aq thick prisms.

Di ethyl ether $\text{Et}_2\text{A}''$ (312° - 318°). Formed by passing nitrous acid gas into an alcoholic solution of its dihydride. Thick oil— $\text{Et}_2\text{A}''\text{HCl}$ long needles, decomposed by water into HCl and $\text{Et}_2\text{A}''$ — $(\text{Et}_2\text{A}'')_2\text{H}_2\text{PtCl}_4$, [208°]. Cubes.

Dihydrate of the diethyl ether

$\text{NH} \langle \text{CMe C(CO}_2\text{H)} \rangle \text{CH C}_2\text{H}_5$, [100°]. Formed by warming acetoacetic ether with isovaleric aldehyde ammonia in alcohol (E). Long prisms, v sol alcohol, ether, and benzene.

METHYL ISOBUTYL SULPHATE $\text{C}_6\text{H}_5(\text{CH}_2)(\text{C}_2\text{H}_5)\text{SO}_3$ *ie* $(\text{CH}_2)(\text{C}_2\text{H}_5)\text{SO}_3$, appears to be formed by the action of methyl alcohol on $\text{C}_6\text{H}_5\text{O SO}_2\text{Cl}$, the product of the action of isobutyl alcohol on SO_2Cl (Behrend, *J pr* [2] 15, 34). Decomposed by water into HOMe and $\text{C}_6\text{H}_5\text{O SO}_2\text{OH}$.

DI METHYL ISOBUTYRYL-ACETIC ETHER *v.* DI ISOPROPYL KETONE CARBOXYLIC ETHER

METHYL CAFFEIC ACID *v.* CAFFEIC ACID

METHYL CAFFURIC ACID *v.* CAFFEINE

METHYL CARBAMATE $\text{NH}_2\text{CO}_2\text{Me}$

Methyl urethane [52°] (G), [56°] (F & K) (177°). Formed from NH_2COCl and excess of methyl alcohol (Gattermann, *A* 244, 39). Formed also from $\text{C}_6\text{H}_5(\text{N(NO}_2)_2\text{CO}_2\text{Me})$, by the action of NH_3 (Franchimont & Klobbe, *R T C* 7, 343).

METHYL CARBAMIC ACID

NHMeCO_2H . The methylammonium salt $\text{NHMeCO}_2\text{NH}_2\text{Me}$ appears to be formed by pass-

ing CO₂ into dry methylamine (Wurtz, *A Ch* [3] 80, 450, 461)

Methyl ether NHMe CO₂Me (158°) SG 15 1065 Formed by treating methyl chloroformate Cl CO₂Me with an aqueous solution of methylamine (Franchimont a Klobbie, *R T C* 7, 358) Colourless liquid, with characteristic odour With pure HNO₃ it yields a characteristic nitro-derivative

Ethyl ether NHMe CO₂Et (170°) Formed from methylamine and chloroformic ether (Schreiner, *J pr* [2] 21, 124) Oil, with pleasant odour When boiled with potash or baryta it yields alcohol, methylamine, and K₂CO₃

Chloride NHMe COCl (90°) (94°) From COCl₂ and methylamine hydrochloride (Gattermann a Schmidt, *B* 20, 118) Decomposed on boiling

Di methyl-carbamic acid *NMe₂ CO₂H

Methyl ether NMe₂ CO₂Me (131°) SG 15 1012 Formed by treating Cl CO₂OMe with an aqueous (33 p c) solution of dimethylamine (Franchimont a Klobbie, *R T C* 8, 299) Colourless liquid, smelling like menthol Pure HNO₃ forms NMe₂(NO₂) CO₂Me

Ethyl ether NMe₂ CO₂Et (140°) (Schreiner, *J pr* [2] 21, 125), (147°) (Franchimont a Klobbie, *R T C* 3, 223) SG 15 978 From Cl CO₂Et and dimethylamine With ammonia it does not form NMe₂ CO₂NH₂

Chloride NMe₂ COCl (165°) From dimethylamine and a solution of COCl₂ in benzene (Michler a Escherich, *B* 12, 1162) Liquid, with peculiar odour, v sol ether and CS₂ Decomposed slowly by water into HCl, dimethylamine, and CO₂

METHYL CARBAMINE C₂H₇N 1 e CH₃ N O

Methyl isocyanide *Methyl cyanide* *Methyl carbonylamine* *Isocetonitrile* Mol w 41 [-45°] (59.6°) VD 144 (calc 142) SG 4 756 S 10 at 15° Formed by the action of MeI (1 mol) on silver cyanide (2 mols) the product being decomposed by KOH and dried over CaCl₂ (Gautier, *C R* 63, 924, 65, 468, 862, 66, 1214, A 152, 222, *A Ch* [4] 17, 203) Formed also by the action of MeI on mercuric fulminate (Calmels, *J pr* [2] 80, 319, *C R* 99, 794)

Properties—Liquid with powerful odour, producing nausea

Reactions—1 *Aqueous acids* decompose it into methylamine and formic acid—2 HOAc forms CH₃NH CHO and Ac₂O—3 Oxidised by HgO at 50° to methyl cyanate CH₃N CO and a compound Me₂N₂C₂O₄(NH₂CHO) [175°] HgO in ethereal solution appears also to form diformamide NH(CHO)₂ or, rather, a compound Me₂N₂C₂O₄(NH(CHO))₂ [163°] (Gautier, *C R* 67, 804)—4 With AgCy it forms a compound MeNCAGCy (80°–90°) (Meyer, *J* 1856, 523) Salt—(MeNC)₂3HCl. Formed by passing HCl into an ethereal solution of methyl carbamine Crystalline Decomposed by water, partially reproducing methyl carbamine—5 MeI acts in a complicated manner (Lubavine, *Bl* [2] 45, 246, Gautier, *A Ch* [4] 17, 148)

Tri-methyl-carbamine. A name used by Rudnief (*Bl* [2] 83, 297) to denote *tert*-BUTYLAMINE

METHYL-SEMI-CARBAZIDE C₂H₇N₂O 1 e NH₂CO NH NHMe [118°]. Formed from

methyl-hydrazine acid sulphate KHO and KONO (Brüning, A 253, 10) Prismatic tables, v sol water and EtOH, m sol ether

METHYL-CARBAZOLE C₈H₁₁N 1 e

C₈H₁₁ > NMe [87°] Formed by heating potassium carbazole with MeI at 180° (Graebe, A 202, 23) Micaceous leaflets or needles, insol water, sol ether Conc H₂SO₄ containing a little HNO₃ gives a green colour—Picrate C₁₅H₁₁NC₂H₃(NO₂)₂OH [141°] Dark red needles

METHYL CARBIMIDO ALLYL THIO UREA

SC < NH > C NMe or C₂H₅ NH CS N C NMe

Allyl-thio-carbamine-methyl cyanide [110°] Formed by the action of methyl iodide upon sodium carbimido allyl thio urea Crystalline V sol water and alcohol Indifferent body (Wunderlich, B 19, 448)

METHYL-CARBIMIDO-ETHYL-THIO-UREA

SC < NH > C NMe or EtNH CS N C NMe

Ethyl-thio-carbamine-methyl cyanide [162°]

Formed by the action of methyl iodide upon sodium carbimido ethyl-thio-urea (Wunderlich, B 19, 448)

TRI-METHYL CARBINOL v *Tert* BUTYL

ALCOHOL

METHYL CARBONATES

Hydrogen methyl carbonate *MeHCO₂ The Ba salt, Ba(MeCO₂)₂, appears to be the white pp formed when CO₂ is passed into a solution of BaO in anhydrous methyl alcohol (Dumas a Pélignot, A Ch [2] 74, 6) It is insol alcohol, but dissolves in water, the solution soon depositing BaCO₃

Di-methyl carbonate C₂H₆O₂ 1 e (CH₃)₂CO₂ [0.5°] (Röse, A 205, 231) (91°) SG 2 1069 HF p 138,390 HF v 136,360 (Thomsen, Th) Formed by boiling methyl chloroformate ClCO₂Me with lead oxide (Counciler, B 13, 1697) Formed also from ClCO₂Me and NaOMe (Schreiner, J pr [2] 22, 354) Oil, insol water, sol alcohol and ether Converted by dry chlorine into hexa-chloro di methyl carbonate (q v)

METHYL-CARBOPYRIDIC ACID v METHYL-

PYRIDINE CARBOXYLIC ACID

METHYL-CARBOSTYRIL v *Methyl ether* of

CARBOSTYRIL and Oxy METHYL QUINOLINE

METHYL CETYL KETONE C₁₈H₃₈O 1 e

C₁₈H₃₈ CO OH₂ [52°] (252° at 100 mm) Formed by distilling a mixture of barium margarate and barium acetate (Krafft, B 15, 1707). Yields palmitic and acetic acids on oxidation

METHYL-CHELIDAMIC ACID v CHELIDONIC

ACID

METHYL-CHLORACETOL v. DI CHLORO-

PROPANE

METHYL CHLORIDE CH₃Cl *Chloro-*

methane Mol w 50.5 (-23.7°) (Regnault) VD 173 (calc 175) SG 2 9915, 2 9523 (Vincent a Delaachanal, A Ch [5] 16, 429) S 5.03 at 7°, 8.46 at 20° (Baeyer, A 107, 181) S. (alcohol) 35 S (HOAc) 40 HF p 22,550 HF v 21,970 (Thomsen) SV 50.8 (Lowsen, A 254, 68) Latent heat of evaporation 96,900 at 0° (Chappuis, A Ch [6] 15, 507)

Formation—1 By heating a mixture of NaCl (2 pts), wood spirit (1 pt) and H₂SO₄ (8 pts) and collecting the gas over water

(Dumas & Péligot, *A Ch* 61, 193, *A* 15, 17) — 2 By exposing a mixture of equal volumes of methane and chlorine to daylight (Berthelot, *A Ch* [3] 52, 97)

Preparation — 1 HCl is passed into MeOH (2 pts) containing ZnCl₂ (1 pt) The gas is passed through KOH and H₂SO₄ (Groves, *C J* 27, 641, *A* 174, 378) — 2 By heating trimethylamine hydrochloride at 260°, the reaction being perhaps 3NMe₃·HCl = 2NMe₃ + NH₃ + 3MeCl (Vincent, *J Ph* [4] 30, 132)

Properties — Colourless gas, with ethereal odour Burns with a bright flame, edged with green

Reactions — 1 When passed through a red-hot tube it deposits charcoal and yields HCl, methane, ethylene, CO, and naphthalene (Perrot, *A* 101, 375) — 2 When passed over heated potash-lime it yields potassium formate, KCl, and hydrogen — 3 Not attacked by chlorine in diffused daylight, but when the mixture is exposed to direct sunlight, substitution occurs, the products being methylene chloride, chloroform, and CCl₄ The chlorination may also be effected by passing a mixture of chlorine and methylchloride over animal charcoal at 250°–350° (Damoiseau, *C R* 92, 42) — 4 When equal volumes of methyl chloride and of an amine are submitted to a pressure of 25 atmospheres for 48 hours the following reactions occur — (a) Ammonia forms NMe₃·HCl, NMe₃·HCl, and 2NH₃·Cl — (b) Methylamine forms 2NMe₃·Cl and NMe₃·Cl — (c) Dimethylamine reacts with formation of NMe₃·HCl and NMe₃·Cl — (d) Trimethylamine forms NMe₃·Cl (Vincent & Chappuis, *Bl* [2] 45, 496)

Hydrate MeCl 9aq Formed by cooling below 0° water into which MeCl is passed Its vapour-tension has been studied by De Forcrand & Vilard *C R* 106, 1357, 1404)

METHYL-CHLORO-ALLYL CARBINOL *v* CHLORO PENTYL ALCOHOL

DI-METHYL-CHLORO-ANILINE *v* CHLORO DI METHYL ANILINE

TRI-METHYL CHLORO-AURO-PHOSPHITE Me₃PAuClO₂ [101°] Formed by the action of pure methyl alcohol on aurous chloride and phosphorus (Lindet, *C R* 103, 1014) Slender colourless needles, insol water, sol MeOH

METHYL CHLORO-BUTYL CARBINOL *v* CHLORO HEXYL ALCOHOL

METHYL - CHLORO - CARBOSTYRIL *v* CHLORO OXY METHYL QUINOLINE

METHYL CHLORO-ETHYL OXIDE C₂H₅ClO *v* CH₃·CHCl·OMe (72°–75°) SG 12 996 Formed from aldehyde (1 vol), methyl alcohol (1½ vols), and dry HCl at 0° (Geuther, *A* 225, 270)

METHYL TETRA-CHLORO-ETHYL OXIDE C₂H₂Cl₄O *v* CCl₃·CHCl·OCH₃ (178°) SG 2 184 Formed by the action of PCl₅ on chloral methylate (Magnanini, *G* 16, 330) Colourless liquid

METHYL CHLORO-FORMAMIDE *v* CHLORO FORMIC ACID

METHYL CHLORO - PROPYL KETONE C₃H₇ClO *v* CH₃·CO·CHClEt (180°) Formed by heating CH₃·CO·CClEt·CO₂Et with dilute HCl at 180° (Conrad, *A* 186, 241)

Methyl tri-chloro-propyl ketone C₃H₂Cl₃O *v* CH₃·CO·C₂H₂Cl₂ (193°) Formed by oxidising

C₂H₅Cl, CHMe OH with chromic acid mixture (Garzaroli Thurnlackh, *A* 223, 152) Heavy oil Does not combine with NaHSO₄ Yields CO₂, acetic acid, and HCl on oxidation

DI METHYL-CHLORO - QUINALDINE *v*

CHLORO TRI METHYL QUINOLINE

METHYL-TRI CHLORO-QUINOLINE *v* TRI-CHLORO METHYL-QUINOLINE

METHYL DI - CHLORO - VINYL OXIDE C₂H₂Cl₂O *v* CCl·CH·OCH₃ (110°) SG 2 12934, 122 11574 Prepared by heating tri-chloro ethylene with KOMe, being ppt on addition of water (Denaro, *G* 14, 117) Colourless oil, with an odour resembling that of acetal Decomposed on exposure to air, with evolution of HCl When heated with dilute H₂SO₄ it yields di chloro acetic orthoaldehyde

Methyl tri chloro-vinyl oxide CCl₂·CCl·OCH₃ Readily absorbs dry oxygen, forming the compound CCl₂·CCl·O(OMe), which, by the action of water, gives oxalic acid (Henry, *B* 12, 1838)

METHYL CHRYSOIDINE *v* Benzene-Azo methyl phenylene diamine

METHYL-CINCHENIC ACID *v* CINCHENE

METHYL - CINCHONAMINE *v* CINCHONA

BASES

DI-METHYL-CINCHONIC ACID *v* DI

METHYL QUINOLINE CARBOXYLIC ACID

METHYL-CINCHONIDINE *v* CINCHONIDINE

METHYL-CINCHONINE *v* CINCHONINE

METHYL CINNAMIC ACIDS *v* PHENYL-

METHACRYLIC ACID AND TOLYL ACRYLIC ACID

METHYL - CINNAMIC ALDEHYDE *v*

PHENYL METHACRYLIC ALDEHYDE

METHYL CINNAMYL KETONE *v* BENZYL-

IDENE ACETONE

METHYL CINNOLINE CARBOXYLIC ACID

C₁₀H₇N₂O₂ *v* C₆H₅·(CO·H) $\begin{matrix} \text{CMe CH} \\ \diagdown \\ \text{N}=\text{N} \end{matrix}$ [c 230°]

Small tables or fine needles Sol acetic acid, sl sol hot alcohol and hot water, insol cold water Dissolves both in aqueous acids and alkalis, forming easily soluble salts Obtained, as a yellow pp, by adding NaNO₂ to a cold solution of the hydrochloride of amido propenyl benzoic acid C₆H₄·(CMe CH)·(NH₂)·CO₂H [2 4], very probably di azo-propenyl-benzoic acid C₆H₄·(CO₂H) $\begin{matrix} \text{CMe CH} \\ \diagdown \\ \text{N}=\text{N OH} \end{matrix}$ is first formed, which then splits off H₂O (Widman, *B* 17, 722)

METHYL-CITRACONIC ACID C₇H₈O₄ [100°]

A product of the dry distillation of methyl-paraconic acid (Frankel *A* 255, 25) Long needles (from CHCl₃) Reduced by sodium amalgam to ethyl succinic acid — CaA''aq — BaA''4aq — Ag A'

METHYL COCAINE C₁₇H₂₁NO₄ [47°] Extracted from synthetic cocaine (cf vol 1 p 230), in which it occurs in very small quantity (Liebermann & Giesel, *B* 23, 508, 926) An oil, which crystallises after standing some hours Ppd by Na₂CO₃ from its salts Very sol ether, chloroform, benzene, and ligroin It can be separated from ordinary cocaine by means of its nitrate A 5 4 pc solution of the hydrochloride in a 2 dm tube gave a rotation of +4 5° By heating with water it is not so easily decomposed as ordinary cocaine By boiling with hydriodic acid MeI is split off By hydrochloric acid it is split

up finally into methyl alcohol, benzoic acid, and methyl ecgonine hydrochloride. This reaction is also more difficultly brought about than with ordinary cocaine. Methyl cocaine appears to be identical with 'dextro cocaine,' a body got by heating the methyl ether of dextro ecgonine with BzCl (Einhorn & Marquardt, *B* 23, 468).

Salts— B^+HCl [210°] needles or columns forming rosettes (from hot alcohol), much less soluble than the hydrochloride of ordinary cocaine—Sulphate plates, v sol water, sl sol alcohol—Nitrate crystals, sl sol water—Platino-chloride $(\text{C}_{15}\text{H}_{23}\text{NO}_2\text{HCl})_2\text{PtCl}_6$ glittering yellowish needles (from hot water)—Auro-chloride $\text{C}_{15}\text{H}_{23}\text{NO}_2\text{HClAuCl}_4$ [148°] needles, melts under water to a yellow oil.

METHYL CODEINE $\text{C}_{15}\text{H}_{21}\text{NO}_2$, $\pm e$
 $\text{C}_{15}\text{H}_{21}\text{MeNO}$, or $\text{C}_{15}\text{H}_{21}\text{Me}_2\text{NO}_2$ [118.5°]
 $[\alpha]_D^{20} = -209^\circ$ in a 4 c solution (in 97 p c alcohol) at 15°. Formed by boiling codeine methyl iodide $\text{C}_{15}\text{H}_{21}\text{NO}_2\text{MeI}$ with potash, and extracting with benzene (Grimaux, *A Ch* [5] 27, 276, Hesse, *A* 222, 218). Anhydrous prisms (from alcohol or ether), or hydrated crystals (containing aq) (from water). Insol water, v e sol hot alcohol, m sol ether. Its solution in conc H_2SO_4 is violet, but becomes blue on heating. Boiling Ac_2O forms a compound $\text{C}_{15}\text{H}_{21}\text{O}_3$ [131°]. Its hydrochloride forms a brown solution in H_2SO_4 , which becomes violet on warming, and finally blue on exposure to the air— B^+HCl 2aq needles S 9 at 18°. V e sol alcohol, sl sol NaCl aq— $\text{B}^+\text{H}_2\text{PtCl}_6$ aq.

Acetyl derivative $\text{C}_{17}\text{H}_{23}\text{AcMe}_2\text{NO}_2$ [66°] Formed by heating methyl codeine with Ac_2O at 85°, adding aqueous NH_3 , and extracting with ether (Hesse, *A* 222, 222). Glittering tables (from ether), v sol alcohol and ether, sl sol water, insol KOH aq. Alcoholic potash forms potassium acetate and methyl codeine— $(\text{C}_{17}\text{H}_{23}\text{AcMe}_2\text{NO}_2)\text{HCl}$ 2aq satiny plates, sl sol cold water— $(\text{C}_{17}\text{H}_{23}\text{AcMe}_2\text{NO}_2)_2\text{H}_2\text{PtCl}_6$ 4aq yellow laminae— $(\text{C}_{17}\text{H}_{23}\text{AcMe}_2\text{NO}_2)\text{HNO}_3$ 3aq satiny plates— $(\text{C}_{17}\text{H}_{23}\text{AcMe}_2\text{NO}_2)_2\text{H}_2\text{SO}_4$ 8aq.

(a) **Methyl iodide** $\text{C}_{15}\text{H}_{21}\text{NO}_2\text{MeI}$ 3aq prisms, formed at once on dissolving methyl codeine in a mixture of MeI and methyl alcohol (Hesse).

(b) **Methyl iodide** $\text{C}_{15}\text{H}_{23}\text{NO}_2\text{MeI}$ Slender crystals, formed by boiling the preceding with aqueous NaOH , and ppg with KI . It is less soluble in water than its (a) isomeride. Ag_2SO_4 forms $(\text{C}_{15}\text{H}_{23}\text{NO}_2\text{Me})_2\text{SO}_4$, crystallising in plates.

(a) **Methyl chloride** $\text{C}_{15}\text{H}_{23}\text{NO}_2\text{MeCl}$ From the (a)-iodide and AgCl . Amorphous. With platonic chloride it yields a yellow flocculent pp of $(\text{C}_{15}\text{H}_{23}\text{NO}_2\text{MeCl})_2\text{PtCl}_6$ 8aq. With Ac_2O it yields $\text{C}_{15}\text{H}_{23}\text{AcNO}_2\text{MeCl}$ 2aq, which crystallises in long satiny needles, v e sol alcohol, sl sol cold water. Its platinochloride $(\text{C}_{15}\text{H}_{23}\text{AcNO}_2\text{MeCl})_2\text{PtCl}_6$ 4aq is a yellow crystalline pp.

(b) **Methyl chloride** $\text{C}_{15}\text{H}_{23}\text{NO}_2\text{MeCl}$ 3aq Radiating crystalline mass, v sol water and alcohol. With conc H_2SO_4 it gives a purple colour. With platonic chloride it yields an orange pp of small needles of the platinochloride $(\text{C}_{15}\text{H}_{23}\text{NO}_2\text{MeCl})_2\text{PtCl}_6$ aq. With Ac_2O it yields amorphous $\text{C}_{15}\text{H}_{23}\text{AcNO}_2\text{MeCl}$, which forms $(\text{C}_{15}\text{H}_{23}\text{AcNO}_2\text{MeCl})_2\text{PtCl}_6$ 8aq, a yellow powder.

Methyl hydroxide $\text{C}_{15}\text{H}_{23}\text{NO}_2\text{MeOH}$. Formed from the (b) methyl sulphate and baryta. Colourless plates, insol ether, v sol alcohol. Strongly alkaline in reaction. Conc H_2SO_4 forms a blue solution. Gives off tri-methylamine when strongly heated.

METHYL COLCHICINE $\text{C}_{25}\text{H}_{35}\text{NO}_5$, $\pm e$
 $\text{C}_{15}\text{H}_2(\text{OMe})_2(\text{NAcMe})\text{CO}_2\text{Me}$. Occurs in the mother liquor from which the compound of colchicine with chloroform has separated (vol u p 234). Decomposed by boiling dilute HCl into methyl chloride and methyl colchicine $\text{C}_{15}\text{H}_2(\text{OMe})_2(\text{NAcMe})\text{CO}_2\text{H}$ (Johanny & Zeisel, *M* 9, 871).

DI METHYL-COLCHICINIC ACID v *D*.
 methyl derivative of COLCHICINIC ACID.

Tri methyl-colchicinic acid $\text{C}_{15}\text{H}_2\text{NO}_5$, $\pm e$
 $\text{C}_{15}\text{H}_2(\text{OMe})_3(\text{NH}_2)\text{CO}_2\text{H}$ [150°] Formed by the action of HCl on colchicine (*q* v). Forms a platinochloride $\text{B}^+\text{H}_2\text{PtCl}_6$ 2aq. On warming with acetic anhydride it yields colchicine $\text{C}_{15}\text{H}_2(\text{OMe})_2(\text{NHAc})\text{CO}_2\text{H}$. With MeOH it forms an addition product $\text{C}_{15}\text{H}_2\text{NO}_5\text{MeOH}$. On warming with MeOH , methyl iodide, and sodium it yields 'tri methyl colchidimethinic acid' $\text{C}_{15}\text{H}_2(\text{OMe})_3(\text{NMe}_2)\text{CO}_2\text{H}$ [125°]. The methyl ether of this body forms an iodomethylate $\text{C}_{25}\text{H}_{35}\text{NO}_5\text{Iaq}$ (Johanny & Zeisel, *M* 9, 877).

METHYL-CONINE v CONINE.

METHYL-COUMARIC ACID v *Methyl derivative of COUMARIC ACID*.

Di-methyl di coumaric acid so called v *Di METHYL DI-COUMARIN*.

METHYL COUMARILIC ACID $\text{C}_{10}\text{H}_8\text{O}_3$, $\pm e$
 $\text{C}_6\text{H}_5\text{C}(\text{OMe})_2\text{CO}_2\text{H}$ [126°] Formed by the action of dilute KOH upon the methyl derivative of *exo* bromo coumaric acid (Perkin, *C J* 39, 423). Needles (from CS_2).

(b) **Methyl coumarilic acid** $\text{C}_{10}\text{H}_8\text{O}_3$, $\pm e$
 $\text{C}_6\text{H}_5\text{C}(\text{OMe})_2\text{CO}_2\text{H}$ [189°] Formed by saponifying its ethyl ether with alcoholic potash (Hantzsch, *B* 19, 1290). Feathery needles (from dilute alcohol). Decomposed by heat into CO_2 and (b) methyl coumarone— KA aq needles— NH_4A aq needles (from water)— BaA , 3aq— AgA minute prisms.

Ethylether EtA [51°] (290°) Formed by the action of sodium phenylate NaOC_6H_5 upon chloro aceto acetic ether, the resulting phenoxy aceto acetic ether being condensed by cold conc H_2SO_4 (Hantzsch).

Amide $\text{C}_6\text{H}_5(\text{CH}_2)_2\text{O CO NH}_2$; [145°], needles (Hantzsch, *B* 19, 2401).

Di-methyl-coumarilic acid $\text{C}_{11}\text{H}_{10}\text{O}_3$, $\pm e$

$\text{C}_6\text{H}_5(\text{CH}_2)_2\text{C}(\text{CH}_3)\text{CO}_2\text{H}$ *Di methyl coumarone-a carboxylic acid* [225°]

Formation—1 By the action of hot alcoholic KOH upon bromo di methyl cumarin

$\text{C}_6\text{H}_5(\text{CH}_2)_2\text{C}(\text{CH}_3)\text{CBr}$
 $\text{O} \quad \text{O} \quad \text{CO}$
 —2 By saponification

of the ethyl ether obtained by the reaction of sodium *p* cresol and chloro aceto acetic ether.

Properties—Short prisms or tables. On heating the sodium salt with lime di methyl coumarone is obtained.

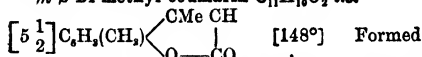
Ethyl ether AEt [55°], (300° at 728 mm) (Hantzsch & Lang, *B* 19, 1299).

Reference — COUMARILIC ACID and OXY-

METHYL COUMARILIC ACID

METHYL COUMARIN *v* Anhydride of OXY-
PHENYL CROTONIC ACID

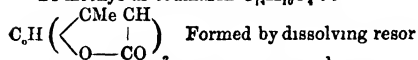
m β Di methyl coumarin $C_{11}H_{10}O_2$, *z* a.



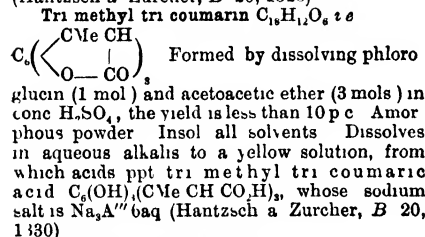
by the action of H_2SO_4 on a mixture of acetoacetic ether and *p* cresol (Pechmann & Duisberg, *B* 16, 2127, Pechmann & Cohen, *B* 17, 2187) Long colourless needles

Reference — Brom. di methyl coumarin

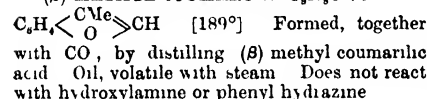
Di methyl di coumarin $C_{14}H_{12}O_4$, *z* e



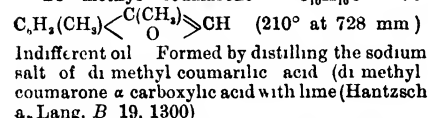
Tri methyl tri coumarin $C_{18}H_{14}O_6$, *z* e



(β) **METHYL COUMARONE** C_9H_8O , *z* e



Di methyl coumarone $C_{10}H_{10}O$, *z* e



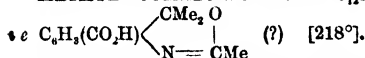
METHYL-CREOSOL *v* Methyl ether of CREOSOL

METHYL CROTONIC ACIDS *v* ANGELIC ACID and TIGLIC ACID

METHYL ISOCROTYL OXIDE *v* Sec ISO BUTENYL ALCOHOL

METHYL CUMARIN *v* METHYL COUMARIN

METHYL-CUMAZONIC ACID $C_{12}H_{11}NO_4$, *z* e



Formation — 1 By boiling (341) amido-oxypropyl benzoic acid with acetic anhydride 2 By boiling acetyl amido oxypropyl benzoic acid with HCl — 3 By boiling acetyl amido propenyl benzoic acid with HCl Small colourless trimetric tables V sol alcohol, insol water Its N is tertiary

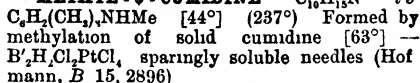
Reactions — By reduction with sodium-

amalgam it yields the acetyl derivative of amido cuminic acid

Salts — $A'H.HCl^x$ very soluble white needles — $(A'H.HCl)_2PtCl_6$ tables, cubes, or prisms — $A'H.H_2SO_4$ aq very soluble white silky needles (Widmann, *B* 16, 2576)

METHYL-CUMENE *v* CYMENE

METHYL-ψ-CUMIDINE $C_{10}H_{11}N$, *z* e



Di methyl-ψ cumidine $C_6H_5(CH_3).NMe_2$ (222°) Oil Formed by methylation of solid cumidine [63°] — $B'_2H_2Cl_2PtCl_6$ (Hofmann, *B* 15, 2897)

Methylo-iodide $C_6H(CH_3).NMe_2.I$

Prisms Yields $(C_6H_5(CH_3).NMe.Cl)_2PtCl_6$

METHYL CUMYL ETHER KETONE

$C_6H_5.C_6H_5.CH_2.CH_2.CO.CH_3$ Cumyl acetone (260°–265°) Is one of the products of the action of NaOEt and cumyl chloride on acetoacetic ether (Widmann, *B* 22, 2271) Colourless liquid, with fragrant odour Is not acted upon by NaOBr Oxidised by $KMnO_4$ to cuminic acid

Oxim $C_6H_5.C_6H_5.CH_2.CH_2.C(OH)CH_3$ [57°] Long shining prisms (from ligroin)

METHYL CUMYL KETONE

$CH_3.CO.C_6H_5.Pr$ [14] (253°) SG $d_{20}^{25} 9755$ Formed by the action of $AcCl$ on cymene in presence of $AlCl_3$ (Widmann, *B* 21, 2225)

Oxim $CH_3.C(OH)C_6H_5.Pr$ [71°] Four-sided tables (from petroleum ether)

Phenyl hydrazide

$CH_3.C(NHPh)C_6H_5.Pr$ [82°] Colourless thick six sided tables (from petroleum ether)

METHYL CYANAMIDE *v* Cyanamide in the article CYANIC ACIDS

METHYL CYANATE *v* CYANIC ACIDS

METHYL-CYANETHINE *v* CYANETHINE

METHYL CARBIMIDE is Methyl isocyanate *v* CYANIC ACIDS

METHYL CYANIC ACID *v* CYANIC ACID

METHYL CYANIDES *v* ACPIONITRILE and

METHYL CARBAMINE

Di methyl di cyanide C_6H_5N , *z* e

$NH.CMe.CH_2.CN$ or $NH_2.CMe.CH.CN$ [53°] VD 42.5 Formed by the action of dry sodium on acetonitrile dissolved in ether Methane is evolved in the reaction, and the product is decomposed by water (Holtzwarth, *J pr* [2] 38, 343, 39, 240) White needles, v sol ether, alcohol, chloroform, and benzene, m sol water, sl sol petroleum ether

Reactions — 1 Boiling water liberates ammonia forming $C_6H_5N_2O$, a body which is converted by PCl_5 into crystalline $C_6H_5N_2Cl_2$ [175°], which, on recrystallisation from water, becomes $C_6H_5N_2$ — 2 Acetyl chloride forms a compound $(C_6H_5N_2).CH_3.COCl$, which on decomposition by water yields $C_6H_5N_2$, crystallising in beautiful white needles [223°] — 3 Warm dilute (25° c) HCl aq forms NH_4Cl and an oil which has the composition of cyano acetone It solidifies to a glassy mass, carbonises above 230°, and forms with phenyl hydrazine a condensation product [97°]

Tri-methyl tri-cyanide *v* CYANMETHINE

METHYL-CYANO FORMAMIDE *v* Methyl-amide of Para-Cyanopformic ACID

METHYL-CYANO-SUCCINIC ETHER

$C_7H_{11}NO$, \pm CO_2Et CH_2 $CMcCy$ CO_2Et An oil formed by treating cyano succinic ether with Na and MeI successively (Barthe, *C R* 108, 297)

\pm -Di-methyl cyano-succinic ether $C_{11}H_{11}NO$, \pm CO_2Et $CHMe$ $CMcCy$ CO_2Et (273°) S G. 24.5 1.0577 A product of the action of alcoholic KCy on a bromo propionic ether (Zelinsky, *B* 21, 3164) Formed also by adding a bromo-propionic ether to cyano-propionic ether mixed with KCy (Z)

METHYL CYANURIC ACID *v* *Cyanuric acid* in the article *CYANIC ACIDS*

METHYL CYMYL KETONE $C_{15}H_{18}O$ \pm CH_3 CO C_6H_5 Me C_6H_5 [2 14] (247°) An oil, formed by the action of AcCl on cymene in presence of $AlCl_3$ (Claus, *B* 19, 233)

METHYL-DAMBOSE *v* *BORNESITE*, vol 1, p 524

Di-methyl-dambos *v* *DAMBONITE*

METHYL-DAPHNETIN *v* *DAPHNETIN*

METHYL DECYL KETONE $C_{12}H_{22}O$ \pm CH_3 CO $C_{10}H_{21}$ [21°] (247°) Formed by distilling a mixture of barium acetate and barium hendecate (undecylate) (Krafft, *B* 15, 1708) Yields acetic and decic acids on oxidation

METHYL-DESOXYBENZOLIN *v* *TOLYL BENZYL KETONE*

Dimethyl-desoxybenzol *v* *BENZYL XYLIL KETONE*

METHYL DODECYL KETONE $C_{14}H_{28}O$ \pm CH_3 CO $C_{12}H_{25}$ [34°] (206° at 100 mm) Formed by distilling barium tridecoteate ($C_{13}H_{26}O_2$) Ba with barium acetate (Krafft, *B* 15, 1708) Yields lauric and acetic acids on oxidation

METHYL c-DURYL KETONE $C_{12}H_{18}O$ \pm CH_3 CO C_6H_5 Me [1 2 3 4 5] (259°) From c-durene, AcCl, and $AlCl_3$ (Claus & Fohlisch, *J pr* [2] 38, 230) Oil

Phenyl hydrazide [129°] Lamine

Isomerides *v* *DURYL METHYL KETONES*

METHYL-EGGONINE $C_{15}H_{17}NO$ [264°] A product of the action of HCl on methyl cocaine, the resulting methyl eggonine hydrochloride being decomposed by Ag_2O (Liebermann & Giesel, *B* 23, 510) May be crystallised from methyl alcohol containing a trace of water. Extremely sol water, insol absolute alcohol, v sl sol absolute MeOH. Decomposed by fusion. A 96 p c solution of the hydrochloride exhibits $\alpha = +2^\circ$ —B'HCl [236°], needles (from methyl alcohol ether) —B'HAuCl₄ [220°], lemon yellow needles

Benzoyl derivative Hydrochloride $C_{17}H_{19}NO_2$ HCl The first product of the action of hydrochloric acid on methyl cocaine [47°], the base being just dissolved in HCl and then heated $\frac{1}{2}$ hour at 90° Glassy columns (from hot water) The base is not ppt by carbonate of soda —Aurochloride $C_{17}H_{19}NO$ HCl AuCl₄ The nitrate is v sl sol water

METHYLENE The radicle CH_2 , which is not known to exist in the free state

Dimethylene C_2H_4 , \pm CH_2 CH_2 is called **ETHYLENE** (*q v*)

Trimethylene C_3H_6 , \pm CH_2 $\begin{matrix} \diagup CH_2 \\ | \\ CH_2 \end{matrix}$ H F p

—8470 H.F.v. —4630 This gas is formed

when trimethylene bromide CH_2Br C_2H_4 CH_2Br (140 g) is boiled with sodium (5 g) The contents of the flask become pasty, but still contain much of the bromide (120 g), which can be mostly recovered by filtration (Freund, *J pr* [2] 26, 367) It is also formed by heating tri-methylene bromide with zinc dust and 75 p c alcohol (Gustavson, *J pr* [2] 36, 300) Trimethylene burns with a bright flame and smells like butylene

Reactions —1 Trimethylene is readily absorbed by fuming HI The product is *n* propyl iodide, whereas propylene gives *iso* propyl iodide.

2 Trimethylene is very slowly absorbed by bromine, the product is trimethylene bromide (165°) Propylene is readily absorbed, forming propylene bromide —3 Conc H_2SO_4 forms liquid hydrocarbons, and on diluting and distilling *n* propyl alcohol is got (G) —4 $KMnO_4$ does not oxidise it (Wagner, *B* 21, 1230)

References —TRIMETHYLENE CARBOXYLIC ACIDS, TRIMETHYLENYL METHYL KETONE, and PHENYL TRIMETHYLENYL KETONE

Tri-methylene is a term also applied to the divalent radicle CH_2 CH_2 CH_2

Tetra-methylene is a term applied to the ring CH_2 CH_2

$\begin{matrix} | \\ CH_2 \\ | \end{matrix}$, and it is also used to denote the group CH_2 CH_2 CH_2 CH_2 CH_2 CH_2

Pentamethylene is a name used to denote CH_2 CH_2 $\begin{matrix} \diagup CH_2 \\ | \\ CH_2 \end{matrix}$ $\begin{matrix} \diagup CH_2 \\ | \\ CH_2 \end{matrix}$ CH_2 , and also CH_2 CH_2 CH_2 CH_2 CH_2 CH_2

Hexamethylene is benzene hexahydride According to Baeyer (*A* 258, 156), it should be regarded as lying entirely in one plane

METHYLENE ACETATE *v* *Acetyl derivative of FORMIC ORTHALDEHYDE*, vol 11 p 570

TRIMETHYLENE ACETO ACETIC ACID, so called, *v* vol 1 p 24

METHYLENE DIAMINE *Di benzoyl derivative* $CH(NHBz)_2$ [221°] S (alcohol) 47 at 14.5°, 63 at 22°, *Apparaffin*

Formation —From hippuric acid, PbO_2 and HNO_3 or H_2SO_4 (H Schwarz, *A* 75, 201, *Sit. W* 77, 11 762, J Maier, *A* 127, 162, Kautz *A* Y Schwarz, *A* 223, 40)

Preparation —From benzonitrile (15 g), methylal (6 g) and conc H_2SO_4 (100 g) (Hepp & Spiess, *B* 9, 1424)

Properties —Long white felted needles (from alcohol), v sol CS_2 , ether, and chloroform

Reactions —1 Dissolves unchanged in conc H_2SO_4 , and in fuming HNO_3 —2 On distillation some passes over unchanged along with benzoic acid —3 Not affected by bromine —4 Boiled for some time with dilute (32 p c) H_2SO_4 it gives formic aldehyde, NH_3 , and benzoic acid —5 Heated in sealed tubes with dilute HCl it yields benzamide and formic paraldehyde

Trimethylene diamine $C_3H_7N_2$, \pm NH_2 CH_2 CH_2 CH_2 NH_2 (c 140°) Formed by heating trimethylene bromide (1 mol) and NH_3 (20 mols) in alcohol for 10 hours at 100° The liquid is decanted from NH_4Br and evaporated, treated with KOH and distilled (Fischer & Koch, *B* 17, 1799, Lellmann & Wurtthner, *A* 228, 227) Colourless mobile liquid, easily miscible with alcohol, ether, and benzene Fumes in moist air, combining with water to form a

hydrate Readily takes up CO_2 becoming solid
Condenses with benzoic aldehyde forming
 $\text{C}_6\text{H}_5(\text{N OHPh})_2$ Phenanthraquinone forms
 $\text{C}_{12}\text{H}_8\text{N}_2\text{O}_2$ a yellow powder, melting above 250°

Benzil forms $\text{CH}_2\text{<CH}_2\text{N CPh>CH}_2\text{N CPh>}$ a transparent
vitreous mass [α 76°]

Salts — $\text{B}^+\text{H}_2\text{Cl}_2$ Easily soluble prisms —
 $\text{B}^+\text{H}_2\text{Cl}_2\text{PcCl}_2$ Orange prisms — $\text{B}^+(\text{HSCN})_2$
[102°] At 140° it is partially decomposed into
trimethylene thio-urea and NH_4SCN —
 $\text{B}^+\text{H}_2\text{Br}_2$

Diacetyl derivative $\text{C}_6\text{H}_5(\text{NHAc})_2$ [79°]
White needles, v sol water, v sol alcohol,
chloroform, sl sol benzene, insol ether, petro-
leum ether (Strache, *B* 21, 2364)

Dibenzoyl derivative $\text{C}_6\text{H}_5(\text{NHBz})_2$
[148°] White crystalline powder, insol water,
m sol benzene, v sol alcohol and chloroform
On heating in a stream of HCl the product is
 $\text{C}_6\text{H}_5\text{<NH>OPh}$, an oily base which slowly
becomes crystalline and forms crystalline salts
(Hofmann, *B* 21, 2337)

Oxalyl derivative $\text{CH}_2\text{<CH}_2\text{NH CO>CH}_2\text{NH CO>}$

A sparingly soluble white powder, formed on
mixing trimethylene diamine with an alcoholic
solution of methyl oxalate It does not melt at
 250°

Derivative — $\text{V OXY TRI METHYLENE DIAMINE}$

Trimethylene di nitro di amine $\text{C}_3\text{H}_8\text{N}_4\text{O}_2$ i.e.
 $\text{NO}_2\text{NHCH}_2\text{CH}_2\text{CH}_2\text{NHNO}_2$ [67°] Formed by
heating $\text{C}_3\text{H}_8(\text{N(NO)}_2\text{CO}_2\text{Me})_2$ (v TRI METHYLENE
DICARBAMIC AMIDE) with aqueous ammonia (Franchi-
mont a Klobbie, *R T C* 7, 343) Short thick
prisms (from water or alcohol), v sol water and
alcohol, less sol ether and chloroform Boiled
with dilute (2 p c) H_2SO_4 it evolves N_2O
It has no acid reaction, and easily forms metallic
derivatives

Tetra methylene diamine $\text{C}_4\text{H}_{12}\text{N}_4$ i.e.
 $\text{NH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{NH}_2$ [24°] (159°) Ob-
tained by reducing ethylene cyanide in alcoholic
solution with sodium (Ladenburg, *B* 19, 780)
Identical with the ptomaine 'putrescine' (cf L
Brieger, 'Die Ptomaine,' Berlin, 1885-1886, 1,
43, 3, 101) and also with a base obtained from
a morbid urine (Udranszky a Baumann, *B* 21,
2938) Strong base, smelling like piperidine
Solutions of its salts give a crystalline pp with
iodine dissolved in KIaq but no pp with HgCl_2
or KODl_2 When shaken with aqueous NaOH
and BzCl it gives a crystalline pp of the di-
benzoyl derivative Methyl chloroformate
 ClCO_2Me forms $(\text{CH}_2)_4(\text{NH CO}_2\text{Me})_2$, which crys-
tallises in flattened needles [128°], sl sol cold
water, and is nitrated by HNO_3 , yielding
 $(\text{CH}_2)_4(\text{N(NO}_2)_2\text{CO}_2\text{Me})_2$, which crystallises from
ether and melts at 62° (Dekkers, *R T C* 9, 97)

Salts — $\text{B}^+\text{H}_2\text{Cl}_2$ Plates (from alcohol) —
 $\text{B}^+\text{H}_2\text{PcCl}_2$ Needles — Picrate Thin yellow
needles — Aurochloride [210°] Minute
thread like needles, v sol hot water (Giamicani
a Zanetti, *B* 22, 1978)

Di-benzoyl derivative $\text{O}_2\text{H}_5(\text{NHBz})_2$
[175°] (U a B), [178°] (O a Z) Plates, in-
sol water, almost insol ether, v sol hot alcohol
May be sublimed

Tetra-methylene-di-nitro-di-amine $\text{C}_4\text{H}_{12}\text{N}_4\text{O}_2$
i.e. $(\text{CH}_2)_4(\text{NH NO}_2)_2$ [168°] Got by heating
Vol. III.

$(\text{CH}_2)_4(\text{N(NO}_2)_2\text{CO}_2\text{Me})_2$ with conc NH_4Aq at
 100° and ppg by HOAc (D) Small hard crys-
tals (from water)

Penta methylene diamine $\text{C}_5\text{H}_{12}\text{N}_4$ i.e.
 $\text{NH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{NH}_2$ (178.5°) (L),
[178° - 180.5°] (Perkin, *C J* 55, 699) SG $\frac{1}{2}$
9174 (L), $\frac{1}{2}$ 8846, $\frac{3}{2}$ 8784 M M 7493 (P)
Formed from trimethylene cyanide by reduction
in ethereal solution with zinc and HCl , or in
alcoholic solution with sodium (Ladenburg, *B*
16, 1151, 18, 2956, 19, 780, 2585) It is iden-
tical with 'cadaverine' a base discovered by
Drieger in corpses, and among the products of
putrefaction of flesh and fish (Brieger, *B* 16,
1186, 18, 1922, 'Die Ptomaine,' Berlin, 1885,
Ladenburg, *B* 19, 2585) Found also in the
urine of a patient suffering from cystinuria, but
not in normal urine (U a B) Syrup, smelling
like piperidine, v sol water and alcohol, m sol
ether Fumes in the air Absorbs CO_2 from
the air The hydrochloride is converted by dry
distillation into NH_3 , HCl , and piperidine

Salts — $\text{B}^+\text{H}_2\text{Cl}_2$ — $\text{B}^+\text{H}_2\text{PcCl}_2$ Thick orange
prisms (from water), m sol cold water — Per-
iodide Almost black crystals (from alcohol) —
 $\text{B}^+\text{H}_2\text{Cl}_2$, 3 HgCl_2 Crystals (from hot water)
(L) — $\text{B}^+\text{H}_2\text{Cl}_2$, 4 HgCl_2 Crystals (from alcohol)
(B)

Di acetyl derivative

$\text{CH}_3(\text{CH}_2)_3\text{CH}_2\text{NHAc})_2$ Small needles (from al-
cohol) May be distilled

Di benzoyl derivative

$\text{CH}_3(\text{CH}_2)_3\text{CH}_2\text{NHBz})_2$ [130°] (above 360°) Ppd
by adding aqueous NaOH and BzCl to a solution
of the base (Udranszky a Baumann, *B* 21,
2744) Long needles and plates, v sol alcohol,
m sol ether, insol water Not affected by hot
dilute acids or alkalis

Penta methylene di nitro di amine

$\text{CH}_3(\text{CH}_2)_3\text{CH}_2\text{NH(NO}_2)_2$ Formed by treating
 $\text{CH}_3(\text{CH}_2)_3\text{CH}_2\text{N(NO)}_2\text{CO}_2\text{Me}$ with aqueous
ammonia (Franchimont a Klobbie, *R T C* 7,
343) Small oblong plates (from chloroform),
v sol water and alcohol, sol ether, sl sol
 CHCl_3 When boiled with dilute (2 p c) H_2SO_4
it evolves N_2O

Di nitroso penta methylene tetramine (so
called) $\text{C}_5\text{H}_{12}\text{N}_4\text{O}_2$ i.e. $\text{C}_5\text{H}_{12}\text{N}_4(\text{NO})_2$ [207°] (G),
[203°] (M) Formed by the action of nitrous
acid on 'hexamethylene tetramine' (Griess, *B*
21, 2738, Mayer, *B* 21, 2888) Needles (from
alcohol), v sol hot alcohol, m sol chloroform,
insol ether Not affected by boiling with zinc-
dust Dilute HCl decomposes it into nitrogen,
ammonia, and formic aldehyde

Hexa-methylene tetramine (so called)

$\text{C}_6\text{H}_{12}\text{N}_4$ Hexamethylenamine [189°] 8
(alcohol) 7 Mol w 115 (by Raoult's method)
(calc 140) (Tollens a Mayer, *B* 21, 1566)
Formed by passing dry NH_3 over heated tri-
oxy methylene (formic paraldehyde) (Butlerow,
A 115, 322, *Z* [2] 5, 278) Prepared by dis-
solving methylal in dilute H_2SO_4 , and distilling
the product with steam into a receiver containing
ammonia (Wohl, *B* 19, 1842)

Properties — Rhombohedra (from alcohol),
v sol water, chloroform, and CS_2 May be sub-
limed Scarcely attacked by sodium-amalgam
Acts nutritiously upon algae (Loew a Bokorny,
J. pr [2] 86, 272)

Reactions. — 1. Split up by boiling dilute

acids into formic aldehyde and ammonia—2 MeI at 190° forms NMeI (Tollens, *B* 17, 656)
 3 Sodium nitrite (2½ pts) added to a solution of the base (1 pt) in dilute HCl at 0° forms 'tri methylene tri nitrosamine' $C_3H_8N_3O_3$, which crystallises from alcohol in needles or prisms [106°], and is split up by warming with dilute acids into nitrogen and formic aldehyde. It exhibits Liebermann's reaction. When acetic acid is added gradually to a solution containing hexamethylene tetramine and sodium nitrite there is formed the so called di nitroso penta methylene tetramine (*v supra*), which is much less soluble in the usual menstrua than the tri methylene tri nitrosamine.

Salts— $B^+H_2Cl_2$ (dried at 100°). Long prismatic needles, *v* sol water, sl sol alcohol— $B^+H_2PtCl_4$ 4aq— B^+3AgNO_3 . White crystalline pp, formed on adding $AgNO_3$ to an aqueous solution of hexamethylene tetramine (Pratesi, *G* 13, 437). Sl sol cold, m sol hot, water, with partial reduction to silver.

Methylo iodide B^+MeI [190°]. *V* sol water, sl sol alcohol, insol ether, chloroform, and CS_2 . Gives $(B^+Me)PtCl_6$ [205°].

Ethyl iodide B^+EtI [133°].

Methylene iodide $B^+CH_2I_2$ [165°] (Wohl).

Di-bromide $C_3H_8N_2Br_2$. Insol ordinary solvents (Legler, *C C* 1888, 1604).

Tetrabromide $C_3H_8N_2Br_4$. Red crystals, obtained by the action of bromine vapour on hexamethylene tetramine (Horton, *B* 21, 1999). Decomposed by boiling water.

Di iodide $C_3H_8N_2I_2$. Obtained by adding an alcoholic solution of iodine to an aqueous solution of the base. Crystalline, *v* sl sol alcohol.

Tetra iodide $C_3H_8N_2I_4$. Obtained when excess of iodine is used. Minute brown plates, *v* sol acetone, $CHCl_3$, and CS_2 . Decomposed by boiling water.

METHYLENE DI-ISOAMYL DIOXIDE *v* FORMIC ALDEHYDE

METHYLENE - ANILINE $(C_6H_5N)_2$ *ie* $(CH_2NPh)_2$. Anhydro formaldehyde aniline [198°]. Formed by mixing aniline and crude formic aldehyde in the cold (Tollens, *B* 17, 657, 18, 3309, Kolotoff, *Bl* [2] 43, 112). White silky crystals, *v* sol chloroform and benzene, sl sol alcohol, insol water. Decomposed by boiling with water or alcohol— $B^+H_2PtCl_4$.

Methylene-aniline C_6H_5N . Formed, together with di phenyl methylene diamine and the preceding body by the action of formic aldehyde on aniline (Pratesi, *G* 14, 355). Minute crystals, decomposed by heat *v* sl sol alcohol.

Methylene-di-aniline *v* DI-PHENYL METHYLENE DIAMINE

METHYLENE-DIBENZYLAMINE $C_{10}H_{12}N_2$ *ie* $CH_2(NHCH_2Ph)_2$ [46°] (*c* 227°). Formed by the action of methylene chloride on benzyl amine at 100° (Kempff, *A* 256, 220). Rhombic prisms, insol water, *v* sol absolute alcohol and ether.

Salts— $C_{10}H_{12}N_2 \cdot 2HCl$ [242°]. Colourless monoclinic plates— B^+2HBr silky plates— B^+2HI monoclinic plates (from benzene), sl sol water and alcohol— $B^+H_2SO_4$ 2aq colourless prisms— $B^+2H_3PO_4$ [*c* 230°]. Needles— $B^+H_2Cl_4 \cdot PtCl_4$. monoclinic plates—Aurochlor.

ide $B^+H_2Cl_4 \cdot 2AuCl_3$, golden yellow needles.—Oxalate $B^+(H_2C_2O_4)_2$ [*c* 134°]—Picrate $B^+(C_6H_3(NO_2)_3OH)_2$, not stable in a hot solution, sl sol alcohol, and water.

METHYLENE BLUE *v* TETRA METHYL DIAMIDO IMIDO DI PHENYL SULPHIDE

TRIMETHYLENE BROMHYDRIN *v* BROMO PROPYL ALCOHOL

METHYLENE BROMIDE CH_2Br_2 . Dibromo methane (97° cor) (Perkin, *C J* 45, 520), (98.5° 1 V) (Henry) SG $\frac{2}{3}$ 2493 (H₂), $\frac{1}{15}$ 24985, $\frac{25}{35}$ 24775 (P) M M 8110 at 15.9°. Formed by adding bromine to methylene iodide under water (Butlerow, *A* 111, 251, Henry, *A Ch* [5] 30, 266). Formed also by heating methyl bromide with bromine at 250° (Steiner, *B* 7, 507). Found among the bromides obtained by passing into bromine the products got by passing ethylene through a red hot tube (Norton a Noyes, *Am* 8, 362). Colourless mobile liquid, insol water, sol alcohol and ether. Converted by PCl_5 at 190° into CCl_4 and CBR_4 (Holand, *A* 240, 230). When heated with 18 volts water and excess of PbO at 145° it yields ethylene glycol, traces of ethylene oxide, and $PbBr_2$ (Jeltokoff, *B* 6, 558). Slowly converted by $SbCl_5$ into methylene chloride. Potassium phenylate KOC_6H_5 yields $CH(O_2C_6H_5)_2$.

Trimethylene bromide *v* DI-BROMO PROPANE

Tetra-methylene bromide $C_4H_8Br_2$ (189°)

Formed from ethylene cyanide by successive conversion into tetra methylene diamine and $CH_2(OH)CH_2CH_2CH_2(OH)$, the glycol being then treated with HBr (Gustavson a Demjanoff, *J pr* [2] 39, 542). The yield is small.

Penta methylene bromide $C_5H_{12}Br_2$ *ie* $CH_3(CH_2CH_2Br)_2$ (205°). Formed by saturating the corresponding di oxy pentane (derived from tri methylene diamine) with HBr (*G a D*).

METHYLENE BROMO-IODIDE *v* BROMO IODO-METHANE

METHYLENE-TETRA-BUTYL-DIAMINE $CH_3(N(C_4H_9)_2)_2$ (250°). Obtained by distilling trimethylene oxide with di isobutyl amine (Ehrenberg, *J pr* [2] 36, 124)— $B^+H_2PtCl_6$ [198°]. Pale yellow powder—Aurochloride [*c* 195°]— B^+CS_2 [54°].

METHYLENE DI ISOBUTYL DIOXIDE *v* Di isobutyl ether of FORMIC ORTHALDEHYDE, vol 11 p 571

METHYLENE CAFFEIC ACID *v* vol 1 p 659

TRIMETHYLENE DICARBAMIC ACID

* $CH_2(CH_2NHCO_2H)_2$
 Methylene ether $CH_2(CH_2NHCO_2Me)_2$ [75°]

Formed by treating trimethylene diamine with methyl carbonate (Franchimont a Klobbie, *R T C* 7, 343). Oblique rhombic plates *V* sol water, alcohol, and $CHCl_3$, m sol ether, *v* sl sol benzene and light petroleum. Pure HNO_3 yields a dinitro-derivative forming tufts of needles or prisms from alcohol [90°], and this when treated with solution of ammonia gives tri-methylene dinitramine [67°].

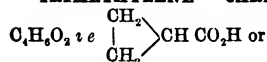
Ethyl ether $CH_2(CH_2NHCO_2Et)_2$. Tri methylene-diurethane [42°] (210° at 30 mm). When a solution of trimethylene diamine in alcohol and ether is added to an ethereal solution of chloro formic ether $ClCO_2Et$, a pp of trimethylene diamine hydrochloride is formed, and the filtrate leaves on evaporation oily

$C_3H_6(NHCO_2Et)_2$ (Fischer & Koch, *A* 232, 225) It crystallises after some time, and separates from ether in colourless prisms *V* sol ether, alcohol, and chloroform, *sl sol* ligroin, *insol* water It dissolves in acids but is reppd by alkalis

Tetra methylene dicarbamie acid *Methyl ether* $C_4H_8(NHCO_2Me)_2$ [128°] Formed from tetra methylene diamine and $ClCO_2Me$ (Dekkers, *R T C* 9, 97) Flattened needles, *sol* warm water Converted by HNO_3 into $C_4H_8(N(NO_2)CO_2Me)_2$, which separates from ether in small brilliant crystals [62°], *v sl sol* cold water, *sl sol* ether, *sol* warm alcohol, and is converted by NH_4Aq into $C_4H_8(NHNO_2)_2$ [163°]

Pentamethylene dicarbamie acid *Methyl ether* $CH_2(CH_2CH_2NHCO_2Me)_2$ [114°] Formed by treating penta methylene diamine with methyl carbonate Fine needles (from boiling water) *V sol* hot alcohol (Franchimont & Klobbie, *R T C* 7, 343) Pure HNO_3 gives a di nitro derivative, forming small brilliant prisms [37°], and this treated with aqueous ammonia gives penta methylene dimtramine [60°] together with methyl carbamate

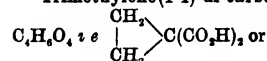
TRIMETHYLENE CARBOXYLIC ACID



$CH_2CHCH_2CO_2H$ *Isomeride of crotonic acid* [19°] (183°) Formed by heating ethylene malonic acid (trimethylene dicarboxylic acid) at 210° (Roder, *A* 227, 24, Perkin, *jun*, *C J* 47, 817, *B* 17, 57) White crystals, *m sol* water Has a powerful odour and a burning taste — CaA' , 6aq Needles, *v sol* water — BaA' , 2aq Needles, *v sol* water — AgA' Small needles (from hot water) Gives off a low boiling oil on distillation

Ethyl ether EtA' (134°) Formed by digesting the Ag salt with an ethereal solution of EtI Volatile oil, with pleasant odour Not affected by bromine in the cold, and only slowly attacked when boiled with bromine, HBr being given off

Trimethylene (1 1)-di-carboxylic acid so called



$CH_2CHCH_2CO_2H$, *Vinamic acid* *Ethylene malonic acid* [141°] Obtained by saponification of its ether Colourless triclinic crystals, $a b c = 0.7712 \times 1.8702$ Crystallises with aq (from water), *v e sol* water, *v sol* ether At 160° it begins to give off CO_2 , leaving trimethylene carboxylic acid It is not attacked by sodium amalgam When heated with bromine it gives off HBr slowly It combines, however, with HBr , forming bromo ethyl malonic acid (Perkin, *C J* 47, 814), and it combines with bromine when this is dissolved in chloroform (Fittig) Boiling dilute H_2SO_4 forms γ -oxy ethyl-malonic acid Alkaline $KMnO_4$ does not attack it (Buchner, *B* 23, 704)

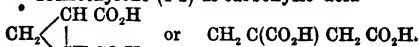
Salts—**Ammonium salt** Four-sided plates — CuA' aq Deep blue octahedra, *sl sol* hot water At 160° it becomes a light green powder — BaA' Needles — BaH_2A' , 4aq Prismatic needles — PhA' (dried at 100°) *Insol* cold, *sl sol* hot, water — AgA' Needles, *v. sl*

sol cold water, *sl sol* hot water — $AgHA''$ Long colourless needles (from water)

Ethyl ether EtA'' (213°) *VD* 632 (calc 643) *SG* $\frac{1}{18}$ 1.0646, $\frac{2}{3}$ 1.0566 Formed by the action of ethylene bromide on sodium-malonic ether (Perkin, *jun*, *C J* 47, 812, *B* 17, 54, Fittig & Roder, *A* 227, 13) Oil Not attacked when successively treated with benzyl chloride and $NaOEt$ (Perkin, *jun*, *B* 18, 1734)

Dibromide $C_3H_4Br_2(COOH)_2$ [110°] From tri methylene di carboxylic acid and Br in chloroform (Fittig & Marburg, *B* 18, 3413)

Trimethylene (1 2)-di-carboxylic acid

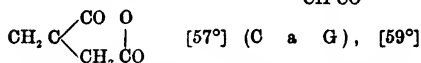
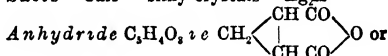


Allo itaconic acid [137°] (*C a G*), [139°] (*B*)

Formation—1 By heating its anhydride with water at 140° (Conrad & Guthzeit, *B* 17, 1187) —2 From sodium propane tetracarboxylic ether $(CO_2Et)_3CNaCH_2CNa(CO_2Et)_2$, by treatment with bromine, the resulting trimethylene tetracarboxylic ether being saponified, and the free acid heated to 230°, and then distilled under 16 mm pressure the oil which passes over at 170° to 180° is heated with water at 140° (Perkin, *jun*, *B* 19, 1056, Dressel, *A* 256, 197)

Properties—Prisms, *v sol* water, alcohol, and ether Not attacked by alkaline $KMnO_4$, or by sodium amalgam (Buchner, *B* 23, 705)

Salts— CaA'' silky crystals — AgA''

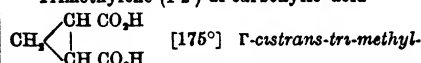


(Buchner) Formed by heating trimethylene

tricarboxylic acid $CH_2 \begin{array}{c} \diagup \\ \diagdown \end{array} \begin{array}{l} CHCO_2H \\ C(CO_2H)_2 \end{array}$ or possibly

$CH_2C(CO_2H)CH(CO_2H)_2$ at 184° to 190° for a long time (*C a G*) Needles, *sl sol* ether

Trimethylene (1 2)-di-carboxylic acid



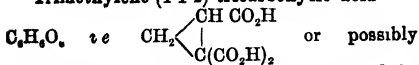
ene-1 2 di carboxylic acid Formed by saponification of the di methyl ether (Eduard Buchner, *B* 23, 705) Compact apparently rhombic crystals (from water), containing no water of crystallisation Grouped needles (from ether) Less *sol* water than the isomeric glutaric acid, but more *sol* ether Distils without splitting off water Acetyl chloride forms no anhydride Not oxidised by permanganate in alkaline solution Not reduced by sodium amalgam Chloride of iron gives a weak red brown colour

Salts—The zinc salt is more *sol* cold than hot water It crystallises in nodular groups of needles A solution of the ammonium salt gives no pp with $CaCl_2$ and $BaCl_2$, but white crystalline pps with silver and lead salts By heating the silver salt a white body sublimes in needles; probably the anhydride

Di methyl ether $C_3H_4(CO_2Me)_2$ (205°–215° at 718 mm) Formed by heating acrylo-di azo acetic ether $C_3H_4N_2(CO_2Me)_2$ for 40 minutes to 160°–185°, Saponified by boiling with aqueous potash.

Isomerides of trimethylene dicarboxylic acid
v ITACONIC, CITRACONIC, MESACONIC, and GLUTA
CONIC ACIDS

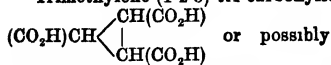
Trimethylene (1 1 2) tricarboxylic acid



$CH_2 \cdot C(CO_2H) \cdot CH(CO_2H)_2$ [184°] Formed by saponification of its ether, which is prepared by the action of $\alpha\beta$ di bromo propionic ether on di sodio malonic ether (Conrad & Guthzeit, *B* 17, 1185) Its ether is also formed by the action of a bromo acrylic ether upon sodio malonic ether $CHNa(CO_2Et)_2$ (Michael, *J pr* [2] 35, 132, 351, *Am* 9, 121) Prisms (from water) At 184°-190° it is split up into CO_2 and the dicarboxylic acid or its anhydride

Tetra-ethyl ether Et_4A'' (276°) SG 15 1 127 Colourless liquid Does not react with $NaOEt$ and $BzCl$

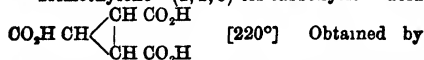
Trimethylene (1 2 3)-tri-carboxylic acid



$CO_2H \cdot CH \cdot C(CO_2H) \cdot CH_2 \cdot CO_2H$ [150°-153°] Formed by heating the tetra carboxylic acid (1 1 2 3) at 200° for a long time (Perkin, *B* 17, 1654, *C J* 47, 826) Colourless crystalline solid V sol water, alcohol, and acetone, sl sol benzene, chloroform, ligroin, and CS_2 On heating it yields a sublimate, probably of an anhydride

Salts — $A''Ag$, white granular pp — $A''Ca$, crystalline pp, soluble in cold water, nearly insoluble in hot The cupric salt is a beautiful light green pp, sl sol water The Ba and Pb salts are white pps

Trimethylene (1, 2, 3) tri-carboxylic acid



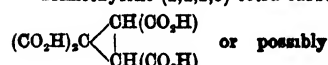
saponifying its methyl ether with alcoholic potash (Buchner, *B* 21, 2641) Small aggregates of needles (from ether), v sol alcohol and water, sl sol ether Not affected by bromine or by $KMnO_4$ Its ammonium salt crystallises in plates

Methyl ether Me_4A'' [61°] (267°) at 732 mm, (224° at 180 mm) Obtained by distilling the compound of methyl fumarate with methyl diazo acetate Needles (from alcohol or water), v sol alcohol, ether, benzene, and petroleum-ether

Anhydride $C_3H_2(CO_2H) \begin{array}{c} \diagup CO \\ \diagdown CO \end{array} O$ [189°]

(265° at 75 mm) Obtained by distilling the acid under reduced pressure Small prisms, v sol water and alcohol, m sol ether Its Pb, Ba, and Ag salts are m sol water

Trimethylene (1,1,2,3) tetra-carboxylic acid



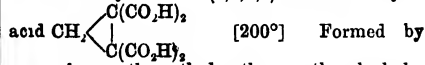
$(CO_2H)_2C \cdot C(CO_2H) \cdot CH_2 \cdot CO_2H$ [95°-100°] The tetra-ethyl ether of this acid is obtained by the action of di bromo succinic ether on di sodio-malonic ether (Perkin, *B* 17, 1652, *C J* 47, 824) Crystalline colourless solid V sol water, alcohol, ether, and acetone, sl sol ligroin and benzene. Very strong acid The acid loses CO_2

on heating to 200° giving the (1, 2, 3) tri carboxylic acid

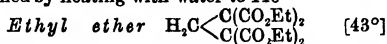
Salts — $A''Ag$, white amorphous pp — $A''Ca$, crystalline pp, more soluble in cold water than in hot

Tetra-ethyl ether Et_4A'' , (246° at 85 mm), thick colourless oil

Tri-methylene (1,1,2,2)-tetra-carboxylic

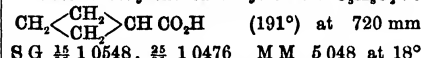


saponifying the ethyl ether with alcoholic $KOHAq$ (Dressel, *A* 256, 196) Large shining crystals (from water) When heated to 230° it gives off $2CO_2$ and H_2O and is converted into the anhydride of trimethylene dicarboxylic acid, and from this the dicarboxylic acid itself can be obtained by heating with water to 140°



(187°) Formed by the action of bromine on the disodium compound of propane tetra carboxylic ether (Dressel, *A* 256, 194, cf Perkin, *B* 19, 1056) Long needles, v sol ether, alcohol, $CHCl_3$, and benzene (D)

Tetra-methylene carboxylic acid $C_4H_4O_6$ is



SG $\frac{1}{15}$ 1 0548, $\frac{2}{15}$ 1 0476 MM 5 048 at 18° μ_D 1 4403 at 26° Formed by heating tetra methylene dicarboxylic acid at 200°, CO_2 being given off (Perkin, *B* 16, 1795, *C J* 51, 8) Colourless oil, smelling like butyric acid, sl sol water, miscible with alcohol and ether Not attacked by bromine below 100°

Reaction—The calcium salt distilled with lime gives C_2H_4 , H_2 , CH_4 , CO , di tetramethylenyl ketone, and tetramethylenyl methyl ketone (Colman & Perkin, jun, *C J* 51, 228, *B* 19, 3112)

Salts — AgA' sparingly soluble white pp — CaA' , 5aq

Ethyl ether Et_4A' (151°) at 720 mm (P), (162°) (Freund, *B* 21, 2694)

Chloride $C_4H_4OCl_2$ (143°) (F) Obtained by heating the amide with PCl_5

Amide $C_4H_4CONH_2$ [198°] (c 240°)

Formed by heating the ammonium salt of the acid to 250° (Freund, *B* 21, 2694) Plates (from alcohol), v sol water, ether, chloroform, benzene May be sublimed Decomposed by treatment with bromine and KOH

Nitrile C_4H_4CN (150°) Formed by distilling the amide of tetra methylene carboxylic acid with phosphoric anhydride (Freund, *B* 21, 2696) Colourless oil, with pleasant odour Is converted into the amine by reducing with sodium and alcohol

Anilide $C_4H_4CONHPh$ [111°] Formed by heating the amide with aniline until no more ammonia is given off (Freund) Long needles (from alcohol), sl sol hot water

Anhydride $(C_4H_4CO)_2O$ (160°) Formed by distilling the sodium salt with the chloride of the acid (F)

Tetra-methylene (1,1)-dicarboxylic acid

$C_4H_4O_6$ is $CH_2 \begin{array}{c} \diagup CH_2 \\ \diagdown CH_2 \end{array} C(CO_2H)_2$ [156°] From the ether (Perkin, *C J* 51, 4) Monoclinic crystals (from ether), $a b c = 1.0324 \ 1.11854$, $\beta = 88^\circ 58'$. V. sol. ether and benzene, nearly

insol chloroform and ligroin, v sol water A few degrees above its melting point it splits off CO_2 , becoming tetramethylene carboxylic acid

Salts — $\text{Ag}_2\text{A}''$ — $\text{CuA}''\text{aq}$ — $\text{PbA}''\text{aq}$ — $\text{BaA}''\text{aq}$

Ethyl ether $\text{Et}_2\text{A}''$ (221°) at 720 mm SG $\frac{1}{4}$ 1.0633, $\frac{20}{4}$ 1.0405. M M 9.940 at 18.9° μ_n 1.433 Formed from malonic ether, trimethylene bromide, and NaOEt (Perkin, B 16, 1793, C J 51, 2) Oil, smelling like camphor

Tetramethylene (1,2)-di-carboxylic acid
 $\text{CH}_2\text{CH}(\text{CO}_2\text{H})$
 $\text{CH}_2\text{CH}(\text{CO}_2\text{H})$ [130°] Formed by heating

the tetra carboxylic acid to 180°–200° (Perkin, B 19, 2042, C J 51, 22) Colourless feathery crystals (from water) V sol water, alcohol, and ether, more sparingly sol benzene and ligroin By alkaline KMnO_4 it is oxidised to oxalic acid

Salts — $\text{A}''\text{Ag}_2$ heavy white pp — $\text{A}''\text{Ba}''$ sparingly soluble six sided transparent tables
Diethyl ether $\text{A}''\text{Et}_2$ (238°–242°), colourless liquid

Anhydride $\begin{array}{c} \text{CH}_2\text{CHCO} \\ \text{CH}_2\text{CHCO} \end{array} \text{O}$ [78°], colour-

less crystals, easily soluble in alcohol, sparingly in ether and benzene Formed by heating the acid to 300° Reconverted into the acid by boiling with water Heated with resorcin it gives a beautifully fluorescent condensation product

Tetramethylene (1,3) dicarboxylic acid

$\text{CO}_2\text{HCH} \begin{array}{c} \text{CH}_2 \\ \text{CH}_2 \end{array} \text{CHCO}_2\text{H}$ *Homotaconic acid* [171°] Formed by boiling its ether with fuming HClAq (Markownikoff a Krestownikoff, A 208, 333) Prisms, v sol hot water and alcohol, sl sol ether May be sublimed Does not combine with bromine, but when heated with Br it gives off HBr and CO_2 It is not reduced by sodium amalgam Does not form an anhydride Does not unite with HBr or HI

Salts — $\text{PbA}''\frac{1}{2}\text{aq}$ Crystalline crusts — AgA'' Amorphous pp, not decomposed by boiling water

Methyl ether $\text{Me}_2\text{A}''$ (220°)

Ethyl ether $\text{Et}_2\text{A}''$ (230°) Formed in very small quantity when $\text{CH}_2\text{CH}(\text{OEt})\text{CO}_2\text{Et}$ is prepared by the action of dry NaOEt on a chloro propionic ether (M a K)

Tetramethylene (1,1,2,2) - tetra - carboxylic acid $\begin{array}{c} \text{CH}_2\text{C}(\text{CO}_2\text{H})_2 \\ \text{CH}_2\text{C}(\text{CO}_2\text{H})_2 \end{array}$ [145°–150°] Obtained by

saponification of the tetra-ethyl ether, which is formed by the action of bromine upon the di sodium derivative of butane tetra carboxylic ether $\text{CH}_2\text{CNa}(\text{CO}_2\text{Et})_2$ (Perkin, B 19, 2041,

$\text{CH}_2\text{CNa}(\text{CO}_2\text{Et})_2$, C J 51, 21) Colourless crystals Easily soluble in water, alcohol, and ether, more sparingly in benzene and ligroin It evolves CO_2 at its melting point, and is converted into the di carboxylic acid

Tetramethylene (1,1,3,3) - tetracarboxylic acid *Ethyl ether*

$(\text{CO}_2\text{Et})_2\text{C} \begin{array}{c} \text{CH}_2 \\ \text{CH}_2 \end{array} \text{C}(\text{CO}_2\text{Et})_2$ Formed by the action of methylene iodide on the di sodium de-

rivative of propane tetra carboxylic ether in an alcoholic solution on the water bath (Dressel, A 256, 198) Colourless viscid oil, distilling with some decomposition between 220° and 250° at 15 mm

Pentamethylene dicarboxylic acid

$\begin{array}{c} \text{CH}_2-\text{CHCO}_2\text{H} \\ \text{CH}_2-\text{CHCO}_2\text{H} \end{array}$ [160°]

Preparation — Disodium pentane tetra carboxylic ether,

$(\text{CO}_2\text{Et})_2\text{CNaCH}_2\text{CH}_2\text{CH}_2\text{CNa}(\text{CO}_2\text{Et})_2$ is converted by Br into $\text{CH}_2 \begin{array}{c} \text{CH}_2-\text{C}(\text{CO}_2\text{Et})_2 \\ \text{CH}_2-\text{C}(\text{CO}_2\text{Et})_2 \end{array}$, which

yields pentamethylene tetracarboxylic acid on saponification, and this acid is decomposed by heat into CO_2 and pentamethylene dicarboxylic acid This is purified by conversion into its ethyl salt (Perkin, jun, C J 51, 244, B 18, 3250)

Properties — Nodules (from water) V sol hot water, alcohol, and acetic ether, sl sol ether

Salts — AgA'' Very stable white pp

Ethyl ether $\text{Et}_2\text{A}''$ (c 250°)

Anhydride $\begin{array}{c} \text{CH}_2-\text{CHCO} \\ \text{CH}_2-\text{CHCO} \end{array} \text{O}$

[64°–67°] Formed by heating the acid to 300° M sol alcohol and ether, sl sol CS_2 , insol cold, but saponified by hot, $\text{Na}_2\text{CO}_3\text{Aq}$ With resorcin and H_2SO_4 it gives the fluorescein reaction

Penta-methylene tetracarboxylic acid

$\begin{array}{c} \text{CH}_2-\text{C}(\text{CO}_2\text{H})_2 \\ \text{CH}_2-\text{C}(\text{CO}_2\text{H})_2 \end{array}$ Formed as above Sol. ether

TETRAMETHYLENE CARBOXYLIC ALDE

HYDE $\text{CH}_2 \begin{array}{c} \text{CH} \\ \text{CH}_2 \end{array} \text{CHCHO}$ (115°–117°) From

calcium tetramethylene carboxylate by distilling with calcium formate (Colman a Perkin, C J 51, 238) Oil, smelling like isobutyric aldehyde Gives a purple colour with rosamine hydrochloride which has been bleached by SO_2 , reduces ammoniacal AgNO_3Aq , combines with NaHSO_4 , reacts with hydroxylamine and with phenyl hydrazine

TRIMETHYLENE CHLORHYDRIN v

CHLORO PROPYL ALCOHOL

METHYLENE CHLORIDE CH_2Cl_2 *Di-chloro methane* Mol w 85 (41.8°) (Thorpe, C J 37, 195) SG $\frac{1}{4}$ 1.3778 (T), $\frac{15}{4}$ 1.3377, $\frac{20}{4}$ 1.3220 M M 4.313 at 11.9° (Perkin C J 45, 527) C E (0°–10°) 001335, (0°–40°) 001416 SV 65.12 H Cp 106800 (Berthe lot a Ogier, Bl [2] 36, 68)

Formation — 1 From chlorine and methyl chloride in sunlight (Regnault, A 33, 328, A Ch [2] 70, 377) — 2 By the action of zinc dust and ammonia on chloroform, the yield being very small (Perkin, C N 18, 106) — 3 By the action of chlorine on methylene iodide (Butlerow, A 107, 110, 111, 251, Z [2] 5, 276) — 4 By the action of AlCl_3 on the compound of $\text{ClCO}_2\text{CCl}_2$ with ClCO_2CH_3 (Hentschel, J pr [2] 86, 474)

Preparation — Aqueous HCl is cautiously added to a mixture of alcohol (5 vols), chloroform (1 vol), and zinc Sufficient heat is pro-

duced in the reaction to distil over much of the methylene chloride. The product is fractionally distilled (Greene, *C R* 89, 1077, *A Ph S* 18, 847, *C N* 50, 75, *A C J* 1, 522)

Properties—Oil. Like CCl_4 (but unlike CHCl_3 and CHCl_2) it exerts a poisonous action when inhaled (Regnault & Villejean, *C R* 100, 1146)

Reactions—1 Converted by ICl or ICl_2 into CHCl_3 , and hexachloro benzene. Converted by IBr into iodoform and di chloro di iodo methane (Holand, *A* 240, 234). Converted by IBr , into CBr_4 , CHBr_3 , and C_2Br_4 .—2 Heated with KI , iodine, and alcohol, it gives CH_3I , EtI , and alcohol (H).—3 Iodine at 200° gives methylene iodide (H).—4 Bromine at 170° forms CHBrCl_2 (91°), and a little CBr_2Cl_2 [38°] (150°) (Arnhold, *A* 240, 204).—5 Alcoholic NaOAc forms $\text{CH}_3(\text{OEt})$, and acetic acid (A).—6 Converted by alcoholic NH_3 at 125° into so called hexa methylene diamine (Holand, *A* 240, 225). Aqueous ammonia at 140° forms NH_4Cl , methylamine hydrochloride and formic acid (André, *C R* 102, 1474).—7 Mixed with benzene it is converted by AlCl_3 into anthracene and other products (Friedel & Crafts, *A Ch* [6] 11, 264).—8 Water at 200° forms HCl , formic acid, MeCl , and MeOH (André).—9 With H_2S it forms a crystalline compound $\text{CH}_2\text{Cl}_2(\text{H}_2\text{S})_2 \cdot 23\text{aq}$ (Forcrand, *A Ch* [5] 28, 17)

TRIMETHYLENE CHLORIDE v ω 8 Di-CHLORO PROPANE

METHYLENE CHLORO BROMIDE v CHLORO BROMO METHANE

Tri methylene chloro-bromide v CHLORO BROMO PROPANE

METHYLENE CHLORO IODIDE $\text{CH}_2\text{CH}_2\text{I}$, *Chloro iodo methane* (109°) VD 88 14 SG $\frac{11}{11}$ 2 447, $\frac{145}{145}$ 2 444. Formed by the action of ICl on methylene iodide or of iodine on $\text{IHg CH}_2\text{Cl}$ (Sakurai, *C J* 41, 361, 47, 198). Oil.

TRIMETHYLENE CYANHYDRIN v *Nitrile of γ Oxy-butyric acid*

TRIMETHYLENE CYANIDE $\text{C}_3\text{H}_6\text{N}_2$ v $\text{CH}_2(\text{CH}_2\text{CN})_2$ *Glutaronitrile* (274°) (H), (286°) (Perkin, *C J* 55, 702), (203° at 100 mm), (142° at 10 mm) (Krafft & Noerdlinger, *B* 22, 817) SG $\frac{11}{11}$ 9952, $\frac{2}{2}$ 9894 MM 5 136 (P). Formed from trimethylene bromide and alcoholic KCy (Henry, *Bl* [2] 43, 618, *C R* 100, 742). Liquid, sol water, alcohol, and chloroform, insol ether and CS_2 . Yields glutaric acid on saponification. Sodium reduces it in alcoholic solution to pentamethylene diamine and piperidine.

TRIMETHYLENE-DI ETHYL ALKINE v ETHYL OXYPROPYL AMINE

METHYLENE-ETHYL AMINE $\text{C}_3\text{H}_7\text{N}$ v EtNCH_2 (208° v) VD 2. Formed by the action of ethylamine on formic aldehyde (tri oxymethylene) (Kolotoff, *Bl* [2] 43, 112, *J R* 17, 231). Liquid, with unpleasant odour, sol cold water, but separates again on warming, v sol alcohol. HCl splits it up into ethylamine and trioxymethylene (formic paraldehyde).— $\text{B}'\text{H}_2\text{PtCl}_6$. Yellow crystalline pp. An isomeride $(\text{CH}_3)_3\text{N}, \text{Et}$, of this base is described by Lermontoff (*B*, 7, 1252) as an oil formed by heating

ethylamine with alcoholic methylene iodide at 100° — $(\text{CH}_3)_3\text{N}, \text{Et}, \text{H}_2\text{PtCl}_6$. Amorphous.

Methylene tetra ethyl-diamine $\text{C}_4\text{H}_{12}\text{N}_2$ v $\text{CH}_2(\text{NEt}_2)_2$ *Tetra ethyl-di-amido methane* (169° v) SG 10. Formed by heating trioxymethylene (formic paraldehyde) with diethylamine in sealed tubes at 100° (Kolotoff, *Bl* [2] 43, 112, Ehrenberg, *J pr* [2] 36, 118). Liquid, with peppery odour, sol water, miscible with alcohol, ether, and CHCl_3 . Split up by dilute acids, even by oxalic acid, into NH_4Et , and formic aldehyde. Combines with CS_2 , forming $\text{C}_4\text{H}_{12}\text{N}_2\text{OS}_2$.

Tetra - methylene tetra - ethyl - tetramine $\text{C}_{12}\text{H}_{28}\text{N}_4$ v $\text{CH}_2 \left\langle \begin{smallmatrix} \text{NEt CH}_2 \text{NEt} \\ \text{NEt CH}_2 \text{NEt} \end{smallmatrix} \right\rangle \text{CH}_2$. Formed by heating methylene iodide with alcoholic ethylamine at 100° (Lermontoff, *B* 7, 1252). Liquid, yielding amorphous salts— $\text{B}'\text{H}_2\text{PtCl}_6$, sol water.

METHYLENE-ETHYL PHTHALIMIDINE

$\text{C}_{11}\text{H}_{11}\text{NO}$ v $\text{C}_6\text{H}_5 \left\langle \begin{smallmatrix} \text{C}(\text{CH}_2) \\ \text{CO} \end{smallmatrix} \right\rangle \text{NEt}$. Obtained by heating $\text{C}_2\text{H}_5\text{N}_3\text{O}$, which is produced by adding aqueous ethylamine to phthalyl acetic acid (Mertens, *B* 19, 2369). Colourless oil, smelling of fresh carrots. Volatile with steam, v sol alcohol and ether.

METHYLENE DI-ETHYL DISULPHIDE

$\text{C}_4\text{H}_{10}\text{S}_2$ v $\text{CH}_3(\text{SEt})_2$ *Formic aldehyde ethyl mercaptal* *Ethyl derivative of di thio formic orthoaldehyde* (178° – 181°) SG 22 987. Formed from methylene chloride and NaSEt in alcoholic solution (Niederst, *A* 186, 391, Fromm, *A* 253, 155).

METHYLENE DI ETHYL DISULPHONE

$\text{C}_4\text{H}_{10}\text{S}_2\text{O}_2$ v $\text{CH}_3(\text{SO}_2\text{Et})_2$ (104°). Formed by the action of KMnO_4 and H_2SO_4 on $\text{CH}_3(\text{SEt})_2$ (Fromm, *A* 253, 156, cf Baumann, *B* 19, 2811). Needles, v sol water and alcohol, sol ether. Chlorine forms $\text{CCl}_2(\text{SO}_2\text{Et})_2$ [99°] crystallising in needles, while bromine produces $\text{CBr}_2(\text{SO}_2\text{Et})_2$ [132°].

Di-methylene di-ethyl trisulphone

$(\text{Et SO}_2\text{CH}_2)_2\text{SO}_2$ [149°]. Formed by treating formic aldehyde with H_2S , dissolving the product in aqueous NaOH , shaking with EtBr , and oxidising the product with KMnO_4 (Baumann, *B* 23, 1876). Sparingly soluble colourless plates.

METHYLENE-FURFURANE TRIHYDRIDE

$\text{C}_5\text{H}_8\text{O}_3$ v $\text{CH}_2 \left\langle \begin{smallmatrix} \text{CH}_2 \text{C}(\text{CH}_2) \\ \text{CH}_2 \text{O} \end{smallmatrix} \right\rangle$ (111°) at 718 mm.

Formed by the action of solid KOH on methyl bromo propyl ketone (Lupp, *B* 22, 1207). Mobile liquid, sol water. Yields a hydrate. When heated with 5 p.c. HCl aq at 100° it yields aceto propyl alcohol (methyl oxypropyl ketone). Probably identical with trimethylenyl methyl ketone of Perkin, jun (*B* 17, 1440).

TRI-METHYLENE GLYCOL $\text{C}_3\text{H}_8\text{O}_3$ v $\text{CH}_2(\text{CH}_2\text{OH})_2$ (214°) SG $\frac{1}{1}$ 10625 (Z), $\frac{186}{1}$ 10536 (F) CE (0° – 10°) 00060 SV 84 (Zander, *A* 214, 178, Lossen, *A* 254, 59). One of the products of the fermentation of glycerin by schizomycetes (Freund, *M* 2, 636).

Formation—1 By saponifying its diacetyl derivative with baryta water (Reboul, *A Ch* [5] 14, 491).—2 By warming trimethylene bromide with moist Ag_2O (Beilstein & Wiegand, *B* 15,

1497)—8 By allowing $\text{CH}_2(\text{CH}_2\text{Br})_2$ to stand for some time with a large excess of water (Niederist, *M* 3, 889)—4 By boiling trimethylene bromide with dilute aqueous K_2CO_3 (Z)

Properties—Viscid liquid, with sweet taste, miscible with water

Reactions—1 Fuming HClAq at 100° converts it into $\text{CH}_2(\text{CH}_2\text{Cl})_2$.—2 Trimethylene glycol (35 g) heated with aldehyde (12 g) at 100° yields the ethyldene derivative (*v infra*)

Di acetyl derivative $\text{CH}_2(\text{CH}_2\text{OAc})_2$ (210° cor) (c 111°) SG d 991, n_D 1.070 S 11 Formed by boiling $\text{CH}_2(\text{CH}_2\text{Br})_2$ with NaOAc (Reboul)

Ethyldene derivative $\text{C}_5\text{H}_{10}\text{O}_2$ v e
 $\text{CH}_2 \begin{smallmatrix} \text{CH}_2\text{O} \\ \text{CH}_2\text{O} \end{smallmatrix} \text{CHCH}_3$ VD 3.62 (calc 3.53)

Obtained by heating the glycol with aldehyde at 100° , the yield being nearly the theoretical (Loohert, *A Ch* [6] 16, 49) Colourless liquid, with slight aldehydic odour. Dissolves in 1½ volumes of water, *v* sol alcohol and ether. Separated from its aqueous solution by CaCl_2 and by KOH . Saponified by boiling water, alkalis, and dilute acids. With PCl_5 it yields aldehyde and $\text{CH}_2(\text{CHCl})_2$

Amyldene derivative $\text{C}_6\text{H}_{10}\text{O}_2$ v e
 $\text{CH} \begin{smallmatrix} \text{CH} \\ \text{CH} \end{smallmatrix} \text{O} \text{CHCH}_2\text{C}_4\text{H}_9$ (c 165°) SG d 995 VD 5.03 (calc 4.98) Formed by heating trimethylene glycol (15 g) with valeric aldehyde (20 g) in a sealed tube at 125° . Colourless mobile liquid, *v* sol water, *v* sol alcohol and ether. Saponified by boiling water

Heptyldene derivative $\text{C}_8\text{H}_{10}\text{O}_2$ v e
 $\text{CH} \begin{smallmatrix} \text{CH}_2\text{O} \\ \text{CH}_2\text{O} \end{smallmatrix} \text{CHCH}_2\text{C}_6\text{H}_{13}$ (c 216°) SG d 933 From the glycol (30 g) and heptioic aldehyde (cyanthol) at 160° (L)

Bromhydrin v Bromo propyl alcohol
TRI METHYLENE IMINE $\text{C}_3\text{H}_5\text{N}$ v e
 $\text{CH}_2 \begin{smallmatrix} \text{CH}_2 \\ \text{CH}_2 \end{smallmatrix} \text{NH}$ (65° – 80°) Formed, together with a polymeride $\text{C}_6\text{H}_9\text{N}$ (160° – 167°), from $\text{CH}_2\text{BrCH}_2\text{CH}_2\text{NH}_2$ and NaOH in the cold (Gabriel & Weiner, *B* 21, 2669) Volatile liquid, smelling like NH_3 and fuming in the air. Eagerly combines with CS_2 , $\text{B}'\text{H}\text{AuCl}_4$, $\text{B}'\text{H}\text{PtCl}_4$, $\text{B}'\text{C}_6\text{H}(\text{NO})_2\text{OH}$ (167°)

METHYLENE IODIDE CH_2I_2 *Di iodo methane* [4°] (152° at 330 mm) SG d $\frac{4}{5}$ 3.2853, $\frac{25}{4}$ 3.2656 MM 18.827 at 15° (Perkin, *C J* 45, 464)

Formation—1 By heating iodoform (4 mols) with NaOEt (9 mols) dissolved in alcohol (Butlerow, *A* 107, 110, 111, 242, *cf* Bruning, *A* 104, 187)—2 By heating chloroform with HIAq at 130° (Bljuduchov, *Z* [2] 7, 91)—3 By heating iodoform (50 g) with conc HIAq (200 g) to boiling (127°) and adding phosphorus (Lieben, *Z* 1868, 712, Baeyer, *B* 5, 1095)—4 From methylene chloride and CaI_2 at 75° (Spindler, *A* 231, 262)—5 By warming a mixture of iodoform (5 pts), water (2 pts) and reduced iron (5 pts), and fractionally distilling *in vacuo* (Cazeneuve, *C R* 98, 869)—6 An alcoholic solution of iodoform is decomposed by light, yielding CHI_3 and iodine. Oxalic acid accelerates the reaction (Mulder, *R T C* 7, 316)

Properties—Yellowish liquid, boiling with partial decomposition at 180° .

Reactions—1 Potassium has no action in the cold, but on heating it acts with explosive violence.—2 Heated with copper and water, the products are cuprous iodide and a mixture of CO , methane, CO , and ethylene (Butlerow, *A* 120, 356)—3 Silver acetate forms the diacetyl derivative of formic orthaldehyde $\text{CH}_2(\text{OAc})_2$ —4 Silver oxalate yields formic paraldehyde (trioxymethylene)—5 Chlorine yields CH_2Cl_2 —6 Bromine forms CH_2Br_2 (Arnhold, *A* 240, 207)—7 Converted by PCl_5 at 70° into methylene chloride and a trace of chloroform (Höland, *A* 240, 227)—8 Alcoholic Na_2S yields thioformic aldehyde—9 NMe_3 combines with formation of $(\text{CH}_2\text{I})\text{NMe}_3\text{I}$ —10 Aniline forms $\text{CH}_2(\text{NHPh})_2$ —11 Mercury forms IHgCH_2I (Sakurai, *C J* 37, 658)

Trimethylene iodide v Di iodo propane
DI-TETRAMETHYLENE KETONE v Di-tetramethylenyl ketone

METHYLENE-MALONIC ETHER $\text{C}_5\text{H}_8\text{O}_4$ v e
 $\text{CH C}(\text{CO}_2\text{Et})_2$ (156°) Formed by heating malonic ether with methylene iodide and NaOEt (Zelnitsky, *B* 22, 3294) Mobile liquid. Isomeric with fumaric and maleic ethers. With bromine it yields $\text{CH}_2\text{Br CBr}(\text{CO}_2\text{Et})_2$ (185° – 190° at 75–85 mm)

Polymeride $[\text{CH}_2\text{C}(\text{CO}_2\text{Et})_2]_n$ [156°] Ac companies the preceding (Z) Minute amorphous granules, insol water, *m* sol ether and alcohol. Its molecular weight determined by Raoult's method is 342 (calc 344)

DI-METHYLENE-DI-METHYL-DIAMINE $\text{C}_4\text{H}_{10}\text{N}_2$ v e (CH_2) $_2\text{N}_2\text{Me}_2$ VD ($H=1$) 41.5 (calc 43) Formed by passing a mixture of trimethylamine and hydrogen through a red hot tube (Romeny, *B* 11, 835) Crystalline. Hot dilute HClAq splits it up into formic aldehyde and methylamine— $\text{B}'\text{H PtCl}_4$

A base, boiling at about 207° , formed by the action of methylamine upon trioxymethylene (formic paraldehyde) is perhaps identical with the preceding (Kolottoff, *Bl* [2] 45, 253)

TRIMETHYLENE-HEXA-METHYL-DIAMINE $(\text{C}_2\text{H}_5)_6\text{N}_2$ Formed by heating trimethylene bromide ($\text{C}_2\text{H}_5\text{Br}$) with trimethylamine

Salts— $\text{B}'\text{H}\text{Br}_2$ aq soluble colourless needles— $\text{B}'\text{HCl}_2\text{PtCl}_4$ sparingly soluble (Roth, *B* 14, 1351)

METHYLENE METHYL BUTENYL DIKETONE $\text{CH}_3\text{COCH}_2\text{COCH}_2\text{C}_4\text{H}_7$ *Acetyl mesityl oxide* *Acetyl angelcyl methane* (205°)

Formed as a secondary product in the preparation of methylenedimethyldiketone by the action of acetone and NaOEt on acetic ether (Claisen & Ehrhardt, *B* 22, 1012, *Bl* [3] 1, 498) Oil, soluble in aqueous alkalis. Its alcoholic solution gives a red colour with FeCl_3 . Cupric acetate in concentrated solutions gives a dark green pp— CuA [123°] Crystalline, *v* sol warm alcohol and ether

METHYLENE DIMETHYL ETHER v Di-methyl ether of Ortho Formic aldehyde

METHYLENE METHYL ETHYL DIKETONE $\text{C}_5\text{H}_8\text{O}_4$ v e $\text{C}_2\text{H}_5\text{COCH}_2\text{COCH}_3$ *Acetyl propionyl methane* (158°) SG d 9538 Formed by the action of EtOAc and NaOAc upon methyl ethyl ketone (Claisen & Ehrhardt, *B* 22, 1014)— $\text{Cu}(\text{C}_2\text{H}_3\text{O}_2)_2$ [179°] Slender blue needles (from hot alcohol)

METHYLENE METHYL HEXYL DIKETONE $C_{16}H_{30}O_2$, ν_e CH_2 , CO , CH_2 , CO , C_6H_{13} (229°). Formed from methyl hexyl ketone, $AcOEt$, and $AcONa$ (Claisen a Ehrhardt, *B* 22, 1015) liquid— $Cu(C_{10}H_{19}O_2)_2$ (122°) Crystalline

METHYLENE DI-METHYL DIKETONE $C_8H_{14}O_2$, ν_e CH_2 , $(CO\ CH_2)_2$ *Di-acetyl methane acetyl-acetone* (136°) SG μ 987 (Combes)

Preparation—1 By slowly adding acetone (1 mol) to a mixture of acetic ether ($3\frac{1}{2}$ mols) and dry $NaOEt$ (1 mol) heated on the water bath. The product is poured into ice cold water, and the aqueous liquid mixed with $HOAc$, and cupric acetate, which ppts $Cu(C_4H_7O_2)_2$. The yield is 35 p.c. of the weight of acetone employed (Claisen, *Bl* [3] 1, 498)—2 Acetone (5.8 pts) mixed with $EtOAc$ (33 pts) is treated in the cold with sodium wire (2.3 pts). When most of the sodium is dissolved, the mixture is heated on a water bath. The product is pptd as cupric salt, the yield of ketone being 55 p.c. of the acetone employed (Claisen a Ehrhardt, *B* 22, 1009). In either case the diketone is obtained from the copper compound by adding dilute H_2SO_4 and extracting with ether—3 From the compound $C_{12}H_{21}O_4AlCl_2$ (obtained from acetyl chloride and $AlCl_3$) by adding water and extracting with chloroform. The yield is 85 p.c. of the theoretical (Combes, *A Ch* [6] 12, 211).

Properties—Colourless liquid with pleasant acetic odour, sol water, v sol aqueous HCl , miscible with alcohol, ether, and chloroform. Not attacked by PCl_5 .

Reactions—1 *Phenyl hydrazine* forms phenyl dimethyl pyrazole $CMe<\overset{CH}{NPh}>CMe$ (270.5°) (Combes, *Bl* [2] 50, 145)—2 *Potash* decomposes it into acetone and $KOAc$ —3 PCl_5 yields HCl and $C_6H_5Cl_2$ (145°), a di chloro amylene which readily combines with bromine—4 *Sodium amalgam* yields isopropyl alcohol and pinacene. But in acid solution the products of reduction by sodium amalgam are di-oxy pentane $CH_3CH(OH)CH_2CH(OH)CH_3$, and a tetrahydric alcohol analogous to pinacene—5 Conc $HIAq$ at 185° reduces it to pure n pentane (38°). At lower temperatures it forms $CH_3(CHICH_2)_2$ and $CH_3CHI(CH_2CH_2CH_3)_2$ —6 Excess of *chlorine* in sunlight yields $(COCl, CO)_2CH_2$ as final product—7 *Bromine* attacks the diketone rigorously, finally producing $(CBr, CO)_2CH_2$ (108°)—8 *Oxidation* by CrO_3 or by $KMnO_4$ yields acetic acid—9 Warm dilute *nitric acid* produces $CH_3(NO_2)_2$ and $HOAc$ —10 Dry *ammonia* passed into its ethereal solution ppts white pearly scales of the ammonium salt $CH(NH_4)(CO\ CH_3)_2$ —11 *Ethylene diamine* (2 mols) forms $C_4H_{12}(N\ CMe\ CH_2\ Ac)_2$ [111°] which yields a violet cupric salt $C_{12}H_{24}N_2O_4Cu$ [137°] and a hydrochloride $C_{12}H_{24}N_2O_4H_2Cl_2$ melting above 280°. Other diamines act in like manner (Combes, *C R* 108, 1252)—12 *Umetes* with *benzidine* (1 mol) forming a base melting at 195° (Combes)—13 *Aldehyde ammonia* ($\frac{1}{2}$ mol) at 100° forms di-acetyl tri-methyl-pyridine dihydride $C_{12}H_{17}NO_2$ [153°] (Combes, *Bl* [3] 1, 14). 14 Toluene *m* diamine at 100°, followed by H_2SO_4 , yields amido trimethyl quinoline $C_8H_{11}N_3$ [191°] (Combes, *C R* 108, 1254)—15 SO_2Cl_2 forms the chloro derivative $C_8H_9ClO_2$ (156°) (Combes, *C R* 111, 272)

Salts— $CH(NH_4)(CO\ CH_3)_2$ Pearly scales, pptd by passing NH_3 into the ethereal solution. Decomposes readily into acetone and acetamide.— Ac_2CHNa Formed by dissolving sodium in the diketone (Combes, *C R* 104, 920). White six-sided prisms, insol ether. Decomposed by water into acetone and $NaOAc$. With ethyl iodide at 140° it gives $CMEt(CO\ CH_3)_2$, a liquid boiling at 171°. Amyl iodide, in like manner, yields $C_5H_{11}CH(CO\ CH_3)_2$. The second atom of hydrogen in the methylene group may, in such compounds, be displaced by Na , and by acting with an alkyl iodide $R'I$ upon $RCN_2(CO\ CH_3)_2$, we may obtain compounds of the form $RCR'(CO\ CH_3)_2$. These reactions take place with hardly any secondary decompositions. These homologues of methylene, dimethyl diketone are decomposed by potash like the diketone itself (Combes, *A Ch* [6] 12, 211). $ClCO_2Et$ acting on the sodium derivative of methylene dimethyl diketone forms $C(OMe)_2(CO_2Et)_2$ (Claisen a Zedel, *B* 21, 3397)— $KCHAc$, White six sided prisms, sl sol alcohol, insol ether. Formed by adding $KOEt$ to an alcoholic solution of the diketone. Decomposed by hot water into acetone and $KOAc$ — $Mg(CHAc)_2$ (dried at 125°). From the diketone and magnesium carbonate (Combes, *C R* 105, 868). Transparent six sided prisms— $Al(CHAc)_3$. A by product in the rectification of the diketone, from which it may be obtained by treatment with $AlCl_3$. Small red crystals, insol water, sl sol alcohol, v sl sol ether. Can be partially volatilised. Acts on polarised light. Not decomposed by alcoholic NH_3 — $Cu(HC_2Ac)_2$ (dried at 125°). Pale blue needles, obtained by adding cupric acetate or chloride to an aqueous solution of the diketone. Insol water, the pptn being complete in dilute solutions. At 65° it forms with $COCl_2$ dissolved in benzene a crystalline compound melting at 121° (Thomas a Lefèvre, *Bl* [2] 50, 193)— $Fe(HC_2Ac)_3$. Red crystals, deposited from the ethereal extract of the red solution obtained by adding $FeCl_3$ to the diketone dissolved in water— $Pb(HC_2Ac)_2$. From the diketone and lead carbonate. Transparent crystals, sol water.

Oxim $CH_3CO\ CH_2C(NOH)CH_3$, Anhydride $CH<\overset{CMe}{O}>CMeN$ (142°) SG μ 985

Formed by the action of hydroxylamine on the diketone (Zedel, *B* 21, 2178). Colourless oil, with peculiar alkaloidal odour.

Oxim $CH_2C(NOH)CH_3$, [150°] Large transparent crystals (from ether)

Phenyl methyl hydrazide $CH_3CO\ CH_2C(N_2Ph)Me$ (Kohlrausch, *A* 253, 22)

Di-methylene di-methyl triketone $(CH_3CO\ CH_2)_2CO$ *Di-acetyl acetone* [49°] Formed from its anhydride (*vide infra*). Plates. Sl sol water, sol alkalis, warm alcohol, and ether. $FeCl_3$ gives a deep-red colour. Converted by NH_3 into oxy-di methyl pyridine [225°]

Anhydride $CO<\overset{CH\ CMe}{O}>O$ *Di methyl pyrone* [182°] (249°) Formed by the action of $HIAq$ on dehydracetic acid at a high temperature (Fest, *B* 22, 1570, *A* 257, 253). The yield is 70 p.c. of the theoretical. Crystals, v sol water. Converted by baryta, followed by HCl , into di-methylene di-methyl triketone. The carboxylic

acid of this anhydride is dehydracetic acid The dicarboxylic ether $\text{CO} \begin{smallmatrix} \diagup \text{C}(\text{CO}_2\text{Et}) \text{CMe} \\ \diagdown \text{C}(\text{CO}_2\text{Et}) \text{CMe} \end{smallmatrix} \text{O}$ [80°]

S 8 at 20° is formed by the action of COCl_2 on copper acetoacetic ether, and is converted by P_2S_5 into $\text{CS} \begin{smallmatrix} \diagup \text{C}(\text{CO}_2\text{Et}) \text{CMe} \\ \diagdown \text{C}(\text{CO}_2\text{Et}) \text{CMe} \end{smallmatrix} \text{O}$ [110°] (Conrad, *B* 19, 22, 20, 152, 2111).

Di phenyl hydrazide

$(\text{CH}_3)_2\text{C}(\text{N} \text{HPh}) \text{CH}_2 \text{CO}$ [142°]

Trimethylene methyl ketone v TRIMETHYL METHYL KETONE.

Tetramethylene methyl ketone v TETRAMETHYL METHYL KETONE

METHYLENE DIMETHYL ETHER v

Methyl ether of formic orthaldehyde

METHYLENE DIMETHYL DIOXIDE v

Methyl ether of formic orthaldehyde, vol 11 p 570

METHYLENE METHYL PHENYL DIKETONE v BENZOYL ACETONE

METHYLENE METHYL-PHTHALIMIDINE

$\text{C}_{10}\text{H}_8\text{ON}$ & $\text{C}_6\text{H}_5 \begin{smallmatrix} \diagup \text{C}=\text{CH}_2 \\ \diagdown >\text{NMe} \\ \diagup \text{CO} \end{smallmatrix}$ Formed by heat

ing phthal methyl imidyl-acetic acid

$\text{C}_6\text{H}_5 \begin{smallmatrix} \diagup \text{CC}=\text{H CO}_2\text{H} \\ \diagdown >\text{NMe} \\ \diagup \text{CO} \end{smallmatrix}$ above 210° Colourless

crystals Volatile with steam V sol alcohol, ether, and chloroform, more sparingly sol water Very unstable body (Gabriel, *B* 18, 2454)

METHYLENE METHYL PROPYL DIKETONE $\text{C}_8\text{H}_{12}\text{O}_2$ & $\text{CH}_3\text{COCHCOCH}_3$, (175°) S G 12 9411 Formed by the action of EtOAc and NaOAc upon methyl propyl ketone (Claisen & Ehrhardt, *B* 22, 1015) Formed also from butyric ether, acetone, and NaOEt Colourless oil Boiling alkalis yield acetic and butyric acids Yields $\text{C}_6\text{H}_5\text{N}$, CHAcCOPr [55°] crystallising in yellow prisms $-\text{Cu}(\text{C}_6\text{H}_5\text{O})_2$ [161°] Blue needles

TETRA METHYL TRIMETHYLENE DI-

PYRROLE $\text{C}_{11}\text{H}_{12}\text{N}_2$ & $\begin{smallmatrix} \diagup \text{CH CMe} \\ | \\ \diagdown \text{CH CMe} \end{smallmatrix} \text{N CH}_2 \end{smallmatrix} \text{CH}_2$

[77°] Formed by heating acetyl acetone with alcoholic tri methylene diamine at 120° (Paal & Schneider, *B* 19, 3157) Crystalline

METHYLENE DI METHYL DISULPHONE

$(\text{CH}_3\text{SO})_2\text{CH}_2$ [141°] Formed from methyl mercaptan and methylene chloride, and oxidation of the product (Baumann, *B* 23, 1875) Plates On treatment with bromine water it gives $\text{CBr}_2(\text{SO}_2\text{Me})_2$ [234°]

Di methylene di methyl trisulphone

$(\text{CH}_3\text{SO}_2\text{CH}_2)_2\text{SO}_2$ [185°] Formed by saturating a solution of formic aldehyde with H_2S , extracting with ether, evaporating, dissolving the residual oil in aqueous NaOH, adding MeI, and oxidising with KMnO_4 (Baumann, *B* 23, 1872) Prisms, v sl sol cold water, alcohol, and ether Yields with bromine-water insoluble $\text{C}_6\text{H}_5\text{Br}_2\text{S}_3\text{O}_4$ [190°]

TRIMETHYLENE TRINITROSAMINE

$\text{C}_6\text{H}_5\text{N}_3\text{O}_2$ & $(\text{CH}_3\text{N NO})_3$ [106°] Formed by the action of nitrous acid upon hexamethyleneamine (F Mayer, *B* 21, 2888) Yellow needles or prisms (from alcohol), v. sol. alcohol, insol.

petroleum ether Decomposed by water with production of formic aldehyde

METHYLENE DI-OCTYL OXIDE v *Octyl ether of formic orthaldehyde*

METHYLENE OXIDE is FORMIC ALDEHYDE (q v)

METHYLENE-DI-OXY compounds v *Methylene derivatives of Di oxy compounds*

METHYLENE DI PHENYL DIAMINE v

Di PHENYL METHYLENE DIAMINE

METHYLENE DI PHENYLENE v *Di PHENYLENE METHANE*

METHYLENE DIPHENYLENE OXIDE v *Di PHENYLENE METHANE OXIDE*

METHYLENE DIPHENYL DIKETONE v

Di PHENYL METHYLENE DIKETONE

TRIMETHYLENE PHENYL KETONE v

PHENYL TRIMETHYLENE KETONE

Tetramethylene phenyl ketone v *Anhydride of PHENYL OXYBUTYL KETONE*

METHYLENE DIPHENYL OXIDE v *Di PHENYLENE METHANE OXIDE*

METHYLENE HEXAPHENYL PHOS-

PHONIUM IODIDE v *Methylene di iodide of TRI-PHENYL PHOSPHINE*

TRIMETHYLENE DI PHTHALAMIC ACID

$\text{C}_{12}\text{H}_{10}\text{N}_2\text{O}_4$ & $\text{CH}_2(\text{CH}_2\text{NH CO C}_6\text{H}_4\text{CO H})_2$ [70°-123°] Obtained by boiling trimethylene di phthalimide with potash (Gabriel, *B* 21, 2670) Crystalline Decomposed by water On boiling with aqueous HCl it is converted into phthalic acid and trimethylene-diamine

METHYLENE PHTHALIDE $\text{C}_8\text{H}_6\text{O}_2$ & a.

$\text{C}_6\text{H}_5 \begin{smallmatrix} \diagup \text{C}=\text{CH}_2 \\ \diagdown \text{C}=\text{O} \\ \diagup \text{C}=\text{O} \end{smallmatrix}$ *Anhydride of o Oxy vinyl-*

benzoic acid [60°] Formed by heating phthalyl acetic acid *in vacuo* (Gabriel, *B* 17, 2521) Small glistening crystals Soluble in hot water, easily in alcohol, ether, benzene, &c Volatile with steam It readily polymerises Combines with Br (1 mol)

Di bromide $\text{C}_6\text{H}_4 \begin{smallmatrix} \diagup \text{CBr CH}_2\text{Br} \\ \diagdown \text{C}=\text{O} \end{smallmatrix}$ [99°] Thick

glistening crystals Formed by the combination of methylene phthalide with bromine On warming with aqueous KOH it yields aceto phenone o carboxylic acid $\text{C}_6\text{H}_4(\text{CO}_2\text{H})\text{CO CH}_3$ Boiled with water it gives methylene phthalide oxide $\text{C}_8\text{H}_6\text{O}_3$ (Gabriel, *B* 17, 2524)

Methylene-phthalide oxide $\text{C}_8\text{H}_6\text{O}_3$ [146°] Long needles Formed by boiling the dibromide of methylene phthalide with water Formed also by the action of water on the product of the bromination of acetophenone o carboxylic acid (Gabriel, *B* 17, 2524)

Di chloro-methylene phthalide $\text{C}_8\text{H}_4\text{Cl}_2\text{O}_2$ & a.

$\text{C}_6\text{H}_4 \begin{smallmatrix} \diagup \text{C}=\text{CCl}_2 \\ \diagdown \text{C}=\text{O} \end{smallmatrix}$ [128°] Formed by passing

chlorine into a mixture of a chloro phenyl-acetic acid (1 pt) and HOAc (10 pts) Formed also by warming di chloro acetophenone carboxylic acid with conc H_2SO_4 (Zincke & Cooksey, *A* 255, 388) Long slender needles, gradually becoming compact when left in the liquid The needles are v sol hot alcohol, benzene, HOAc, and benzoline, the compact crystals are soluble with difficulty Alcoholic potash

converts it into di chloro acetophenone *o*-carboxylic acid Forms a dichloride

$C_6H_4 \begin{matrix} \diagup CCl_2 \\ \diagdown CCl_2 \end{matrix} \begin{matrix} \diagup O \\ \diagdown O \end{matrix}$ which crystallises in colourless prisms [94°], v sol alcohol and HOAc, and converted by alcoholic potash into phthalic acid

Bromo methylene phthalide v vol 1 p 580

Isomeride of methylene phthalide v METHYLENE PHTHALYL

Polymeride of methylene phthalide

$C_{18}H_{12}O_4$ [216°] Obtained by allowing a solution of acetophenone carboxylic acid (1 pt) in H_2SO_4 (2 pts) to stand in the cold, and then ppg with water (Roser, *B* 17, 2620, Gabriel, *B* 17, 2666, cf ACETOPHENONE *o* CARBOXYLIC ACID, reaction 2) Plates Insol water and cold alkalis, sl sol alcohol, v sol HOAc

Oxym $C_{18}H_{12}NO_4$ [180°] Obtained by heating the substance with alcoholic hydroxylamine hydrochloride at 160° Crystalline granules (from dilute HOAc)

METHYLENE-DI-PHTHALIMIDE

$CH_2(N \begin{matrix} \diagup CO \\ \diagdown CO \end{matrix} C_6H_4)_2$ [226°] Formed by heating potassium phthalimide (2 mols) with methylene iodide (1 mol) at 175° (Neumann, *B* 23, 1002) Light brown crystals (from HOAc), insol dilute alkalis Yields phthalic acid on heating with $HClAq$

TRIMETHYLENE-DI-PHTHALIMIDE

$C_{18}H_{12}N_2O_4$ i.e. C_6H_4O , N , CH , CH , CH_2 , N , C_6H_4O , [198°] Formed by the action of tri methylene bromide on potassium phthalimide (Gabriel, *B* 21, 2669) White needles, m sol hot HOAc, sl sol ether, CS_2 , chloroform, and cold alcohol, v sl sol water and petroleum ether Converted by heating with KOH into tri methylene di phthalamic acid $C_{18}H_{12}(NHCOC_6H_4CO_2H)_2$ De composed by heating with HCl at 180° into phthalic acid and tri methylene diamine

METHYLENE-PHTHALYL C_6H_4O , i.e.

$C_6H_4 \begin{matrix} \diagup CO \\ \diagdown CO \end{matrix} CH_2$? [219°] Fine yellow needles

Formed, together with *o* tribenzoylene benzene, by heating phthalic anhydride with malonic ether and sodium acetate (Gabriel, *B* 14, 925)

METHYLENE-DI-PIPERIDINE

$CH_2(NC_4H_8)_2$ (230°) Formed by distilling piperidine with trimethylene oxide (formic par aldehyde) (Ehrenberg, *J pr* [2] 36, 126) Liquid, with pepper like smell Is decomposed by dilute acids into its constituents CS_2 forms an addition product $B''CS_2$, [58°], sol alcohol and ether, insol water

METHYLENE-TETRA-PROPYL-DIAMINE

$CH_2(NPr)_2$ (220°) Formed by distilling di-propylamine with trimethylene oxide (formic paraldehyde) (Ehrenberg, *J pr* [2] 36, 122) Liquid, sl sol water, v sol alcohol, ether, and chloroform

METHYLENE DI-PROPYL OXIDE v Di-propyl ether of FORMIC ORTHALDEHYDE

TRI-METHYLENE SELENO-UREA v SELENIUM COMPOUNDS, ORGANIC

METHYLENE SULPHIDE v. THIOFORMIC ALDEHYDE.

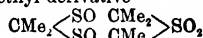
Tri methylene tetrasulphide $C_3H_2S_4$, i.e.

$CH_2 \begin{matrix} \diagup S CH_2 S \\ \diagdown S CH_2 S \end{matrix}$ [84°] Formed by the action of H_2S on formic aldehyde (Baumann, *B* 23, 1869) Colourless needles, insol water, m sol alcohol and ether, v e sol $CHCl_3$ and benzene

METHYLENE SULPHOCYANIDE $C_3H_2S_2N_2$, i.e. $CH_2(SCy)_2$, [102°] Formed by digesting potassium sulphocyanide (2 mols) with methylene iodide (1 mol) in alcoholic solution for 2 or 3 hours on the water bath, ppg with water, and recrystallising from alcohol (Lermontoff, *B* 7, 1282) Crystals, v sol alcohol and ether, m sol hot, nearly insol cold, water Oxidised by conc HNO_3 to methane disulphonic acid

TRI-METHYLENE-TRISULPHONE

$C_3H_2S_3O_6$, i.e. $CH_2 \begin{matrix} \diagup SO_2 CH_2 \\ \diagdown SO_2 CH_2 \end{matrix} SO_2$ Formed by oxidising thioformic paraldehyde (trithio form aldehyde) with $KMnO_4$ (E Baumann a R Camps, *B* 23, 69) Crystalline powder, insol water, dilute acids, alcohol, ether, chloroform, and glacial acetic acid, v sol cold caustic soda, ammonia (by warming), and sodic carbonate Expels CO from $Na CO_3$ on warming Conc HNO_3 and H_2SO_4 have no action even on warming Its solution in H_2SO_4 is ppd by water unaltered It partially sublims The six hydrogen atoms can be displaced by alkyl groups The hexamethyl derivative



is identical with the product obtained by B Jaffé, E Baumann, and Fromm (*B* 22, 2598, 2609) by oxidising trithio acetone

TRIMETHYLENE-DI-SULPHONIC ACID so called v PROPANE DI SULPHONIC ACID

METHYLENE TRI-THIO-CARBONATE

$CH_2 \begin{matrix} \diagup S \\ \diagdown S \end{matrix} CS$ Separates on gently heating $CS(SNa)_2$ with methylene iodide in alcoholic solution (Husemann, *A* 126, 292) Amorphous yellowish white powder, insol water Converted by fuming nitric acid into methane di sulphonic acid

TRIMETHYLENE THIO UREA $C_3H_4N_2S$ i.e.

$CH \begin{matrix} \diagup CH_2 NH \\ \diagdown CH_2 NH \end{matrix} CS$ [198°] Formed, together with ammonium sulphocyanide, by heating the sulphocyanide of trimethylene diamine $C_3H_4(NH_2)_2(HNCS)$, (Lellmann a Wurthner, *A* 228, 232) White needles (from chloroform mixed with light petroleum) Sol water, alcohol, $CHCl_3$, and benzene, m sol aqueous NaOH, insol light petroleum Forms a sparingly soluble compound with $HgCl_2$

Trimethylene-ψ thio-urea

$CH_2 \begin{matrix} \diagup CH_2 S \\ \diagdown CH_2 NH \end{matrix} C NH$ Formed by evaporating a solution of γ bromo propyl aminohydrobromide and potassium sulphocyanide to dryness at 100° (Gabriel a Lauer, *B* 23, 94) Liquid, v sol water forming an alkaline solution, from which it can be extracted by benzene -B'HBr [136°] -B'C₆H₅(NO₂)OH [128°] Long needles

TRIMETHYLENE-UREA $C_3H_4N_3O$ i.e.

$CH_2 \begin{matrix} \diagup CH_2 NH \\ \diagdown CH_2 NH \end{matrix} CO$ *Oxy-pyrimidine tetrahy dride* [260°] Formed by heating trimethylene diamine (1 mol) with carbonic ether (1 mol) for 6 hours at 180° (Fischer a Koch, *A* 232, 224).

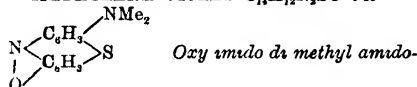
White needles, ν sol water, sl sol alcohol and ether. Its solution is neutral, and is not pptd by HNO_3 or oxalic acid. By chromic acid mixture it is oxidised to a compound $\text{C}_6\text{H}_4\text{N}_2\text{O}_2$ crystallising in plates [275°]

Trimethylene ψ -urea $\text{CH}_2\langle\text{CH}_2\text{O}\rangle\text{C NH}$

or $\text{CH}_2\langle\text{CH}_2\text{O}\rangle\text{C NH}_2$. Formed by evaporating a solution of equivalent quantities of potassium cyanate and γ -bromo propyl aminehydrobromide (Gabriel & Lauer, *B* 23, 95). Thick liquid, ν sol water— $\text{B}'\text{C}_6\text{H}_5(\text{NO}_2)_2\text{OH}$ [200°]. Long yellow needles.

Trimethylene-di-urea $\text{C}_6\text{H}_4\text{N}_2\text{O}_2$ ν e $\text{CH}_2(\text{CH}_2\text{NH CO NH}_2)_2$ [182°]. Formed by warming a dilute aqueous solution of trimethylene diamine hydrochloride with silver cyanate (F & K). White needles, ν sol water, sl sol alcohol, insol ether.

METHYLENE VIOLET $\text{C}_{11}\text{H}_{12}\text{N}_2\text{SO}$ ν e.



diphenylene sulphide. Oxy dimethylamido thio diphenylamide. Formed by boiling a solution of methylene blue (base). Formed also by oxidising a mixture of di methyl di amido phenyl, mercaptan, and phenol (Bernthsen, *A* 230, 171, 251, 96). Needles, ν sl sol water, sl sol alcohol, ether acetone, chloroform, benzene, ligroin, and cumene, with a reddish brown fluorescence, ν sol aniline, without fluorescence— $\text{B}'\text{HCl}$ ν sl sol cold dilute HClAq .

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DI METHYL ENNYLENE DIKETONE

$\text{C}_{11}\text{H}_{16}\text{O}_2$ ν e $\text{CH}(\text{CH}_3\text{CHEt CO CH}_3)_2$. *Di acetyl di ethyl n pentane* (208° at 110 mm). Formed from its dicarboxylic ether by treatment with alcoholic potash (Kipping & Perkin, jun, *C J* 57, 33). Colourless oil, with slight aromatic odour, ν sl sol water, miscible with alcohol and ether. It dissolves without change in conc H_2SO_4 , forming a yellowish brown solution. It does not combine with NaHSO_3 .

Orim $\text{CH}(\text{CH}_3\text{CHEt CMe NOH})_2$ [111°]. Obtained by heating the ketone with an alcoholic solution of hydroxylamine, evaporating, adding water, and extracting with ether. Minute colourless crystals (from benzene ligroin), ν sol alcohol, ether, HOAc , benzene, and aqueous alkalis.

DI-METHYL ENNYLENE DIKETONE DI-CARBOXYLIC ETHER

$\text{CH}_2(\text{CH}_2\text{Cet}(\text{CO CH}_3)_2\text{CO}_2\text{Et})_2$ ν e *Di acetyl ac di ethyl pmetic ether* [45°]. Obtained, together with compounds of low boiling point, when sodium ethyl acetoacetic ether is heated with an alcoholic solution of trimethylene bromide on a water bath (Kipping & Perkin, jun, *C J* 57, 81). Very slender needles (from ether alcohol), ν sol ether and alcohol, ν e sol benzene, light petroleum, xylene, and chloroform, insol cold water. Gives no colour with FeCl_3 . Readily decomposed by warm alcoholic potash.

METHYL ENNYL KETONE $\text{CH}_2\text{CO C}_6\text{H}_5$, [16°] (224°) SG 17.5 8295. The chief constituent of oil of rue, obtained by distilling *Ruta*

graveolens with steam (Greville Williams, *T* 1858 [1] 99, Hallwachs, *A* 113, 109, Harbordt, *A* 123, 293, Giesecke, *Z* [2] 6, 429, cf Gerhardt *C R* 26, 225, 361, Cahours, *C R* 26, 262). Occurs in the essential oil obtained from lime leaves (*citrus Limetta*) (F Watts, *C J* 49, 316). Formed also by distilling a mixture of calcium acetate and calcium decanoate (Gorup Besanez & Grimm, *A* 157, 275, B 3, 518), and by boiling octyl acetoacetic ether with alcoholic potash (Guthzeit, *A* 204, 4). It is oxidised by chromic acid mixture to acetic and ennoic acids. Sodium amalgam reduces it, in alcoholic solution, to *sec* hendecyl alcohol (229°), SG 12 826. It combines with NH_3 . With alkali bisulphites it forms compounds such as $\text{C}_6\text{H}_5\text{CMe}(\text{OH})\text{SO}_2\text{NH}_2\text{aq}$, which crystallises in pearly plates. PCl_5 converts the ketone into $\text{C}_{11}\text{H}_{22}\text{Cl}_2$, which is resolved by distillation into HCl and $\text{C}_{11}\text{H}_{22}\text{Cl}$ (222°).

Orim $\text{C}_6\text{H}_5\text{CMe NOH}$ [42°]. Minute prisms (from alcohol), ν sol ether (Spiegler, *M* 5, 242, B 17, 1575).

METHYL ENNYL KETONE CARBOXYLIC

ACID $\text{CH}_2\text{CO CHEt CH}_2\text{CH}_2\text{CH}_2\text{CHEt CO}_2\text{H}$ ν e *Acetyl-ac di ethyl-caproic acid* (254° at 90 mm). Formed, together with di methyl ennylene diketone and acetic acid, by boiling di ethyl ennylene diketone dicarboxylic acid with alcoholic potash (Kipping & Perkin, *C J* 57, 36). Thick oil, miscible with alcohol, ether, and benzene— AgA' . Amorphous pp, m sol hot water.

Orim $\text{CH}_2\text{C}(\text{NOH})\text{C}_6\text{H}_5\text{CO}_2\text{H}$ [103°]. Minute plates (from benzene ligroin), ν sol alcohol and benzene, sl sol ligroin, sol alkalis and conc HClAq .

TETRAMETHYLENYL-CARBINYL-AMINE

$\text{C}_6\text{H}_{11}\text{N}$ ν e $\text{CH}_2\langle\text{CH}_2\text{CH}_2\text{NH}\rangle_2$ 'Tetra methylenylamine' (83°). Formed from the nitrile of tetramethylene carboxylic acid $\text{CH}_2\langle\text{CH}_2\text{CH}_2\text{CN}\rangle_2$ by reduction in alcoholic solution with sodium (Freund & Gudeman, *B* 21, 2692). Oil, with alkaline reaction, which absorbs CO_2 from the air— $\text{B}'\text{HCl}$ [236°]. Crystallises from alcohol ether, ν sol water and alcohol, insol ether— $\text{B}'_2\text{H}_4\text{PtCl}_4$. Crystalline.

TETRAMETHYLENYLCARBINYL-THIO-UREA $\text{C}_6\text{H}_{11}\text{N}_2\text{S}$ ν e

$\text{CH}_2\langle\text{CH}_2\text{CH}_2\text{NH CS NH}_2\rangle_2$ [68°]. Prepared by the action of ammonium sulphocyanide upon the hydrochloride of tetramethylenyl-carbinyllamine (Freund, *B* 21, 2697). Slender needles (from water or alcohol).

TETRAMETHYLENYLCARBINYL UREA

$\text{C}_6\text{H}_{11}\text{N}_2\text{O}$ ν e $\text{CH}_2\langle\text{CH}_2\text{CH}_2\text{NH CO NH}_2\rangle_2$ [116°]. Obtained by evaporating a solution of tetramethylenylcarbinyllaminehydrochloride with potassium cyanate (Freund, *B* 21, 2697). Needles, sl alcohol, ν sol hot, m sol cold water.

DI-TETRAMETHYLENYL KETONE $\text{C}_6\text{H}_4\text{O}$ ν e $\text{CH}_2\langle\text{CH}_2\text{CH}_2\text{CO CH}_2\text{CH}_2\text{CO}\rangle_2$ (205°)

A product of the distillation of calcium tetramethylene carboxylate with lime (Colman & Perkin, jun, *C J* 61, 235). Oil, smelling of

KOH on the water bath White plates, sol boiling water, v sl sol cold water, v e sol alcohol and ether Forms the salts —(B'MeCl crystallising in rhombohedra, v e sol water — B'Me₂PtCl₂ — Picrate [112°] crystallising in needles, m sol boiling water and alcohol

METHYL ETHER v Di METHYL OXIDE

PENTA METHYL ETHOL A name some times employed to denote the alcohol

CM₅, CM₅OH v HPTYL ALCOHOL

METHYL ETHYL ACETAL v ALDEHYDE

METHYL ETHYL ACETIC ACID v VALERIC

ACID

Di methyl ethyl acetic acid v HEXOIC ACID

METHYL ETHYL ACETOACETIC ETHER

v ACETOACETIC ACID

METHYL ETHYL ACETOXIM v, Oxim of

METHYL ETHYL KPTONF

METHYL ETHYL ACETOXIMIC ACID v

DI ACETYL

METHYL ETHYL ACETYLENE v PENTIN

ENE

METHYL ETHYL ACROLEIN v HFXFNOIC

ALDHYDE

METHYL ETHYL ACRYLIC ACID v HEX

ENOIC ACID

PENTA METHYL ETHYL ALCOHOL v

Tert HEPTYL ALCOHOL

DI METHYL ETHYL ALKINE v Di

METHYL OXY ETHYL AMINE

METHYL ETHYL-ALLYL ALCOHOL v

HEXFNYL ALCOHOL

METHYL DI-ETHYL AMINE C₅H₁₁N 2 e MeNkt, Formed by distilling its methyl chloride, or by distilling triethylamine methyl hydroxide NEt₃Me(OH) (V Meyer a Lecco, A 180, 184, Lossen, A 181, 379) — B'H.PtCl₂. Monoclinic crystals (Hjortdahl, J 1892, 476)

Methyl chloride C₂H₅Cl + Me₂NEt₂Cl Di methyl di ethyl ammonium chloride Obtained from the crystalline iodide Me₂NEt₂I, which is formed by heating dimethylamine with EtI or diethylamine with MeI (Petersen, A 91, 122, V Meyer a Lecco, A 180, 177) — (Me₂NEt₂Cl)₂PtCl₂, yellowish dimetric prisms, m sol water, sl sol alcohol and ether S 1 025 at 15° — (Me₂NEt₂Cl)₂HgCl₂, trimetric crystals (Topsoe, J 1893, 620) — Me₂NEt₂ClHgCl₂ — Me₂NEt₂Cl₂HgCl₂, trimetric crystals — Me₂NEt₂ClAuCl₂, dimetric crystals

Methyl picrate [287°] Needles (Lossen, A 181, 374)

Ethyl hydroxide v Tri ETHYLAMINE methyl hydroxide, vol 11 p 476

Di methyl ethyl amine NMe₂Et (41°) A product of the action of heat on trimethylamine ethylchloride (Collie a Schryver, C J 57, 770) Formed also by distilling NMe₂Et OH

METHYL TRI ETHYL AMMONIUM COMPOUNDS v *Methyl hydroxide* of Tri ETHYL-AMINE

Di methyl di ethyl ammonium compounds v *Methyl chloride* of METHYL DI ETHYL AMINE

Tri methyl-ethyl ammonium compounds v *Ethyl chloride* of Tri METHYL AMINE

METHYL ETHYL ISOAMYL-AMINE

C₅H₁₁N 2 e MeNEtC₅H₁₁ (135°) Formed, together with ethylene, by the dry distillation of methyl di ethyl isoamyl ammonium hydroxide MeNEt₂(C₅H₁₁)(OH), which is obtained by the action of moist Ag₂O on the product of the

union of MeI with di ethyl isoamyl amine (Hofmann, C J 4, 317) Fragrant liquid, sl sol water — B'H₂PtCl₂, orange yellow needles, v sol water

METHYL ETHYL-AMYL-PHENYL-AMMONIUM HYDROXIDE v *Methyl hydroxide* of ETHYL ISOAMYL ANILINE, vol 11 p 476

METHYL-ETHYL-ANILINE C₆H₅N 2 e C₆H₅NMeEt (201° uncor)

Formation — 1 By methylation of ethyl aniline (Hofmann, A 74, 152) — 2 By ethylation of methyl aniline (Claus a Howitz, B 17, 1325) — 3 From di ethyl aniline methyl iodide

Properties — Crystalline, forming extremely soluble salts The hydrochloride melts at 114° (Claus a Hirzel, B 19, 2785)

Methyl iodide B'MeI [125°], identical with di methyl aniline methyl iodide (Claus a Rautenberg, B 14, 620, Hjortdahl, J 1882, 510) Triclinic crystals, v sol water and alcohol De composed by boiling with conc KOHAq, yielding dimethylaniline — B'HCl [114°], very hygroscopic crystals — (B'MeI)₂ZnI₂, monoclinic crystals — (B'Me)₂H₂Fe₂Cy₁₂2aq (Fischer, A 190, 187)

Ethyl iodide B'EtI [102°], identical with di ethyl aniline methyl iodide, by treatment with KOH it gives methyl ethyl aniline

Propyl iodide C₃H₇NMeEtPrI *Methyl propyl aniline ethyl iodide*, ethyl propyl aniline methyl iodide Thick syrup, v sol water Formed by the combination of methyl ethyl aniline with propyl iodide, of ethyl propyl aniline with methyl iodide, or methyl propyl aniline with ethyl iodide By boiling with aqueous KOH the propyl group is split off and methyl-ethyl aniline regenerated (Claus a Hirzel, B 19, 2785)

Reference — Bromo METHYL ETHYL ANILINE **METHYL ETHYL ARSINE** v ARSENIC COMPOUNDS, ORGANIC

DIMETHYL ETHYL AZONIUM CHLORIDE

C₅H₁₁N₂Cl 2 e Me₂NEtClNH₂ Crystalline compound E sol water Prepared by the combination of ethyl chloride with dimethyl hydrazine On reduction with zinc dust and acetic acid it gives dimethyl ethylamine, NH₂, and HCl (Renouf, B 13, 2172) — (C₅H₁₁N₂Cl)₂PtCl₂, crystalline

METHYL ETHYL BENZENE v ETHYL-TOLUENE

Di methyl-ethyl-benzene v ETHYL-XYLENE Laurene C₁₁H₁₆ (LauroI) (188°) S G, 12 887 According to Armstrong a Miller (B 16, 2258) this liquid is di methyl ethyl benzene C₆H₅Me₂Et [1 2 4] It is one of the products of the action of ZnCl₂ on camphor (Fittig, A 145, 149) It yields di methyl benzoic acid on oxidation Montgolfer (A Ch [5] 14, 91) described laurene as C₁₁H₁₆ (195°) Reuter (B 16, 627) described two laurenes (a) laurene (190°) yielding C₆H₅Me₂CO₂H [1 4 2] on oxidation by dilute HNO₃, and (b) laurene (185°) yielding on oxidation C₆H₅Me₂CO₂H [1 3 4] and forming a sulphonic acid whose Ba salt is more soluble than that of (a) laurene According to Uhlhorn (B 23, 2346) laurene contains (1,2,4)- and (1,4,2)-di-methyl ethyl-benzenes

o **METHYL-ETHYL BENZENE HEXAHYDRIDE** C₆H₁₀ 2 e CH₂ < CH₂CHMe > CH₂Et

Methyl-ethyl hexamethylene (151°) Formed by boiling $\text{CH}_2 < \begin{smallmatrix} \text{CH}_2 \text{CHMe} \\ \text{CH}_2 \text{CH}_2 \end{smallmatrix} > \text{CH CH(OH) CH}_2$ with fuming HIAq , diluting with water, extracting with ether, and heating the resulting oily $\text{CH}_2 < \begin{smallmatrix} \text{CH}_2 \text{CHMe} \\ \text{CH}_2 \text{CH}_2 \end{smallmatrix} > \text{CH CHI CH}_2$, (179° at 110 mm) with excess of HIAq (S G 1 96) and amorphous phosphorus for 8 hours at 235° Kipping a Perkin, jun, C J 57, 26) Mobile oil with odour of paraffin, miscible with alcohol and ether

METHYL ETHYL-BENZENE (8) **SULPHONIC ACID** $\text{C}_6\text{H}_4\text{MeEt SO}_3\text{H}$ [1 2 47] Formed by sulphonating *o*-ethyl-toluene (Claus a Pieszek, B 19, 3087) Crystalline — NaA' aq plates, *v* sol water, insol alcohol — KA' aq plates, *v* sol water — CaA' , 2aq plates grouped in nodules, *v* e sol water — BaA' , 3aq plates, *v* e sol water — PbA' , 8aq plates, *v* e sol water — CuA' aq blue plates, *v* e sol water
Chloride $\text{C}_6\text{H}_4\text{MeEt SO}_2\text{Cl}$ Oil
Amide $\text{C}_6\text{H}_4\text{MeEt SO}_2\text{NH}_2$ Yellowish brown oil, *v* sl sol cold water, sol hot water
m Methyl ethyl-benzene sulphonic acid *v* vol 11 p 524

(1, 2, 4) **Di-methyl-ethyl-benzene sulphonic acid** $\text{C}_6\text{H}_2\text{Me}_2\text{Et SO}_3\text{H}$ Formed by sulphonating ethyl-*o*-xylene (O Jacobsen, B 19, 2516, Stahl, B 23, 991) Large tables — BaA' , 4aq — BaA' , 3aq white plates, m sol hot or cold water (S) — NaA' , 1½aq Small white plates, *v* sol water

Amide $\text{C}_6\text{H}_2\text{Me}_2\text{Et SO}_2\text{NH}_2$ [126°] Long slender needles (from warm, very dilute, alcohol), or large prisms (from alcohol)

Di-methyl-ethyl-benzene sulphonic acid $\text{C}_6\text{H}_2\text{Me}_2\text{Et SO}_3\text{H}$ **Laurene sulphonic acid** Formed by sulphonating laurene According to Reuter it is accompanied by an amorphous isomeric which forms a gummy Ba salt Triclinic crystals (Reuter, B 16, 627) When steam is passed into its solution in dilute H_2SO_4 , hydrolysis begins when, through concentration, the temperature has reached 120° (Armstrong a Miller, C J 45, 148) — BaA' , 3aq S (of BaA') 2 5 at 16°

Amide $\text{C}_6\text{H}_2\text{Me}_2\text{Et SO}_2\text{NH}_2$ [127°] (R)

(1, 3, 4) **Di-methyl-ethyl-benzene sulphonic acid** $\text{C}_6\text{H}_2\text{Me}_2\text{Et SO}_3\text{H}$ Formed by sulphonating ethyl *m*-xylene (J) — BaA' , 2aq trimetric laminae — NaA' 2aq prisms

Amide $\text{C}_6\text{H}_2\text{Me}_2\text{Et SO}_2\text{NH}_2$ [148°]

(1, 4, 3) **Di-methyl-ethyl-benzene sulphonic acid** $\text{C}_6\text{H}_2\text{Me}_2\text{Et SO}_3\text{H}$ Formed by sulphonating ethyl *p*-xylene (Jacobsen, B 19, 2516) Large plates (from dilute H_2SO_4) — NaA' aq mass of large tables (from warm saturated solution) — KA' aq flat needles, *v* sol water — BaA' , long six sided plates, sl sol cold, m sol hot, water — CuA' , 8aq light blue needles (Stahl, B 23, 990)

Amide $\text{C}_6\text{H}_2\text{Me}_2\text{Et SO}_2\text{NH}_2$ [117°] Pearly plates (from dilute alcohol), or large transparent crystals (from warm alcohol), *v* sol hot alcohol

METHYL DI ETHYL BORATE MeEt_2BO , (100°–105°) S G 2 904 Formed by heating MeBO , with absolute alcohol at 100°

PENTA METHYL ETHYL BROMIDE *v*.

Tert HEPTYL BROMIDE.

METHYL-ETHYL-BROMANILINE *v* **BROMO-**

METHYL ETHYL ANILINE

METHYL-ETHYL-CARBINOL *v* **Sec BUTYL ALCOHOL**

Methyl-di-ethyl-carbinol *v* **Tert HEXYL ALCOHOL**

Di-methyl-ethyl-carbinol *v* **Tert AMYL ALCOHOL**

METHYL ETHYL CARBONATE MeEtCO_2 , [–14 5°] (109° cor) (Rose, A 205, 230) S G 2 1 002 Obtained by distilling a mixture of potassium methyl carbonate with potassium ethyl sulphate (Chancel, C R 31, 521, A 79, 90) Formed also by the action of NaOMe on ClCO_2Et , or of NaOEt on ClCO_2Me (Schreiner, J pr [2] 22, 354)

PENTA-METHYL-ETHYL CHLORIDE *v* **HEPTYL CHLORIDE**

METHYL ETHYLENE *v* **PROPYLENE**

Di-methyl-ethylene *v* **BUTYLENE**

Tri-methyl-ethylene *v* **AMYLENE**

Tetra-methyl-ethylene *v* **HEXYLENE**

METHYL ETHYLENE DIAMINE **Nitro-derivative** $\text{C}_2\text{H}_4(\text{NHNO}_2)(\text{NMeNO})$ [122°] Formed by the action of methyl iodide on ethylene diamine (Franchimont a Klobbie, R T C 7, 343)

Di-methyl-ethylene-diamine **Nitro-derivative** $\text{C}_2\text{H}_4(\text{NMeNO})_2$ [137°] Formed like the preceding

Di methyl ethylene diamine *v* **ETHYLENE DIAMINE**

METHYL - ETHYLENE - TRI - ETHYL - PHOSPHAMMONIUM BROMIDE *v* vol 11 p 509

DI METHYL-ETHYLENE GLYCOL *v* **DI OXY-BUTANE**

TRI-METHYL-ETHYLENE GLYCOL *v* **DI OXY PENTANE**

DI METHYL ETHYLENE DIKETONE *v* **ACETONYL-ACETONE**

DI-METHYL-ETHYLENE DIKETONE DI CARBOXYLIC ACID *v* **Di aceto succinic ether** described under **ACETYL SUCCINIC ETHER**

TETRA-METHYL-ETHYLENE-DI-PHENYL-DIPHOSPHONIUM BROMIDE *v* **Ethylene-bromide of PHENYL DI METHYL PHOSPHINE**

DI METHYL-ETHYLENE DI PHENYL-DIPYRROLE DICARBOXYLIC ETHER

$\text{C}_6\text{H}_4(\text{C}_2\text{NHMePh CO}_2\text{Et})_2$, [197°] Formed from acetophenone acetoacetic ether and ethylene acetides (Paal a Schneider, B 19, 3156) Scales (containing 4aq) Yields on hydrolysis the corresponding acid [181°]

METHYL-ETHYLENE-PYRIDINE TETRA HYDRIDE $\text{C}_4\text{H}_8\text{N}_4$ *vs* $\text{C}_4\text{H}_8\text{NMe}(\text{C}_2\text{H}_5)_2$ **Tropidine** (162°) S G 2 9665 Formed by heat in atropine or tropine with fuming HClAq and HOAc at 180°, or by heating tropine with H_2SO_4 (Ladenburg, A 217, 117) Liquid, smelling like conine, *v* e sol cold, sl sol hot, water, *v* e sol alcohol and ether Alkaline in reaction Its hydrobromide is decomposed by bromine at 175° yielding di bromo methyl pyridine and ethylene — $\text{B}'\text{H}_2\text{PtCl}_4$, — $\text{B}'\text{HAuCl}_4$, — $\text{B}'\text{HI}$, [98°] Brown prisms — $\text{B}'\text{C}_2\text{H}_5(\text{NO}_2)_2\text{OH}$ yellow needles (from boiling water)

Methyl tropide B'Mel Cubes Decomposed on distillation with KOH yielding tropine $\text{C}_7\text{H}_{10}\text{O}$, an oil $\text{C}_7\text{H}_8\text{O}$ (?) (202°–207°), dimethylamine and methyl tropine With moist Ag_2O it

gives a syrupy base which yields the salts
 $(B'MeCl)_2PtCl_2 - B'MeBr - B'MeC_6H_4(NO_2)_2O$

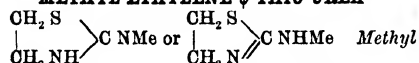
TETRA METHYL, ETHYLENE DI PYR-
ROL $C_{14}H_{20}N_2$ *re*



tained by adding ethylene diamine (1 mol) to a solution of acetylonyl acetone (2 mols) in an equal weight of absolute alcohol. White pearly plates. Sublimable. Volatile with steam. V sol alcohol, ether, benzene, and petroleum spirit, insol water. Colours a chip of pine wood. Carmine red. Gives a purple red colouration with phenanthraquinone and H_2SO_4 . (Paal & Schneider. *B* 19. 3157).

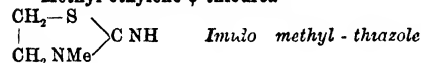
DI METHYL ETHYLENE DISULPHIDE
 $C_2H_4S_2$ *vs* $C_2H_4(SMe)_2$ *Di methyl ether of di thio glycol* (183°) Formed from ethylene bromide and $NaSMe$ (Ewerlof, *B* 4, 716)

DI METHYL ETHYLENE DISULPHONE *v*
Di methyl ether of ETHANE DISULPHINIC ACID
METHYL ETHYLENE \downarrow THIO UREA



Methyl amido
thiazole dihydride [90°] Formed, together
with a crystalline compound $C_6H_{11}N_2O_2$ [70°],
by the action of methyl thiocarbamide on bromo
ethylamine (Gabriel, *B* 22, 1148) Needles, v e
sol water, v sol ordinary solvents Strongly
alkaline Oxidised by bromine water to an acid
(γ -NHMe CO NH, CH $_2$ CO, SO $_2$ H), which is de
composed by fuming HClAq at 155° into CO,
methylamine, and taurine —Picrate [226°]
needles —Aurochloride needles —Platino-
chloride needles

Methyl ethylene ψ thiourea



Formed by heating ethylene ϕ thio urea with methyl iodide (Gabriel). Oil. When it is oxidised by bromine water it yields methyl taurine $\text{NHMeCH}_2\text{CHSO}_3\text{H}$. Salts— B^+H^- [160°]. Colourless crystals, sol water and alcohol. Pirarate [200°–203°]—The aurochloride and platinochloride form long needles, sol boiling water.

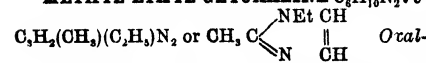
METHYL ETHYL ETHER v METHYL ETHYL
OXIDE

METHYL-ETHYL-ETHYLENE u AMYLENE

METHYL-ETHYL-GLYOXAL 2 METHYL

ETHYL DIKFTONE

METHYL-ETHYL-GLYOXALINE $C_6H_{10}N_2 \cdot 0$



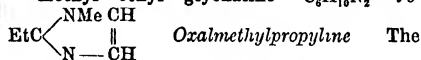
ethyl ethylamine (213°) SG 15 98 Formed by the action of ethyl bromide on methyl glyoxaline (glyoxal ethylamine) (Radziszewski, *B* 16, 489) Formed also from diethyl oxamide NHEt CO CO NHEt by treatment with PCl₅ and heating the hydro iodide (10 g) of the resulting 'chloroethylethylamine' C₄H₁₁ClN, with HIAq (7 g of SG 19) and amorphous phosphorus (1 g) for 6 hours at 140°. The product is dissolved in water, rendered alkaline, and extracted with chloroform (Wallach, *A* 214, 298) Colourless and liquid, with narcotic smell. Sol. water and

alcohol AgNO₃ gives a crystalline pp, HgCl₂, a white pp The zinc double chloride forms crystals melting at [160°] Burns with a blue flame It is a strong base and ppts metallic oxides from their salts Its zinc double salt distilled with lime yields pyrrole, ammonia, HCl, ethylene, and methyl glyoxaline (para oxal methylene) (Wallach, *A* 214, 305) It acts physiologically like atropine (Schulz, *B* 13, 2353) When heated in a sealed tube with dilute H₂SO₄ at 240° it yields ethylamine KMnO₄ yields oxalic acid, NH₃, and acetic acid When passed through a red hot tube it yields HCl and methyl glyoxaline (para oxal methyl ene) H₂O₂ oxidises it to ethyl oxamide (Radzi szewski, *B* 17, 1290)

Salts — $\text{B}'\text{HCl}$ deliquescent crystals — $\text{B}'_2\text{H}_2\text{PtCl}_6$ — $\text{B}'_2\text{H}_2\text{ZnCl}_4$ [160°] — $\text{B}'_2\text{AgNO}_3$.
Methyl iodide $\text{C}_5\text{H}_{10}\text{N}_2\text{MeI}$ Crystals, ν e sol water and alcohol. Not decomposed by aqueous NaOH — $\text{C}_5\text{H}_{10}\text{N}_2\text{MeI}_3$

Benzyl chloride $C_6H_5CH_2Cl$

Methyl - ethyl - glyoxaline $C_6H_{10}N_2$ 2 e



methylo iodide B'MeI of this body is formed by the action of MeI on para ethyl glyoxaline (Radziszewsky, *B* 16, 490)

References — DI BROMO, and CHLORO,
METHYL ETHYL GYOXALINE

METHYL ETHYL GLYOXIM *v Oxim of*
METHYL ETHYL DIKETONE

METHYL ETHYL HYDROXYLAMINE v

METHYL ETHYLIDENE ETHYL DI-KETONE $C_6H_{12}O_2$ *re* $CH_3COCHMeCOCH_3$
Acetyl propionyl ethane (167°-170°) Formed by the action of NaOEt on a mixture of diethyl ketone and acetic ether (Claisen & Ehrhardt, B 22, 1009) Colourless oil— $Cu(C_6H_{11}O_2)_2$, [192°] Blue crystals (from benzene ligroin)

METHYL ETHYLIDENE DI ETHYL
SULPHONE v DI ETHYL PROPYLIDENE DI
SULPHONE

DI METHYL ETHYLDENE DI-INDOLE
 $C_2H_5N_{20}$, τ_e CH_3 , $CH(C_6H_5, MeN)_2$ or
 $(NH < \overset{C_6H_5}{CMe})_2$, $CHMe$ *Ethyldene methyl*
ketole [191*] Formed by heating methyl
indole (methyl ketole), paraldehyde, and $ZnCl_2$
on the water-bath (Fischer, *A* 242, 376).
Colourless prisms, ν sol alcohol, ether, and
acetone

DIMETHYL ETHYLIDENE DIKETONE
(CH₃CO)₂CHMe (165°) Formed from sodium
methylene dimethyl diketone and MeI (Combes,
C R 105, 868)

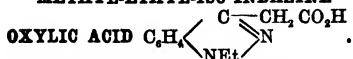
METHYL ETHYL INDAZINE $C_{10}H_{12}N_2$ *ts*
 C_6H_5 $\begin{array}{c} \text{CMe} \\ \diagup \quad \diagdown \\ \text{N} \quad \text{NEt} \end{array}$ Formed by heating methyl-
 indazine with EtI at 100° (Fischer & Tafel, *A*
 227, 303) Liquid, m sol water, v e sol alco-
 hol and ether, volatile with steam—B'HCl
 needles

Methyl ethyl-iso indazine $\text{C}_4\text{H}_7\text{N}_2$ $\begin{smallmatrix} \text{CMe} \\ \text{NEt} \end{smallmatrix} \text{N}$
Ethyl-quinazole [80°] (235°) Formed by
 heating its carboxylic acid at 160°-190° (Fischer
 & Kuzel, *A* 221, 285, *B* 16, 655) Formed

also by reducing the nitrosamine of ethyl amido acetophenone, dissolved in dilute acetic acid, with zinc dust, adding NaOH, and distilling with steam. The oily distillate is dissolved in dilute H_2SO_4 , mixed with $NaNO_2$, and extracted with ether (Fischer & Tafel, *A* 227, 303). Plates, sl sol water, v sol alcohol and ether. Volatile with steam, giving off a pungent odour. Not affected by nitrous acid or by Ag_2O . Does not reduce Fehling's solution. Forms crystalline pps with $AgNO_3$ and $HgCl_2$, which may be recrystallised from hot water in slender needles. Its salts are extremely soluble in water. — $B'H_2SO_4$ long needles — $B'H_2PtCl_6$ sparingly soluble orange prisms — Picrate yellow needles

Methyl-iodide $B'MeI$ [192°]

METHYL-ETHYL-ISO INDAZINE **CARB-**



Formed by treating the nitrosamine of ethyl-o-amido cinnamic acid with zinc dust and acetic acid (Fischer & Kuzel, *A* 221, 285). Colourless plates [131°] (from water) or groups of smaller plates [126°] (from chloroform ligroin). Sl sol water, v sol alcohol, ether, and chloroform. Combines with alkalis and with acids. Does not reduce boiling alkaline solutions of silver or copper. With bromine in HOAc it forms a bromo derivative [173°] and a di bromo-derivative [196°].

METHYL-ETHYL-INDOLE $C_{11}H_{13}N$ $\frac{1}{2}$ aq

$C_6H_5\left\langle\begin{array}{c} CH \\ \backslash \\ NH \end{array}\right\rangle CMe$ (292° i V) Formed by heating the phenyl hydrazide of methyl propyl ketone with $ZnCl_2$ at 180° (E Fischer, *B* 19, 1565, *A* 236, 132). Yellowish oil, v sl sol water, v sol alcohol and ether. Its picrate crystallises from hot benzene in dark red needles. With $NaNO_2$ and HOAc it yields a nitrosamine

Methyl-ethyl-indole $C_6H_5\left\langle\begin{array}{c} CH \\ \backslash \\ NH \end{array}\right\rangle CMe$

Ethyl-methyl ketole (288° i V) Formed, together with a base boiling at 256°, by heating methyl indole (methyl ketole) with EtI and alcohol at 100° (E Fischer & Steche, *B* 20, 2199). Oil, insol dilute acids. Colours pine-wood red — Picrate [146°] Slender dark-red needles (from benzene)

Methyl-ethyl-indole $C_6H_5Me\left\langle\begin{array}{c} CH \\ \backslash \\ NH \end{array}\right\rangle CH$

Ethyl p tolindole (254°) Formed by heating its carboxylic acid for some time at 205° (Hegel, *A* 232, 218). Oil, volatile with steam. Resinified by conc H_2SO_4 . Colours pine wood moistened with HCl red. Fuming HNO_3 gives a red colour and a pp

Di-methyl-ethyl-indole $C_{12}H_{15}N$ $\frac{1}{2}$ aq

$C_6H_5\left\langle\begin{array}{c} CMe \\ \backslash \\ NH \end{array}\right\rangle CMe$ (281°) Formed by heating bromo levulic (bromo acetyl propionic) acid with ethyl aniline (Wolff, *B* 21, 3563). Yellowish oil, v sol ether, alcohol, and benzene, sl sol water — Picrate $B'C_6H_5(NO_2)_2OH$ [105°] Red needles, m sol benzene

METHYL-ETHYL-INDOLE CARBOXYLIC

ACID $C_6H_5\left\langle\begin{array}{c} CH \\ \backslash \\ NH \end{array}\right\rangle\begin{array}{c} CH_2 \\ \backslash \\ CO_2H \end{array}$ [202°] Formed from the p tolyl-ethyl-hydrazide of pyruvic acid by warming with dilute (10 p.c.)

$HClAq$, or with phosphoric acid (Hegel, *A* 232, 218). V sol alkalis, ether, and benzene, insol ligroin. With $NaOCl$ it yields a chloro derivative, which is converted by boiling water into methyl ethyl ψ -isatin

PENTA-METHYL-ETHYL IODIDE v **HEPTIL**

IODIDE

METHYL-ETHYL-ISATIN v **ISATIN**

METHYL-ETHYL-KETOLE v **METHYL**

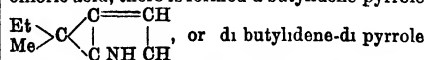
ETHYL INDOLE

METHYL-ETHYL KETONE C_6H_5O $\frac{1}{2}$ aq $CH_3COCH_2CH_3$, *Methyl acetone* Mol w 72 (81°) (F & D), (78° at 740 mm) (Schramm, *B* 16, 1581) S G $\frac{1}{2}$ 8125 (F & D), $\frac{2}{3}$ 8045 (S).

Formation — 1 By the action of $ZnEt_2$ on acetyl chloride, or of $ZnMe_2$ on propionyl chloride (Freund, *A* 118, 3, Popoff, *A* 145, 289) — 2 By boiling methyl acetoacetic ether with potash (Frankland & Duppa, *A* 138, 336), or dilute H_2SO_4 (Böcking, *A* 204, 17) — 3 Among the products of the distillation of crude calcium acetate (Fittig, *A* 110, 18) — 4 By distilling a mixture of calcium acetate and calcium propionate (Schramm, *B* 16, 1581) — 5 By the oxidation of *sec* butyl alcohol (Kanonnikoff & Saytzeff, *A* 175, 377) — 6 By the action of H_2SO_4 on crotonylene (Lwoff & Almédingen, *Bl* [2] 37, 493)

Properties — Colourless liquid, smelling like acetone, miscible with water and alcohol. With hydrogen sodium sulphite it forms the compound $MeCEt(OH)SO_3Na$ $\frac{1}{2}$ aq (Fittig)

Reactions — 1 Sodium converts it into homologues of mesityl oxide, of phorone, and of pinacone (Lawrenovitch, *B* 8, 767). Sodium, added to its solution in benzene, forms an oil $C_{12}H_{22}O$, smelling like camphor — 2 *Chromic acid* in HOAc oxidises it to acetic acid — 3 PCl_5 yields di chloro-butane $CH_3CH_2CCl_2CH_3$. On treating this body with alcoholic potash at 170° there is formed $CH_3CH_2CCH_3$, which ultimately changes into the isomeric $CH_3CCH_2CH_3$, which yields hexa methyl benzene on condensation with H_2SO_4 (Favorsky, *Bl* [2] 43, 112) — 4 On warming with *pyrrol*, MeOH, and a little hydrochloric acid, there is formed a butylidene pyrrole



$Et\left\langle\begin{array}{c} C=CH \\ \backslash \\ C-NHCH \end{array}\right\rangle\begin{array}{c} C=CH \\ \backslash \\ C-NHCH \end{array}CMe$ This body, when crystallised as $C_{16}H_{22}N$ $\frac{1}{2}$ aq, melts at 80°, but when anhydrous it melts at 142°. With alcoholic $AgNO_3$ it yields slender needles of $B'AgNO_3$ (Dennstedt & Zimmermann, *B* 20, 2454)

Oxim $CH_3CH_2\left\langle\begin{array}{c} CH \\ \backslash \\ C-NH \end{array}\right\rangle$ (153° cor) S G

$\frac{1}{2}$ 9195 Formed by the action of an aqueous solution of hydroxylamine on methyl ethyl ketone (Janny, *B* 15, 2779). Colourless liquid, soluble in ten volumes of water, miscible with alcohol and ether. Concentrated NaOHAq forms a crystalline Na salt

Reference — **HEXA-BROMO-METHYL-ETHYL KETONE**

Methyl ethyl diketone $CH_3COCOCH_2CH_3$, *Diketopentane Acetyl propionyl* (108°) S 7 at 15° Formed by the action of amyl nitrite on its mono-oxim (nitroso-di-ethyl ketone) (Manasse,

B 21, 2177) Formed also from ethyl acetoacetic ether by saponifying with dilute (3 p c) alkali, treating the product with NaNO_2 and H_2SO_4 , removing alcohol by distillation, adding dilute H_2SO_4 (15 p c) and distilling with steam (Von Pechmann, *B 21*, 1411, 2140) Dark yellow liquid, smelling like quinone, v sol ordinary solvents Forms an unstable compound with alcohol Reacts with aniline, forming a compound crystallising in needles and melting at 137° Aqueous NaOH converts it into a 'quinogen,' and finally into a quinone $[112^\circ]$, which may be reduced to a hydroquinone $[220^\circ]$, and yields a phenyl hydrazide $[225^\circ]$

(a) Oxim $\text{CH}_3\text{C}(\text{NOH})\text{CO}\text{C}_2\text{H}_5$ *Isonitroso di ethyl ketone* $[59^\circ-62^\circ]$ Formed by the action of amyl nitrite and NaOEt upon di ethyl ketone (Claisen a Manasse, *B 22*, 528) Colourless plates

(b) Oxim $\text{CH}_3\text{CO}\text{C}(\text{NOH})\text{C}_2\text{H}_5$ *Isonitroso-methyl propyl ketone* $[55^\circ]$ ($183^\circ-187^\circ$ cor) Formed by the action of KNO_2 and H_2SO_4 on ethyl acetoacetic ether (V Meyer a Zublin, *B 11*, 323, 695) Crystals, v sol alcohol and ether in sol cold water Dissolves readily in alkalis, forming a yellow solution

Di oxim $\text{CH}_3\text{C}(\text{NOH})\text{C}(\text{NOH})\text{C}_2\text{H}_5$ $[170^\circ]$ Formed by the action of hydroxylamine on the mono oxim (Sohramm, *B 16*, 180, 2187) Small white needles May be sublimed $-\text{C}_3\text{H}_7\text{N}_2\text{O}$ Na white pp

Di acetyl derivative of the *di oxim* $\text{CH}_3\text{C}(\text{NOAc})\text{C}(\text{NOAc})\text{C}_2\text{H}_5$ Prisms, v sol alcohol and ether, sl sol hot water

(a) Oxim (b) phenyl hydrazide $\text{CH}_3\text{C}(\text{NOH})\text{C}(\text{N HPh})\text{C}_2\text{H}_5$ $[128^\circ]$ Formed from the (a) oxim by treatment with phenylhydrazine (Otte a Pechmann, *B 22*, 2119) Crystallises from benzene and ligroin

(b) Oxim (a) phenyl hydrazide $\text{CH}_3\text{C}(\text{N HPh})\text{C}(\text{NOH})\text{C}_2\text{H}_5$ $[131^\circ]$ Formed from the (b) oxim and phenyl hydrazine (Otte a Pechmann, *B 22*, 2118) Crystallises from benzene and ligroin

(a) Phenyl hydrazide $\text{CH}_3\text{C}(\text{N HPh})\text{CO}\text{C}_2\text{H}_5$ $[103^\circ]$ Formed by the action of phenyl hydrazine on the diketone (Otte a Pechmann, *B 22*, 2115)

(b) Phenyl hydrazide $\text{CH}_3\text{CO}\text{C}(\text{N HPh})\text{C}_2\text{H}_5$ $[117^\circ]$ Formed by saponifying ethyl acetoacetic ether by allowing it to stand with aqueous KOH , then adding di azobenzene chloride, and ppg by sodium acetate solution (Japp a Klingemann, *C J* 53, 519, *B 21*, 550, *A 247*, 220) Yellow radiating needles or prisms (from benzene)

Di phenyl hydrazide $\text{CH}_3\text{C}(\text{N HPh})\text{C}(\text{N HPh})\text{C}_2\text{H}_5$ $[162^\circ]$ (J), $[166^\circ-169^\circ]$ (C a M) Formed by the action of phenyl hydrazine on the diketone, on the (b) phenyl hydrazide, or on the (a) oxim (Von Pechmann, *B 21*, 1414, Japp, *A 247*, 221, Claisen a Manasse, *B 22*, 528) Yellow needles (from benzene), sol cold conc H_2SO_4 , forming a brown solution

METHYL-ETHYL KETONE CARBOXYLIC ACID v *Methyl acetoacetic acid*

METHYL-ETHYL MALONIC ACID
 $\text{C}_6\text{H}_{10}\text{O}_4$ v $\text{CH}_3\text{CH}_2\text{CMe}(\text{CO}_2\text{H})_2$ Mol. w. 146
 $[119^\circ]$ H O v 67,200 H Cp 673,300 H F
 Vol. III

236,700 (Stohmann, Kleber, a Langbein, *J pr.* [2] 40, 210)

Formation—1 By saponifying its ether which is obtained by treating ethyl malonic ether with MeI and NaOEt , or methyl-malonic ether with EtI and NaOEt (Conrad a Bischoff, *B 13*, 596, *A 204*, 146)—2 Together with a larger quantity of the isomeric s di methyl succinic acid $[194^\circ]$, by reducing the anhydride of di methyl maleic acid (pyrocinchonic acid) with HI or sodium amalgam (Otto a Beckurts, *B 18*, 841)

Properties—Prisms, v sol water, alcohol, and ether At 180° it splits off CO , giving valeric acid $\text{CH}_3\text{CH}_2\text{CHMeCO}_2\text{H}$ A solution of its ammonium salt gives pps with FeCl_3 and with salts of Hg and Pb

Salt— AgA' Sparingly soluble white crystalline pp

Ethyl ether EtA' (208°) $\text{SG } \frac{1}{12}$ 994
 Oil, sol alcohol and ether

METHYL-ETHYL-METHANE v BUTANE

Methyl di ethyl-methane v HEXANE

Tri-methyl-ethyl-methane v TERT HEXANE

METHYL-ETHYL-PENTAMETHENYL

METHYL KETONE TRIHYDRIDE $\text{C}_{10}\text{H}_{16}\text{O}$ v s

$\text{CH}_3\text{CO} \begin{array}{c} \text{CMe CHEt} \\ | \\ \text{CH}_2\text{CH}_2 \end{array}$ *Ethyl methyl-dihydro-*

pentene methyl ketone $(210^\circ-215^\circ)$ Formed from di butylene diketone dicarboxylic ether (di acetyl adipic ether) by heating with EtI and alcoholic NaOEt The product is freed from alcohol by distillation, mixed with water, and the oil, which can then be extracted by ether, hydrolysed by boiling alcoholic potash (Marshall a Perkin, *C J* 57, 252)

Oxim $\text{C}_{10}\text{H}_{16}\text{NO}$ Oil

METHYL-ETHYL-PENTAMETHYLENE

C_8H_{16} v s $\text{CH}_2 \begin{array}{c} \text{CH}_2\text{CHMe} \\ | \\ \text{CH}_2\text{CHEt} \end{array}$ (124°) Formed

by heating methyl iodo ethyl pentamethylene (obtained from methyl pentamethylenyl methyl carbinol and HI) with amorphous phosphorus and HI at 245° (Marshall a Perkin, jun, *C J* 57, 250) Limpid oil, attacked by bromine with difficulty Rapidly oxidised by boiling dilute HNO_3

METHYL ETHYL OXIDE $\text{C}_4\text{H}_8\text{O}$ v s

$\text{CH}_3\text{O}\text{C}_2\text{H}_5$ Mol w 60 (11°) $\text{SG } \frac{1}{10}$ 7252 (Dobner, *A 243*, 2) $\text{SV } 840$ (D) $\text{VD } 2158$ HF p 58,450 (Thomsen, *Th*) HF v 56,420

Formation—1 By the action of MeI on NaOEt or of EtI on NaOMe (Williamson, *P M* [3] 37, 350 *C J* 4, 106, 229)—2 By distilling potassium methyl-sulphate with KOEt (Chance, *C R* 31, 152)—3 By the action of Ag_2O on a mixture of MeI and EtI (Wurtz, *J* 1856, 563)—4 Together with Me_2O and Et_2O by etherification of a mixture of methyl and ethyl alcohols by H_2SO_4 (Williamson, cf Norton a Prescott, *Am 6*, 244)

Properties—Colourless mobile liquid, with ethereal odour

METHYL-ETHYL-OXY- v. **OXY METHYL-**

ETHYL
METHYL-ETHYL-PHENOL $\text{C}_9\text{H}_{10}\text{O}$ v s
 $\text{CH}_3\text{C}_2\text{H}_4\text{EtOH}$ (215°) Prepared by fusing *p* methyl-ethyl-benzene sulphonic acid with

potash, dissolving the product in water, acidifying, and extracting with ether (Mazzara, *G* 10, 256). Oil, sl sol water. Gives an azure blue colouration with FeCl_3 .

Di methyl ethyl-phenol $\text{C}_9\text{H}_{10}\text{Me}_2\text{Et}(\text{OH})$ [6 8 1 2] ? [87°] (245°) Obtained by potash-fusion from the corresponding di methyl ethyl benzene sulphonic acid (Stahl, *B* 23, 990) FeCl_3 gives an intense green colour in alcoholic (not in aqueous) solution

METHYL - ETHYL - PHENYL - AMINE *v* AMIDO TOLYL ETHANE

Tri-methyl-di-ethyl-phenyl-amine $\text{C}_{15}\text{H}_{19}\text{N}$ *v* $\text{C}_6\text{Me}_2\text{Et}_2\text{NH}_2$ [1 2 5 3 6 4] (286°-290°) SG 971 Colourless oil Obtained by heating a mixture of mono and di ethyl pseudo cumidine with an equal weight of ethyl iodide at 260°-280° for 8-10 hours (Ruttan, *O J* 49, 813, *B* 19, 2383)

Salts— $\text{B}'\text{HCl}$ needles, easily sol alcohol, *v* sl sol hot water, insol cold water. The acetate and sulphate form easily soluble needles. The oxalate forms very sparingly soluble quadratic prisms

Acetyl derivative $\text{C}_6\text{Me}_2\text{Et}_2\text{NHAc}$ [182°], needles

DI-METHYL-DI-ETHYL-*p* PHENYLENE-DIAMINE $\text{C}_{12}\text{H}_{16}\text{N}_2$ *v* $\text{C}_6\text{H}_4(\text{NMe}_2)(\text{NEt}_2)$ (265°) Prepared by heating di ethyl *p* phenyl ene diamine with methyl alcohol and HCl (Lippmann a Fleissner, *M* 4, 788) Light yellow oil

Di-methylo di iodide $\text{B}'\text{Me}_2\text{I}_2$ [218°] Formed by heating di ethyl amido benzene azo di ethyl aniline with MeI Transparent four-sided prisms— $\text{B}'\text{Me}_2\text{I}_2\text{CdI}_2$ silky needles— $\text{B}'\text{Me}_2\text{PtCl}_4$ orange red needles— $\text{B}''(\text{MeAuCl}_4)_2$ yellow leaflets— $\text{B}''2\text{MeOC}_6\text{H}_4(\text{NO}_2)_2$ [235°] Yellow needles

DI METHYL ETHYL PHOSPHATE $\text{C}_4\text{H}_{11}\text{PO}_4$ *v* Me_2EtPO_4 (203° cor) SG 2 1 1752 SV 1611 (Lossen, *A* 254, 74) Formed from Me_2AgPO_4 and EtI (Weger, *A* 221, 90)

METHYL-DI ETHYL-PHOSPHINE $\text{C}_4\text{H}_{13}\text{P}$ *v* Et_2PMe (111°) Formed by heating tri ethyl phosphine methylo chloride Et_3PMeCl at 300° (Collie, *C J* 53, 719) Combines with sulphur and with oxygen Forms red crystals with CS_2

Methylo-chloride v Ethylo chloride of Di methyl ethyl phosphine

Ethylo iodide v TRI ETHYL PHOSPHINE

TRIETHYL IODIDE
Di methyl ethyl phosphine Me_2PEt (84°) Formed by heating Me_2PEtHCl (the product of the distillation of its ethylo chloride) with aqueous NaOH (Collie, *C J* 53, 720)

Methylo chloride v Ethylo chloride of TRI METHYL PHOSPHINE

Ethylo chloride $\text{Me}_2\text{PEt}_2\text{Cl}$ Formed from $\text{Me}_2\text{PEt}_2\text{I}$, the product of the union of Et_2PMe and MeI Split up by heat into ethylene and Me_2PEtHCl (Collie)

METHYL - TRI - ETHYL - PHOSPHONIUM CHLORIDE *v* *Methylo iodide of TRI-ETHYL-PHOSPHINE*

Di-methyl-di-ethyl-phosphonium chloride *v* *Ethylo-chloride of Di-methyl ethyl phosphine*

Tri-methyl-ethyl-phosphonium chloride *v* *Ethylo-chloride of TRI METHYL-PHOSPHINE*

METHYL-ETHYL PINACONE *v* **DI-OXY-OTANE.**

METHYL-ETHYL-PIPERIDINE *v* **METHYL ETHYL PYRIDINE HEXAHYDRIDE**

METHYL-ETHYL-PROPIONIC ACID *v* **HEXOID ACID**

METHYL-ETHYL-PROPYL ALCOHOL *v* **HEXYL ALCOHOL**

METHYL-ETHYL-PROPYL-AMINE $\text{C}_9\text{H}_{11}\text{N}$ *v* $\text{C}_6\text{H}_5\text{NMeEt}$ A product of the destructive distillation of the methylo hydroxide of ethyl-codeine (Von Gerichten a Schrötter, *B* 15, 1484) Volatile base, smelling like trimethylamine— $\text{B}'_2\text{H}_2\text{PtCl}_4$ Long orange needles, *v* sol water, sl sol alcohol

METHYL ETHYL-PROPYL-CARBINOL *v* **Tert HEPTYL ALCOHOL**

METHYL-ETHYL-PROPYLENE *v* **HEXYLENE**

METHYL-ETHYL-PROPYL-METHANE *v* **HEPTANE**

DI-METHYL-DI-ETHYL-PYRAZINE

$\text{C}_{10}\text{H}_{12}\text{N}_2$ *v* $\text{N} \begin{smallmatrix} \text{CMe} \text{Cet} \\ \text{Cet} \text{CMe} \end{smallmatrix} \text{N}$ *Di ethyl ketone* (216° cor) VD 5 63 (calo 5 68) Prepared by reduction of the oxim of methyl ethyl diketone (methyl nitroso propyl ketone) $\text{Me CO C}(\text{NOH})\text{Et}$ with tin and HCl , or with sodium amalgam (Treadwell, *B* 14, 1461) Colourless oil with narcotic smell and alkaline reaction It is a weak poison Combines with water forming a crystalline hydrate Not affected by MeI , acetic anhydride, or HI Bromine added to its solution in HOAc ppts $\text{C}_{10}\text{H}_{12}\text{N}_2\text{Br}_2$ as an unstable yellow compound Very dilute potassium permanganate oxidises it to methyl pyrazinedi carboxylic acid $\text{N} \begin{smallmatrix} \text{CMe} \text{C}(\text{CO}_2\text{H}) \\ \text{C}(\text{CO}_2\text{H}) \text{CMe} \end{smallmatrix} \text{N}$, which does not yield an anhydride (Economidis, *B* 19, 2524) — $\text{B}'\text{HCl}$ Large colourless crystals — $\text{B}'\text{H PtCl}_4$ Soluble red prisms — $\text{B}'\text{AgNO}_3$ Crystals, almost insol cold water, *v* sol dilute HNO_3 (Treadwell, *B* 14, 2158)

Hydrate $\text{B}'\text{zaq}$ [43°] Large prisms (by sublimation) Gives up its water in dry air, becoming liquid

METHYL ETHYL PYRIDINE $\text{C}_8\text{H}_{11}\text{N}$ *v* $\text{N} \begin{smallmatrix} \text{CH} \text{CH} \\ \text{CMe} \text{CH} \end{smallmatrix} \text{Cet}$ (*a*) *Collidine* Mol w 121

(179°) SG 2 929 (Richard, *Bl* [2] 32, 488), 10 929 (Weidel a Pick, *M* 5, 659) SV 157 9 (Ramsay) A base found, together with many others, among the products of the dry distillation of bones (Anderson, *P M* [4] 9, 145, 214, *A* 94, 360), of bituminous shale (Greville Williams, *C J* 7, 97) and of peat (Church a Owen, *P M* [4] 20, 110) It occurs in coal tar Colourless oil, with unpleasant odour Fumes with HCl *V* sl sol water, separating again when warmed, *v* sol alcohol, ether, and oils Ppts ferric, aluminium, chromium, and mercurous oxides from their salts, but not manganese and nickel oxides Its salts are deliquescent and gummy KMnO_4 oxidises it to pyridine dicarboxylic (lutidmic) acid [219°], which is converted by heat into pyridine carboxylic acid [302°] — $\text{B}'_2\text{H}_2\text{PtCl}_4$ Orange yellow flakes

Ethylo-iodide $\text{B}'\text{EtI}$ Oil — ($\text{B}'\text{EtCl}_3$) PtCl_4 Sparingly soluble crystalline pp (Anderson, *P M* [4] 9, 221)

'*a*'-*Collidine* $\text{C}_8\text{H}_{11}\text{N}$ (179°-183°) This base, according to Oechsner de Coninck (*A Ch* [5] 27, 468), accompanies (*B*) collidine in the

product of the distillation of cinchonine with KOH. It is perhaps identical with (a) collidine.

When 'a' collidine is heated with *ethylene chlorhydrin* and a little water for a few hours at 100° combination takes place. After removing uncombined collidine and chlorhydrin by shaking with ether, the residue is evaporated *in vacuo*. PtCl₄ forms with the product an orange yellow pp of $\{C_8H_{11}N(C_2H_4OH)Cl\}_2PtCl_4$. Boiling water removes HCl, forming oxethyl- α collidine chloro platinite $\{C_8H_{11}N(C_2H_4OH)Cl\}_2PtCl_4$. This forms brilliant scales (from alcohol). Oxethyl- α collidine forms a gold salt, B'AuCl₄, crystallising in thin golden needles (Wurtz, *Pr* 33, 450, *C R* 95, 263, *Bl* [2] 39, 536).

Methyl-ethyl pyridine $N \begin{smallmatrix} \text{CH CEt} \\ \text{CH CH} \end{smallmatrix} > \text{CMe}$

(*B*) *Collidine* (196°) SG $\frac{2}{966}$ VD 4 25 (calc 4 19). Obtained by distilling cinchonine or brucine with KOH (Oechsner de Coninck, *C R* 91, 296, *A Ch* [5] 27, 469, *Bl* [2] 37, 457, 42, 100, *R T C* 1, 132, *Cf* Greville Williams, *Tr E* 21, Part 2, *A Ch* [3] 45, 488). Colourless, mobile, highly refractive liquid, rapidly turning yellow on exposure to air. Sl sol water, sol alcohol and ether. When left in contact with moist air it takes up water (1 mol) but does not form a well defined hydrate. Potassium permanganate oxidises it to homo nicotinic acid $N \begin{smallmatrix} \text{CH C(CO}_2\text{H)} \\ \text{CH CH} \end{smallmatrix} > \text{CMe}$, cinchomeronic acid $N \begin{smallmatrix} \text{CH C(CO}_2\text{H)} \\ \text{CH CH} \end{smallmatrix} > \text{C CO}_2\text{H}$, and formic acid. It is very poisonous, a sub cutaneous injection of 1 g paralysing the nerve centres. HI reduces it to the hydride $C_8H_{11}N$, and, at 250°, forms a brown oil $C_8H_{11}NI_4$. Sodium in alcohol reduces it to the hexahydride $C_8H_{11}N$ (175°-180°).

Salts - B'HCl. White deliquescent tables - B'H PtCl₄. S 21 at 60° (Richard, *Bl* [2] 32, 488). Orange red powder, converted by boiling water into B' PtCl₄, a yellow crystalline powder - B'HAuCl₄ - B' HCl $\frac{1}{3}$ AuCl₃ - B' H₂HgCl₂. Minute white needles, sl sol water, insol alcohol.

Methyl-ethyl pyridine $N \begin{smallmatrix} \text{CMe CH} \\ \text{CH CEt} \end{smallmatrix} > \text{CH}$

Aldehyde collidine (176°) SG $\frac{2}{9389}$ (D). Occurrence - As acetate in fusel oil (Kramer a Pinner, *B* 3, 77).

Formation - 1 By heating ethyldene chloride with alcoholic or aqueous ammonia for 12 hours at 160° (Kramer, *Z* [2] 6, 568, *B* 3, 202, Durkopt, *B* 18, 920). Ethyldene bromide may be used instead of the chloride, the temperature employed being then 125°-140° (Tavildaroff, *A* 176, 12) - 2 By heating an alcoholic solution of aldehyde ammonia at 120° (Baeyer a Ador, *A* 155, 297). 'Para' collidine, picoline, and lutidine are also formed in this reaction (Vohl, *J* 1870, 807) - 3 By distilling aldol ammonia (aldehyde) (Wurtz, *Bl* [2] 31, 433) - 4 By heating glycol with ammonium chloride for 8 hours at 185°, the yield being from 15 to 20 p.c. of the theoretical quantity (Hofmann, *B* 17, 1905) - 5 By heating ethylene chloride with ethylamine at 190° (Hofmann, *B* 17, 1907) - 6 By heating paraldehyde with acetamide and P₂O₅ at 160° (Hesekiel, *B* 18, 3091) - 7 By heating aldehyde-ammonia with paraldehyde at 220° (Ladenburg a Durkopt, *A* 247, 42).

Properties - Oil, with aromatic odour, insol water and dilute acids, v sol alcohol and ether. Fumes with HCl. Oxidised by KMnO₄ to methylpyridine carboxylic acid $N \begin{smallmatrix} \text{CMe CH} \\ \text{CH C(CO}_2\text{H)} \end{smallmatrix} > \text{CH}$

and isocinchomeronic acid $N \begin{smallmatrix} \text{C(CO}_2\text{H)} \text{CH} \\ \text{CH C(CO}_2\text{H)} \end{smallmatrix} > \text{CH}$ (Durkopt a Schlaugk, *B* 21, 294, *cf* Wischnegradsky, *B* 12, 1506). Bromine forms an oily compound. Fuming HIAq and amorphous phosphorus at 140° yield brownish blue prisms of $C_8H_{11}NI_4$ or $C_8H_{11}NI_4$ (Ladenburg, *B* 14, 232). Sodium, acting on its alcoholic solution, forms a hexahydride. Heated for some days at 100° with glycolic chlorhydrin and a little water it forms oxethyl aldehyde, of which the platinochloride, $\{C_8H_{11}N(C_2H_4OH)Cl\}_2PtCl_4$, forms orange crystals (from dilute alcohol). It may be decomposed by H₂S, the hydrochloride produced would not crystallise, but it is converted by Ag₂O into a caustic base (Wurtz, *Pr* 33, 448).

Salts - B'₂H₂PtCl₄ [180°] Very unstable trimetric crystals - B'HAuCl₄ [72°] Yellow needles, v sol water (Hesekiel, *cf* Herzog, *M* 2, 404) - B'C₆H₅(NO₂)₂OH [157°] Greenish-yellow four sided tables.

Ethylo iodide B'EtI. Trimetric tables, v sol water and alcohol. Yields (B'EtCl)₂PtCl₄, crystallising in needles or prisms.

Methyl ethyl pyridine $N \begin{smallmatrix} \text{CMe CH} \\ \text{CH CEt} \end{smallmatrix} > \text{CH}$

(161°) SG $\frac{2}{9361}$. Formed by heating (a)-methyl pyridine (a picoline) with EtI at 280°-300° (Ladenburg a Schultz, *A* 247, 46, *B* 20, 2720). Liquid, almost insol water, but takes up water from the air. Very volatile with steam. Oxidised by KMnO₄ to pyridine dicarboxylic acid $N \begin{smallmatrix} \text{C(CO}_2\text{H)} \text{CH} \\ \text{C(CO}_2\text{H)} \text{CH} \end{smallmatrix} > \text{CH}$ [226°]. Sodium reduces it, in alcoholic solution, to a hexahydride.

Salts - B'₂H₂PtCl₄ [174°] Triclinic tables, sl sol cold, v sol hot, water, insol alcohol - B'HAuCl₄ [110°] Yellow needles, sl sol water, v sol ether alcohol.

Methyl ethyl pyridine $N \begin{smallmatrix} \text{CMe CH} \\ \text{CH CH} \end{smallmatrix} > \text{CEt}$

(173°) SG $\frac{2}{9353}$, $\frac{20}{9218}$. Formed by heating (a) methyl pyridine with EtI at 280°-300°, and separated from the preceding by fractional distillation (Schultz, *B* 20, 2720, Ladenburg, *A* 247, 46). Colourless hygroscopic liquid, with unpleasant odour, sl sol water. Oxidised by KMnO₄ to pyridine dicarboxylic acid $N \begin{smallmatrix} \text{C(CO}_2\text{H)} \text{CH} \\ \text{CH} \end{smallmatrix} > \text{C CO}_2\text{H}$. Sodium, added to its alcoholic solution, reduces it to the corresponding hexahydride.

Salts - B'₂H₂PtCl₄ [190°] Reddish yellow plates, sl sol cold, v sol hot, water, insol alcohol - B'HAuCl₄ [90°] Yellow needles, sl sol cold, m sol hot, water, v sol alcohol - The phosphomolybdate and bismuthiodide are amorphous, the cadmium iodide and periodide are only.

Paracollidine $C_8H_{11}N$ (220°-230°) Formed, together with the collidine boiling at 176°, by heating aldehyde ammonia (Baeyer a Ador, *A* 155, 307). Liquid, with pungent aromatic odour. Its salts crystallise with difficulty. Its ethyloiodide is syrupy. The platinochloride is amorphous.

Isomeride of collidine—Obtained by the putrefaction at 40° of pancreas (200 g) mixed with gelatin (600 g) and water (10,000 g) After 5 days the liquid is distilled, first with H_2SO_4 , and afterwards with baryta. The alkaline distillate is neutralised by HCl , evaporated to dryness, and extracted with alcohol. The hydrochloride of the base crystallises from alcohol in trimetric needles. The free base is got by adding NaOH to the hydrochloride and shaking with ether (Nencki, *J pr* [2] 26, 49). —($\text{B}'\text{HCl}$), PtCl_4 . Flat needles. Gives off an odour of xylene when distilled. The base is not identical with collidine (from NH_3 and ethyldene chloride), for it differs in odour, in the crystalline form of the platinum chloride, and in being more soluble in water.

Isomeride of collidine $\text{C}_6\text{H}_7\text{N}$ (202°) SG 2 987. Occurs in the flesh of putrid cuttle fish (Oechsner de Connick, *C R* 106, 858, 1604). Mobile liquid, v sol water, alcohol, and ether. Absorbs moisture from the air, becoming brown. — $\text{B}'\text{HCl}$. Deliquescent radiating masses. — $\text{B}'\text{H}_2\text{PtCl}_6$. Deep yellow crystals, almost insol in cold water. Decomposed by hot water, with formation of $\text{B}'_2\text{PtCl}_6$, a pale brown powder. — $\text{B}'\text{H}_2\text{HgCl}_4$. Small white needles. — $\text{B}'\text{H}_2\text{Hg}_2\text{Cl}_6$. Long yellowish needles.

Methyl iodide $\text{B}'\text{MeI}$. Slender needles.

Isomeride of collidine v TRI METHYL PYRIDINE

Di-methyl-ethyl pyridine $\text{C}_6\text{H}_7\text{N}$ *Parvoline* (199° cor) (D & G) SG 2 9419, 2 9289 (D & S).

Formation—1 By heating propionic aldehyde ammonia (Waage, *M* 4, 718). —2 By heating methyl-ethyl acrolein ammonia (Hoppe, *M* 9, 643). —3 By heating propionic aldehyde with acetamide and P_2O_5 (Hesekiel, *B* 18, 3097). —4 By heating propionic aldehyde ammonia with paraldehyde in sealed tubes for 10 hours at 210° (Durkopf & Schlaug, *B* 21, 832). —5 By heating propionic aldehyde ammonia (4 g) with propionic aldehyde (6.5 g) for 6 hours at 210° (Durkopf & Gottsch, *B* 23, 685).

Properties—Colourless oil, not turned brown by light and air, with faint and not unpleasant smell. M sol cold water, sl sol hot water, v sol alcohol and ether. KMnO_4 oxidises it to methyl pyridine dicarboxylic acid [225°] and pyridine tricarboxylic acid [318°].

Salts— $\text{B}'\text{H}_2\text{PtCl}_6$ [189°]. Large monoclinic crystals, sl sol cold water. — $\text{B}'\text{HAuCl}_4$ [82°]. Glittering lemon yellow spangles, sl sol cold water. Not hygroscopic. — $\text{B}'\text{HHg}_2\text{Cl}_6$ [119°]. Long pointed needles, almost insol in cold water. — $\text{B}'\text{C}_6\text{H}_5(\text{NO}_2)_2\text{OH}$ [152°]. Yellow plates.

Di-methyl ethyl pyridine $\text{C}_6\text{H}_7\text{N}$ (217°). Formed, together with the preceding, by heating propionic aldehyde ammonia with propionic aldehyde at 200° (Durkopf & Gottsch, *B* 23, 692). Colourless liquid, not affected by air and light. Has a faint odour of nicotine. V sol alcohol and ether, sl sol water. Yields a di-methyl pyridine carboxylic acid on oxidation.

Salts— $\text{B}'\text{H}_2\text{PtCl}_6$ [above 270°]. Yellowish-red needles, sl sol water. — $\text{B}'\text{HAuCl}_4$ [140°]. Slender lemon-yellow needles, sl sol. water containing HCl . — $\text{B}'\text{HHg}_2\text{Cl}_6$ [159°]. Long

glittering spangles or well formed monoclinic crystals, sl sol cold water.

Di methyl-ethyl pyridine $\text{C}_6\text{H}_7\text{N}$ e

$\text{N} \begin{smallmatrix} \text{CHMe} \\ \text{CH} \end{smallmatrix} \text{CH} \text{CEt}$ (187°) SG 14 916 S

1.8 at 0°. Obtained by the action of propionic aldehyde and ammonia on acetoacetic ether (Jaekle, *A* 246, 45). Formed also by heating to redness a mixture of potassium di-methyl ethyl pyridine dicarboxylate (1 mol) with lime (2 mols) (Engelmann, *A* 231, 44). Liquid, more soluble in cold than in hot water. Its solution has a very bitter taste. KMnO_4 oxidises it to methyl ethyl pyridine carboxylic acid and uvonic acid.

Salts— $\text{B}'\text{H}_2\text{PtCl}_6$ [211°]. Crystalline pp, v sol, hot water and alcohol. — $\text{B}'_2\text{H}_2\text{Cr}_2\text{O}_7$. Crystalline pp. —Picrate [120°]. Needles.

METHYL-ETHYL PYRIDINE DIHYDRIDE

$\text{C}_6\text{H}_7\text{N}$ (8) *Di-hydro collidine* (200°–205°). Formed by heating (8) collidine with HIAq and red phosphorus at 140° (Oechsner de Connick, *Bl* [2] 42, 116).

Methyl ethyl pyridine hexahydride $\text{C}_6\text{H}_7\text{N}$ (c 177°). Formed by adding sodium to an alcoholic solution of (8) collidine (Oechsner de Connick, *Bl* [2] 42, 116).

Methyl ethyl pyridine hexahydride $\text{C}_6\text{H}_7\text{N}$

e $\text{NH} \begin{smallmatrix} \text{CHMe} \\ \text{CH}_2 \end{smallmatrix} \text{CH} \text{CH}_2$ *Copellidine* (164°)

SG 2 8653, 2 8546. Formed from the collidine derived from aldehyde, by reduction in alcoholic solution with sodium (Durkopf, *B* 18, 920, Ladenburg, *A* 247, 90). Colourless alkaline liquid, sl sol water. Its physiological action resembles that of conine, but is weaker.

Salts— $\text{B}'\text{HCl}$ [171°]. Colourless needles, v sol water. — $\text{B}'\text{HBr}$ [165°]. Tufts of needles. — $\text{B}'\text{HAuCl}_4$ [104°]. Dimetric prisms, v sol hot water. — $\text{B}'\text{H}_2\text{PtCl}_6$ [147°]. Long yellow needles, v e sol water.

Acetyl derivative $\text{C}_6\text{H}_7\text{N} \cdot \text{Ac}$ (254°)

SG 2 9787, 2 966

Methyl ethyl pyridine hexahydride

$\text{NH} \begin{smallmatrix} \text{CHMe} \\ \text{CH}_2 \end{smallmatrix} \text{CH} \text{CH}_2$ (147°–151°) SG 2

8550, 2 8410. Formed by reducing the corresponding methyl-ethyl pyridine in alcoholic solution with sodium, the product being purified by means of its oily nitrosamine (Schultz, *B* 20, 2723, Ladenburg, *A* 247, 95). Colourless alkaline liquid, smelling strongly like ammonia and like piperidine, sl sol water.

Salts— $\text{B}'\text{HCl}$. Needles, v e sol water and alcohol. — $\text{B}'\text{HBr}$. Needles.

Methyl ethyl pyridine hexahydride

$\text{NH} \begin{smallmatrix} \text{CHMe} \\ \text{CH}_2 \end{smallmatrix} \text{CH} \text{CH}_2$ *Copellidine* (155°–

160°) SG 2 8515, 2 8389. Obtained by reducing the corresponding methyl ethyl pyridine in alcoholic solution with sodium, the product being purified by means of the nitrosamine (Ladenburg & Schultz, *B* 20, 2723, *A* 247, 96). Colourless, strongly alkaline liquid. — $\text{B}'\text{HCl}$ [c 218°]. White needles, v e sol water and alcohol.

Methyl ethyl-pyridine hexahydride

$\text{NMe} \begin{smallmatrix} \text{CH} \\ \text{CH}_2 \end{smallmatrix} \text{CH} \text{CH}_2$ (c 149°) SG 2

8495. Formed by the action of MeI on the corresponding ethyl-pyridine hexahydride dis-

solved in MeOH, the product being distilled with strong aqueous NaOH (Ladenburg, A 247, 71) Liquid, smelling like ν methylpyridine hexahydride, sl sol water Its hydrochloride crystallises in small needles

Di-methyl-ethyl pyridine hexahydride
 $C_8H_{11}N$ *Parpevoline* (177°) SG 2 8628, 2 8542, 2 8474 Formed at the same time as di methyl ethyl pyridine (109°), when propionic aldehyde ammonia and propionic aldehyde are heated together at 200° Formed also by reducing the same di methyl ethyl pyridine by adding sodium to its alcoholic solution (Dürkopf a Gottsch, B 23, 690) Very mobile colourless liquid, with penetrating odour like rotten hay, v sol ether and alcohol, sl sol water Strongly alkaline in reaction — $B'H_2Cl$, [120°–130°] White sparingly soluble needles (from hot water) — The hydrochloride and hydroiodide are hygroscopic Chlorides of gold and of platinum give no pp in dilute solution, and only pps in concentrated solutions Picric acid acts in like manner

Di methyl ethyl pyridine hexahydride

$C_8H_{11}N$ ν NMe $\langle \begin{smallmatrix} CHMe \\ CH_2 \end{smallmatrix} \rangle CH_2$ ν Methylcopellidine (165°) VD (H=1) 139 5 (obs) SG 2 8519, 1 841 Formed from methyl-ethyl pyridine hexahydride (164°) and MeI in the cold (Ladenburg a Dürkopf, B 18, 925, A 247, 90) Strongly alkaline colourless oil, sl sol water — $B'HBr$ [151°] White needles — Picrate [162°] Yellow needles

Methyl iodide B'MeI [268°] Long white needles, m sol water, v sol alcohol, ppd by ether from alcoholic solution With moist Ag_2O it yields a methyl hydroxide which is split up on distillation into water and tri methyl ethyl pyridine

Methyl chloride B'MeCl Formed from the iodide by shaking with $AgCl$ — ($B'MeCl$), $PtCl_4$, Yellow octahedra — ($B'MeCl$), $HgCl_2$, [108°] Long white needles

Di methyl ethyl pyridine hexahydride

$C_8H_{11}N$ ν NH $\langle \begin{smallmatrix} CHMe \\ CH_2 \end{smallmatrix} \rangle CH_2$ *Parperoline* *Ethyl lupetidine* (167°) at 725 mm Formed by reducing di methyl ethyl pyridine (187°) in alcoholic solution with sodium (Jaechle, A 246, 45) Liquid Gives, in a dilute alcoholic solution, dark brown pps with silver or mercurous nitrate — $B'H_2PtCl_4$, Orange prisms — $B'H_2Cr_2O_7$, Long prisms

Tri methyl ethyl pyridine hexahydride

$C_{10}H_{13}N$ ν C_2H_5 , MeNMe *Di methyl collidine hexahydride* *Di methyl copellidine* (172°) SG 2 7816 Formed by distillation of the methyl hydroxide of ν methyl copellidine (di methyl ethyl pyridine hexahydride) (165°) (Dürkopf, B 18, 927, Ladenburg, A 247, 94) Colourless alkaline liquid, sl sol water. Smells like trimethylamine — $B'HAnCl$, Yellow needles or glistening plates — $B'H_2PtCl_4$, [98°] Silky yellow needles

METHYL ETHYL PYRIDINE CARBOXYLIC

ACID $C_8H_9NO_2$ ν N $\langle \begin{smallmatrix} C(CO_2H) \\ C(CH_3) \end{smallmatrix} \rangle CH$ \rangle CEt Formed by oxidising the corresponding di-methyl-ethyl-pyridine with $KMnO_4$ (Altar, A 228, 189) — (HA'), H_2PtCl_4 , [195°]. Red prisms

Di methyl ethyl pyridine dicarboxylic acid

$C_{11}H_{13}NO_4$ ν N $\langle \begin{smallmatrix} CMe \\ CMe \end{smallmatrix} \rangle \langle \begin{smallmatrix} C(CO_2H) \\ C(CO_2H) \end{smallmatrix} \rangle$ CEt *Parvoline dicarboxylic acid* [290°] Obtained by saponifying its ether with alcoholic potash (Engelmann, A 231, 40) Thick prisms, m sol cold water and alcohol, decomposed on fusion On distillation with lime it yields di methyl-ethyl pyridine — $BA'A'$ 3aq Plates, v e sol water — (H_2A'), HCl aq Prisms

Ethylether Et.A' (305°–308°) Formed by passing nitrous acid gas into an alcoholic solution of the dihydride Thick liquid — (Et_2A'), H_2PtCl_4 , [139°] Long prisms (from alcohol)

DI-METHYL-ETHYL PYRIDINE DIHYDRIDE DICARBOXYLIC ETHER $C_{13}H_{21}NO_4$ ν e

NH $\langle \begin{smallmatrix} CMe \\ CMe \end{smallmatrix} \rangle \langle \begin{smallmatrix} C(CO_2Et) \\ C(CO_2Et) \end{smallmatrix} \rangle$ CHEt *Hydroparvoline dicarboxylic ether* [110°] Formed by boiling acetoacetic ether (45 g) with propionic aldehyde (10 g) and alcoholic ammonia (Engelmann, A 231, 38) Prisms (from alcohol), v e sol hot alcohol, ether, chloroform, and benzene

METHYL ETHYL PYRIMIDINE $C_8H_9N_2$ ν e

CEt $\langle \begin{smallmatrix} N \\ N \end{smallmatrix} \rangle \langle \begin{smallmatrix} CMe \\ CH \end{smallmatrix} \rangle CH$ (160°) Formed by reducing oxy methyl ethyl pyrimidine with zinc dust (Pinner, B 22, 1619) Prisms (Py 2 3) **METHYL ETHYL QUINOLINE**

$C_{12}H_{13}N$ ν e C_2H_5 $\langle \begin{smallmatrix} CH \\ N \end{smallmatrix} \rangle \begin{smallmatrix} CMe \\ CMe \end{smallmatrix}$ [56°] (268° at 710

mm) Formed by the action of propionic aldehyde and HCl upon aniline (Dabner a Miller, B 17, 1714) Large monoclinic prisms V sol alcohol, ether, and benzene, sl sol water On oxidation with CrO_3 and H_2SO_4 it yields (Py 2)-methyl quinoline (Py 3) carboxylic acid

Salts — $B'HCl$, $PtCl_4$, fine yellow needles. — Picrate $B'_2C_6H_4(NO_2)_4OH$ yellow crystals — $B'_2H_2Cr_2O_7$, small yellowish brown prisms

Methyl iodide B'MeI [196°], yellow needles, insol ether

(B 2) Methyl (Py 3) ethyl-quinoline

$C_8H_9(CH_3) \langle \begin{smallmatrix} CH \\ N \end{smallmatrix} \rangle \begin{smallmatrix} CMe \\ CMe \end{smallmatrix}$ *Ethyl toluquinoline*

[60°] (270° at 718 mm) Formed by heating its (Py 2) carboxylic acid (Harz, B 18, 3395) White needles V sol ether or petroleum-ether Yields on oxidation ethyl quinoline carboxylic acid (Daniel, B 23, 2267)

Salts — $B'HCl$ * small white concentric crystals — $B'HNO_3$ * fine white needles — $B'H_2SO_4$ * colourless crystals — $B'H_2Cl_2PtCl_4$ * fine orange needles, sol hot water, sparingly in cold — $B'_2C_6H_4(NO_2)_4OH$ [244°], very sparingly soluble yellow microscopic crystals

(B 4-Py 2) Di methyl-(Py 3)-ethyl quinol-

$C_8H_9(CH_3) \langle \begin{smallmatrix} CH \\ N \end{smallmatrix} \rangle \begin{smallmatrix} C(CH_3) \\ C(CH_3) \end{smallmatrix}$ [44°] (280° at

717 mm) Formed by the action of propionic aldehyde and HCl upon o-toluidine (Harz, B 18, 3400) Large colourless monoclinic crystals It is reduced by tin and HCl to a tetra hydride, which is a yellowish oil of boiling point 275° at 724 mm Yields methyl ethyl-quinoline (B 4)-carboxylic acid on oxidation

Salts — $B'HCl$ * easily soluble fine colour-

less needles — $B'HI^*$, very long colourless needles, ν sol. hot water, very sparingly in cold — $B'H_2Cl_2PtCl_4$, large plates — $B'C_6H_5(NO_2)_2OH$; [187°], nearly insoluble yellow needles

Methylo-iodide $B'MeI$ 2aq white needles
Gives $(B'MeCl)_2PtCl_4$, small glistening orange red needles

(*B* 1or3 *Py* 2) Di-methyl (*Py* 3) ethyl-quinoline $C_6H_5(CH_3)_2$ $\begin{matrix} \text{CH} \\ | \\ \text{C}(\text{CH}_3) \\ | \\ \text{N} \text{C}(\text{C}_6\text{H}_5) \end{matrix}$ [41°] (288°–

292° at 720 mm) Formed by the action of propionic aldehyde and HCl upon *m* toluidine (Harz, *B* 18, 3397) Colourless trimetric hexagonal plates It is reduced by tin and HCl to a tetrahydride, which is a colourless oil of boiling point 282°–285° at 720 mm

Salts — $B'HCl^*$ easily soluble colourless prisms — $B'HI^*$ long colourless needles, soluble in hot water, very sparingly soluble in cold — $B'H_2Cl_2PtCl_4$ 2aq orange plates or fine needles, sl sol cold water — $B'C_6H_5(NO_2)_2OH$ [220°], yellow needles, sol hot alcohol, nearly insol water

Methylo-iodide $B'MeI$ aq yellow needles, ν sol hot water, gives $(B'MeCl)_2PtCl_4$, small glistening orange red crystals

(*B* 2 *Py* 2) Di methyl (*Py* 3) ethyl quinoline $C_6H_5(CH_3)_2$ $\begin{matrix} \text{CH} \\ | \\ \text{C}(\text{CH}_3) \\ | \\ \text{N} \text{C}(\text{C}_6\text{H}_5) \end{matrix}$ [54°] (288° at

720 mm) Prepared by adding propionic aldehyde (60 g) to a cooled mixture of *p* toluidine (50 g) and strong HCl (90 g), finally heating for a short time on the water bath White trimetric crystals Sparingly volatile with steam ν sol alcohol, ether, and benzene, insol water It is reduced by tin and HCl to a tetrahydride (286°) It combines with bromine, forming a dibromide, which crystallises in yellow needles [91°] By CrO_3 and dilute H_2SO_4 it is oxidised to (*B* 2) methyl (*Py* 3) ethyl quinoline (*Py* 2) carboxylic acid

Salts — $B'HCl$ syrup, which slowly crystallises — $B'HBr$ easily soluble white silky needles — $B'HI$ long colourless needles, ν sol hot water, sparingly in cold — $B'H_2Cr_2O_7$, sparingly soluble red needles — $B'H_2Cl_2PtCl_4$ 2aq orange-red needles [208°] (Jungmann, *B* 23, 2273) — $B'C_6H_5(NO_2)_2OH$ [177°], small yellow needles

Methylo-iodide $B'MeI$ aq [218°], monoclinic yellow crystals

Methylo-chloride $B'MeCl$, colourless soluble needles — $(B'MeCl)_2PtCl_4$

Ethylo-iodide $B'EtI$ aq [114°], small yellow crystals — $(B'EtCl)_2PtCl_4$ aq glistening crystals (Harz, *B* 18, 3384)

References — AMDO, DI BROMO, NITRO, and OXY-METHYL ETHYL QUINOLINES

(*B* 2,4 *Py* 2)-Tri methyl (*Py* 3) ethyl-quinoline $C_6H_5(CH_3)_3$ $\begin{matrix} \text{CH} \\ | \\ \text{C}(\text{CH}_3) \\ | \\ \text{N} \text{C}(\text{C}_6\text{H}_5) \end{matrix}$ [62°] (291°)

Formed from (1,3,4) xylidine, propionic aldehyde and HCl (Waldbott, *B* 23, 2270) Triclinic tables (from alcohol), gives on oxidation (*B* 2, *Py* 3) di-methyl- (*Py* 3)-ethyl quinoline (*B* 4) carboxylic acid [183°] — $B'H_2PtCl_4$ — $B'HCl$ 3aq — $B'H_2SO_4$ Monoclinic crystals — $B'H_2Cr_2O_7$ — $B'INO$, Monoclinic crystals (from alcohol-

ether) — $B'C_6H_5(NO_2)_2OH$ [183°]. Yellow needles

Methylo-iodide $B'MeI$ Needles
(*Py* 2,4) Methyl ethyl-quinoline dihydride

C_6H_5 $\begin{matrix} \text{CH} \\ | \\ \text{C}(\text{CH}_3) \\ | \\ \text{N} \text{C}(\text{C}_6\text{H}_5) \end{matrix}$ (255°) Formed by heating methyl ethyl-indole with $MeOH$ and MeI for 15 hours at 120° (Fischer a Steche, *A* 242, 363) Oil, forming very soluble salts

Py (2,3) Methyl ethyl quinoline tetrahydride $C_{12}H_{17}N$ $\begin{matrix} \text{CH}_2 \\ | \\ \text{CH} \\ | \\ \text{NH} \text{CH} \text{Et} \end{matrix}$ (262° at 720

mm) Formed by reduction of (*Py* 2,3) methyl ethyl quinoline with tin and HCl (Doebner a Miller, *B* 17, 1716) Colourless liquid $FeCl_3$ gives a red colouration — $B'HCl$ sparingly soluble colourless concentric needles

(*Py* 3,4) - Methyl - ethyl - quinoline tetrahydride $C_{12}H_{17}N$ $\begin{matrix} \text{CH}_2 \\ | \\ \text{CH} \\ | \\ \text{NH} \text{CH} \text{Me} \end{matrix}$

(256°) Obtained by reducing (*Py* 3) methyl quinoline ethylo iodide with tin and hydrochloric acid (Moller, *A* 242, 321) Colourless liquid — $B'H_2PtCl_4$ Red granular crystals, decomposed by boiling water

Methylo-iodide $B'MeI$ [187°] Not acted on by KOH aq
(*Py* 2,3,4) Di-methyl-ethyl-quinoline di-

hydride $C_{12}H_{17}N$ $\begin{matrix} \text{CH} \\ | \\ \text{CH} \\ | \\ \text{NH} \text{CH} \text{Me} \end{matrix}$

(256° ν V) Formed by heating methyl ketole (methyl indole) with ethyl alcohol and ethyl iodide for 15 hours at 100° (E Fischer a Steche, *A* 242, 360, *B* 20, 2200) Oil The salts are ν sol water

Methylo-iodide $B'MeI$ [189°] Colourless crystals, sol water and alcohol

Di-methyl-ethyl-quinoline dihydride $C_{12}H_{17}N$ (255° ν V) at 750 mm Formed by heating methyl ethyl ketole (methyl ethyl indole) (1 pt) with MeI (2 pts) and $MeOH$ (1 pt) at 120° (F a S)

(*B* 4, *Py* 2) Di-methyl- (*Py* 3)-ethyl-quinoline tetrahydride $C_{12}H_{17}N$ $\begin{matrix} \text{CH}_2 \\ | \\ \text{CH} \\ | \\ \text{NH} \text{CH} \text{Et} \end{matrix}$ (275°)

at 724 mm Obtained by reducing the corresponding di methyl ethyl quinoline with tin and HCl (Harz, *B* 18, 3400) Yellowish oil, sparingly volatile with steam

(*B* 3? *Py* 2) Di-methyl- (*Py* 3)-ethyl-quinoline tetrahydride $C_{12}H_{17}N$ $\begin{matrix} \text{CH}_2 \\ | \\ \text{CH} \\ | \\ \text{NH} \text{CH} \text{Et} \end{matrix}$ (c 284°)

at 720 mm Obtained by reducing the corresponding di methyl ethyl quinoline with tin and HCl (Harz) Oil

(*B* 2, *Py* 2) Di-methyl- (*Py* 3)-ethyl-quinoline tetrahydride $C_{12}H_{17}N$ $\begin{matrix} \text{CH}_2 \\ | \\ \text{CH} \\ | \\ \text{NH} \text{CH} (\text{C}_6\text{H}_5) \end{matrix}$ (286°

at 720 mm) Formed by reduction of di methyl ethyl quinoline with tin and HCl (Harz, *B* 18, 3387) Colourless refractive fluid

Salts — $B'HCl$ colourless needles, sl sol cold water — $B'H_2Cl_2PtCl_4$ 2aq brown plates

Nitrosamine $C_{12}H_{17}N(NO)$ crystalline solid, sparingly volatile with steam

(B 2-Py 24). Tri-methyl-(Py 3)-ethyl-quinoline tetrahydride $C_8H_8(CH_3)_3$ $\begin{matrix} CH_2-CH(CH_3) \\ | \\ N(CH_3)_2CH(C_2H_5) \end{matrix}$

(275°-280° at 720 mm) Oil Formed by heating (B 2 Py 2) di methyl-(Py 3)-ethyl-quinoline tetrahydride with methyl iodide— $B_2H_5Cl, PtCl_2$ 2aq needles (Harz, B 18, 3388)

(B 24, Py 2) Tri methyl (Py 3) ethyl-quinoline tetrahydride C_8H_8N (288°) Formed by reducing the corresponding tri methyl ethyl-quinoline (Waldbohm, B 23, 2272) Oil— $B^*C_2H_5(NO_2)OH$ [146°]

(B 2). METHYL-(Py 3)-ETHYL-QUINOLINE CARBOXYLIC ACID

$C_8H_8(CH_3)_3$ $\begin{matrix} CH C(CO_2H) \\ | \\ N C(C_2H_5) \end{matrix}$ [143°] Formed by

oxidation of (B 2 Py 2) di methyl (Py 3) ethyl quinoline with CrO_3 and dilute H_2SO_4 (Harz, B 18, 3392, Daniel, B 23, 2266) Trichinic needles, prisms, or plates (containing aq) Sol water and alcohol Yields on distillation methyl ethyl quinoline [60°]

Salts— A^*Na 3aq easily soluble glistening crystals— A^*Ag granular powder— A^*Ba 1/2 aq soluble needles— A^*Cu microcrystalline blue pp

Ethyl ether EtA^*aq [170°-190°] Decomposed on fusion (Harz, B 18, 3394)

Methyl ethyl quinoline (B 4) carboxylic acid $CO_2H C_8H_8$ $\begin{matrix} CH CMe \\ | \\ N CEt \end{matrix}$ [216°] Formed by

oxidation of (B 4, Py 2) Di methyl (Py 3) ethyl quinoline (Miller & Daniel, B 23, 2266) Insol water sl sol cold alcohol Gives on distillation methyl ethyl quinoline [56°]— BaA^* 1/2 aq (B 2, Py 2) Di methyl (Py 3) ethyl quinoline (B 4) carboxylic acid $C_8H_8NO_2$ $\begin{matrix} CH CMe \\ | \\ N CEt \end{matrix}$ [183°] Formed by

oxidising (B 24, Py 2) tri methyl (Py 3) ethyl-quinoline by CrO_3 and H_2SO_4 (Jungmann, B 23, 2273) Needles (from alcohol), v e sol alcohol and water, sl sol benzene On dry distillation it gives CO_2 and (B 2, Py 2) di methyl (Py 3) ethyl quinoline (254°)

(B 2) METHYL-(Py 3)-ETHYL-QUINOLINE CARBOXYLIC ALDEHYDE C_8H_8NO $\begin{matrix} CH CCHO \\ | \\ N CEt \end{matrix}$ [57°] (above 300°)

Formed, in small quantity, in the preparation of the preceding acid (Harz, B 18, 3397) Large trimetric prisms (from ether), not very volatile with steam Reduces ammoniacal $AgNO_3$, Ag_2O oxidises it to the corresponding acid (Daniel, B 23, 2267)

(B 2 Py 2) DI METHYL (Py 3) ETHYL QUINOLINE SULPHONIC ACID $C_8H_8N(SO_3H)$ [above 290°] Formed by heating di methyl ethyl quinoline with fuming sulphuric acid (20 p.c. SO_3) (Harz, B 18, 3389) White silky scales or thick trimetric crystals Easily soluble in hot water, sparingly in cold, insol strong alcohol

Salts— A^*Ba aq easily soluble white needles— A^*H_2Pb 6aq easily soluble needles

METHYL-TRI ETHYL-RESORCIN

$OMeEt_3(OH)_2$ [144°] Obtained by heating with HCl its mono-ethyl derivative $C_6MeEt_2(OH)(OEt)$

which is formed, together with the di ethyl ether of orcin, by treating orcin with EtI and KOH (Herzig & Ziesel, M 11, 318) Needles, insol water, v sol ether Neutralises only 1 mol $NaOH$

Acetyl derivative $C_6MeEt_2(OAc)(OH)$ [c 73°] Monoclinic needles

METHYL TRI-ETHYL SILICATE $C_6H_8SiO_3$ $\begin{matrix} MeEt_3SiO_3 \end{matrix}$ (156°) SG 2 989 Formed from $MeO SiCl_3$ by the action of $EtOH$ (Friedel & Crafts, A Ch [4] 9, 32)

Di methyl di ethyl silicate $Me_2Et_2SiO_3$ (143°-146°) VD 6 18 (calc 6 23) SG 2 1 004 Formed from $MeOH$ and Et_2SiO_3 Formed also by the action of $EtOH$ on $(MeO)_2Si_2Cl_2$ Liquid

Tri methyl ethyl silicate Me_3EtSiO_3 (134°) SG 2 1 023 Formed from $(MeO)_3SiCl$ and $EtOH$

METHYL ETHYL STIBONIUM SALTS v AN TIMONY, vol 1 p 294

METHYL ETHYL SUCCINIC ACID

$C_6H_8O_4$ $\begin{matrix} CO_2H CHEt CHMe CO_2H \end{matrix}$ [168 5°] Formed by the action of H_2SO_4 on pentane tricarboxylic ether (Bischoff & Walden, B 22, 1817) It is accompanied by the isomeric 'meso' acid [88°] (Bischoff & Mintz, B 23, 647) The same acid [160°] appears to be formed in small quantities in the saponification of $CH_3CO Me(CO_2Et)CHEtCO_2Et$ with HCl (Young, C J 43, 180) White crystals

Di methyl ethyl succinic acid $C_6H_8O_4$ $\begin{matrix} CO_2H CHEt CMe CO_2H \end{matrix}$ Formed by the action of H_2SO_4 on the product of the action of a bromo isobutyric ether on sodium ethyl malonic ether (Bischoff & Mintz, B 23, 651) It occurs in two varieties, one melting at 63° the other at 105°

DI METHYL DI ETHYL-SULPHAMIDE

$C_6H_8N SO_2$ $\begin{matrix} NMe_2 \\ | \\ NEt_2 \end{matrix}$ (229°) Formed

by the action of di ethyl amine on di methyl amido sulphonic chloride, or of di methyl amine on di ethyl amido sulphonic chloride (Behrend, B 15, 1610, A 222, 125, 136) Volatile with steam Heavy oil Insol water, sol alcohol, ether, and benzene

METHYL ETHYL SULPHIDE C_6H_8S $\begin{matrix} MeSEt \end{matrix}$ (67° 1 V) SG 22 837 VD 2 61 (calc 2 63) Formed by distilling an alcoholic solution of $NaSEt$ with MeI (Kruger, J pr [2] 14, 193, Claesson, J pr [2] 15, 174, B 20, 8413) Occurs also among the products obtained by heating di thio phosphoric ether $Et_2PO_2S_2$ with $MeOH$ at 150° (Carus, A 119, 313, 120, 61) Liquid, with unpleasant odour Yields ethane sulphonic acid on oxidation with HNO_3 — $EtSMeHgCl$, laminae (C)— $(EtSMe)_2HgI_2$

Yellow crystalline powder
Methylo iodide $EtSMeI$ v DI METHYL-SULPHIDE ethylo-iodide
Ethylo iodide Et_2SMeI v Methylo compounds of DI ETHYL SULPHIDE, vol 11 p 516

METHYL-ETHYL SULPHONAMIDE v METHYL ETHYL-SULPHAMIDE

METHYL ETHYL SULPHONE $C_6H_8SO_2$ $\begin{matrix} MeEtSO_2 \end{matrix}$ [36°] Formed by oxidising $MeSEt$ with $KMnO_4$ (Beckmann, J pr [2] 17, 455)

Formed also by heating $C_2H_5SO_2CH_3$, SO_2 , CH_3CO_2H to 190° (Otto, B 21, 993) White needles (from ether) V sol water, alcohol, benzene, chloroform, and strong acids Sl sol CS_2 and ether Not affected by reducing agents

METHYL ETHYL SULPHONE CARBOXYLIC ACID $C_2H_5SO_3$ *see* Et $SO_3CH_2CO_2H$ *Ethylsulphono acetic acid* Obtained by saponifying its ether with cold conc KOH aq. Formed also by oxidising the barium salt of the ethyl derivative of thio glycolic acid with $KMnO_4$ (Claesson, *Bt* [2] 23, 447) Thick colourless syrup At 190° it splits up into CO_2 and methyl ethyl sulphone [36°] Bromine added to its aqueous solution gives a di bromo methyl ethyl sulphone Zinc and HCl reduce it to ethyl mercaptan

Salts — NaA' Pearly plates, v sl sol hot alcohol, v sol water — KA' Small tables (from alcohol) — BA'_2 Nodules — CUA'_2 2aq Broad tables

Ethyl ether EtA' Formed by the action of chloro acetic ether on sodium ethane sulphinate (Claesson, *J pr* [2] 15, 223, Otto, *B* 21, 993) Oil

METHYL ETHYL-THIAZOLE

$\begin{matrix} S & OEt \\ \diagup & \diagdown \\ CH & OMe \end{matrix} N$ (161° cor) Formed by the action of thiopropionamide on chloro acetone in alcoholic solution (Hubacher, *A* 259, 230) Liquid, smelling like pyridine, miscible with alcohol, insol hot water, sl sol cold water — $B'_2H_2PtCl_6$ [177°]

METHYL ETHYL DITHIOCARBONATE v

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METHYL ETHYL THIO UREA $C_2H_5N_2S$ *see*

NHMe CS NHEt [54°] Formed by addition of methylamine to ethyl thio carbimide (Hofmann, *B* 1, 27, *Z* 1868, 655) Fine crystals, sol water, alcohol, and HCl aq

Methyl di ethyl-thio- ψ urea $C_2H_5N_2S$ *see*

NHETC(SMe)NEt Formed from di ethyl thio urea and MeI (Noah, *B* 23, 2196) Oil With alcoholic NH_3 at 100° it yields di ethyl guanidine and MeSH — $B'HI$ — $B'_2C_2H_4(NO_2)_2OH$ [116°] — $B'_2H_2PtCl_6$ laminae

METHYL-ETHYL TOLUQUINOLINE v Di

METHYL-ETHYL-QUINOLINE

METHYL ETHYL UREA $C_2H_5N_2O$ *see*

NHMe CO NHET [53°] (267°) (Wurtz, *Rép chim pure*, 4, 199) [92°–112°] (Schreiner) Formed from methylamine and cyanic ether (Wurtz) When methylamine acts on ethyl carbamic ether there is formed a methyl ethyl urea which melts at 105° and solidifies again at 101° When, on the other hand, ethylamine acts on methyl carbamic ether the methyl ethyl urea which is produced melts at 75° and solidifies again at 72° (Schreiner, *J pr* [2] 22, 360). After frequent meltings and solidifications, both these forms begin to melt at 92° and end at 112°

METHYL-EUGETIC ACID v Methyl deriva-

tive of EUGETIC ACID

METHYL-ISO-FERULIC ACID v Iso FERULIC

ACID and the dimethyl derivative of CAFFEIC ACID, vol I p 659

METHYL-FLAVOLINIUM HYDRATE v

methyl hydrate of (Py 31) PHENYL METHYL-QUINOLINE (flavoline)

METHYL-FLUORESCIN v so-called 'fom-

fluorescein,' vol II p 568

METHYL FLUORIDE CH_3F VD 1 22

(calc 119) S 166 at 15° (D & P) Formed, together with Me_2O , by the action of KF on $KMeSO_4$ (Dumas & Péligot, *A* 15, 59) Formed also, in small quantity, together with NMe_3 , by

heating NMe_3F at 180° *in vacuo* (Lawson & Collie, *C J* 53, 628, 55, 110) Prepared by the action of MeI upon silver fluoride (Moissan & Meslans, *C R* 107, 1155) Gas, sl sol water, v sol alcohol and MeI Liquefied by a pressure of 30 atmospheres It burns with a blue flame, yielding HF Saponified with difficulty by heating in sealed tubes with water or dilute aqueous KOH at 120° In the presence of a little water it forms a crystalline hydrate, decomposing at 18.8° (Villard, *C R* 111, 184) Chlorine, acting upon it in sunlight, forms CH_2ClF , a gas which is decomposed by water, and is hardly inflammable

METHYL FORMAMIDE v Formamide in the article on FORMIC ACID

Di methyl formamidide $C_2H_5N_2$ *see*

NMe_2CHNH Form-imid di methyl amide

Formed by the action of an alcoholic solution of di methyl amine on the hydrochloride of formimido ether (Pinner, *B* 16, 1650) — $B'HI$ thick prisms, [169°], v sol water and alcohol

s Di methyl-formamidide $NHMeCHNMe$

Form methyl imid methyl amide Formed by the action of an alcoholic solution of methyl amine on the hydrochloride of formimido ether $NHCH OEt$ — $B'_2H_2Cl PtCl_6$ short red prisms, [172°] (Pinner, *B* 16, 358, 1648)

METHYL FORMANILIDE v FORMIC ACID

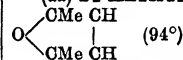
DI METHYL-FUMARIC ACID v DI METHYL

MALEIC ACID

METHYL-FURFURALDEHYDE C_5H_4O *see*

$C_5H_4MeOCHO$ Methyl furfural (187° i V) S 33 Appears to accompany furfuraldehyde in the product of the dry distillation of wood (Hill, *B* 22, 607) Formed also by distilling isodulcote with dilute H_2SO_4 (Maquenne, *C R* 109, 604) According to Bieler & Tollens (*A* 258, 110, *B* 22, 3062) fucosol or fucus aldehyde (*q v*) is a mixture of furfuraldehyde and methyl furfuraldehyde Combines with $NaHSO_3$ Aqueous ammonia converts it into crystalline methyl furfuralamide $N_2(C_5H_4O)_2$ [87°] Yields, on oxidation, methyl pyromucic acid $C_6H_4O_6$ [109°] Reduces Ag_2O Gives Schiff's reaction with isosamine and SO_2 Its phenyl hydrazide is liquid With resorcin and HCl it yields an orange yellow condensation product Pyrogallol and HCl form a crimson compound Paper moistened with aniline acetate is coloured yellow, and finally deep orange red by an aqueous solution of methyl furfuraldehyde

(aa) DI METHYL FURFURANE C_6H_4O *see*



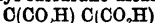
Formation — 1 By dry distillation of pyrotritic or carbopyrotritic acid, the yield being 5–7 p c — 2 By distilling acetonyl acetone $CH_3COCH_2CH_2COCH_3$ with $ZnCl_2$, the yield being 25 p c (Dietrich, *B* 20, 1085) — 3 Occurs also in the product obtained by distilling sugar (1 pt) with lime (3 pts) (E Fischer & Laycock, *B* 22, 101)

Properties — Colourless mobile very volatile liquid of characteristic odour Insol water, miscible with other solvents Does not react with phenyl-hydrazine. By PCl_5 or $POCl_3$ it is resinified By heating with acidified water it is converted back into acetonyl-acetone

METHYL-FURFURANE CARBOXYLIC ACID**METHYL PYROMUCIC ACID**

Di methyl-furfurane carboxylic acid v PYRO-TRITARIC ACID

Di methyl furfurane dicarboxylic acid



$\text{C}_6\text{H}_4\text{O}_4$, 1:2 \parallel (Knorr, cf Fittig,

B 18, 3410) *Carbopyrotritaric acid* [230°] Formed by boiling diacetyl succinic ether for a long time with dilute H_2SO_4 (Harrow, C J 33, 425, A 201, 152) Formed also by the action of fuming HCl on acetonyl aceto acetic ether $\text{CH}_3\text{COCH}_2\text{CH}(\text{CO}_2\text{Et})\text{COCH}_3$ (Paal, B 17, 2765)

Properties — Needles (by sublimation), m sol hot water, v sol alcohol and ether, volatile with steam On heating above its melting point it is decomposed into CO and di methyl furfurane carboxylic acid (pyrotritaric acid) Potash fusion yields acetic and succinic acids

Salts — K.A" silky needles — BaA" — Ag A" white pp HAgA" slender white needles

Mono methyl ether MeHA" [129°] Formed by leaving a mixture of the di methyl ether (1 pt) and fuming HClAq (10 pts) to stand in the summer time (K a C) Slender needles — AgMeA" Ppd by adding ammoniacal AgNO_3 to a solution of MeHA"

Di methyl ether MeA" [64°] (258°) From Ag A" and MeI at 100° under pressure (Knorr a Cavallo, B 22, 155) Snow white crystals, v sol alcohol, ether, chloroform, and benzene, v sl sol water, dilute acids, and alkalis

Methyl ethyl ether MeEtA" (268°) From AgMeA" and EtI, or from AgEtA" and MeI (K a C) The identity of the ethers piepared in these two ways shows that the carboxyls are similarly situated in the molecule of di methyl furfurane di carboxylic acid MeEtA" exhibits blue fluorescence With HCl it yields H A", and the ethers MeA" and EtA"

Mono ethyl ether EtHA" [83°] Formed by heating di acetyl succinic ether at 200°, or by allowing it to stand with conc HClAq for some hours in the cold (Knorr, B 17, 2463) Formed also from AgHA" and EtI (Harrow) Flat glistening needles, v sol alcohol and ether, sl sol water May be distilled AgEtA" Bulky white pp (Knorr a Cavallo, B 22, 154)

Di ethyl ether EtA" (275°) at 735 mm Formed by dissolving di acetyl succinic ether in cold H_2SO_4 , or by heating it with conc phosphoric acid (Knorr) Formed also, together with pyrotritaric and iso carbo pyrotritaric ethers, by heating diacetyl succinic ether at 175° (Knorr, B 22, 158) Oil Does not react with hydroxylamine or with phenyl hydrazine Easily saponified by alcoholic potash

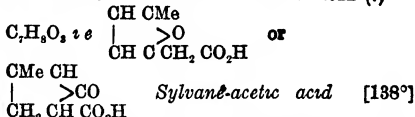
Di methyl-furfurane dicarboxylic acid v. METHRONIC ACID

METHYL FURFURINE

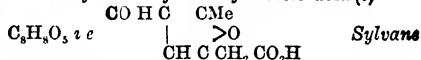
Formed by heating (5 g) methyl furfuramide (v METHYL FURFURALDEHYDE) with NaOH (2.5 g) dissolved in water (25 cc) at 100° (Bieler a Tollens, A 258, 123) Oil, which solidifies as an amorphous resin — B'HNO, lustrous crys-

als — B'H $_2$ SO $_4$ minute needles — B'H $_2$ PtCl $_4$, yellow crystalline pp

METHYL-FURFUROL v METHYL FURFUR-ALDEHYDE

METHYL-FURFURYL-ACETIC ACID (?)

Formed by heating its carboxylic acid (Polonowsky, A 246, 14) Prisms (from water), v e sol alcohol, m sol ether, almost insol cold water Somewhat volatile with steam It gives no colouration with FeCl_3 , and is dissolved without alteration by conc H_2SO_4 Nitric acid oxidises it, yielding acetic and oxalic acids — AgA' 3:4q. bulky white pp — BaA' 2:4aq transparent plates or spherical aggregates

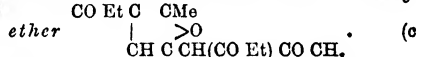
Methyl-carboxy-furfuryl-acetic acid (?)

carboxyacetic acid [207°] Formed, together with methyl furfuryl-carbinyl methyl ketone carboxylic ether and a neutral substance $\text{C}_{12}\text{H}_{18}\text{O}_4$ [139°], by adding a conc solution of ZnCl_2 to a mixture of glyoxal and aceto acetic ether in the cold (Polonowsky, A 246, 1) Needles (from hot water), v sol alcohol — (NH $_4$)A' 3:4q minute needles — BaA' 2:4aq transparent needles — AgA' 1:4q

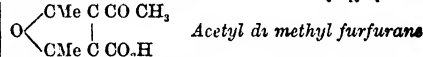
Mono methyl ether MeHA" [98°]: needles — AgMeA'

Di methyl ether MeA" Oil

Mono ethyl ether EtHA" [76°]

METHYL-FURFURYL-CARBINYL METHYL KETONE DI CARBOXYLIC ACID Ethyl

295°) One of the products formed by the condensation of aceto acetic ether (2 mol) with glyoxal (1 mol) in presence of aqueous ZnCl_2 (Polonowsky, A 246, 18) Light yellow oil, miscible with alcohol, ether, benzene, and chloroform Insol water and alkalis, resinsified by warming with conc H_2SO_4

DI METHYL - FURFURYL METHYL KETONE CARBOXYLIC ACID C₆H₁₀O₄ 1:2.

carboxylic acid [152°] Formed by heating at 225° in a sealed tube a mixture of β acetyl propionic acid (3 g) with Ac_2O (15 g), the product being freed from Ac_2O by distillation *in vacuo* and the residue extracted with water (Magnanim, B 21, 1523) Small needles (from hot water) — AgA' crystalline pp, sl sol water — BaA' 2:4q crystalline pp

DI - METHYL GENTISIC ALDEHYDE v. Di methyl derivative of Di oxy benzoic ALDEHYDE

METHYL GLUTACONIC ACID C₆H₈O₄ 1:2 $\text{CH}_2\text{CH}(\text{CO}_2\text{H})\text{CH}(\text{OH})\text{CO}_2\text{H}$ [187°] Formed by boiling methyl-dicarboxy-glutaconic ether (vol 1 p 706) with conc alcoholic potash (Conrad a Guthzeit, A 222, 259) Nodules (from water), v sol water, alcohol, and ether

α -METHYL GLUTARIC ACID $C_6H_{10}O_4$ *rac.*
 CO_2H $CHMe$ CH_2 CH_2 CO_2H *Butane dicarboxylic acid* Mol w 146 [77°] HF 238,200
 H C v 670,500 H C p 670,800 (Stohmann, Kleber & Langbein, *J pr* [2] 40, 214) Formed by saponifying with conc alcoholic KOH the product of the action of β iodo propionic ether on sodium methyl acetate ether (Wislizenus & Limpach, *A* 192, 133) Formed also by the action of HI and P on γ oxy α methyl glutaric acid (Krekeler, *B* 19, 3270), and on saccharonic acid CO_2H $CHMe(OH)$ $CH(OH)$ $CH(OH)$ CO_2H , a product of the decomposition of glucose (Kiliani, *A* 218, 369) Obtained by boiling γ cyano valeric acid (*q v*) with aqueous NaOH (Wislizenus, *A* 238, 101) Prisms, v sol water, alcohol, and ether The zinc salt is a viscid mass — Ag_2A''

β Methyl glutaric acid $C_6H_{10}O_4$ *rac*
 CH_3 $CH(CH_2CO_2H)_2$ *Ethylidene diacetic acid* [85°-86°] Formed from malonic acid, paraldehyde, and Ac_2O , by heating at 100° for several days On rectification the anhydride (282°-284°) is obtained (15 p c of weight of malonic acid). Boiling water changes it into the acid (Komnenos, *A* 218, 150) Colourless glass like prisms or tables (from CS_2 mixed with $CHCl_3$) V sol water, alcohol, or ether, m sol cold benzene or CS_2 , v sl sol CS_2 , or benzoline On distillation it splits up into H_2O and its anhydride On distilling its sodium salt with P_2S_5 , there is formed, as an oil, the homologue of thiophene $CH_2 \begin{smallmatrix} < CMe & CH \\ & CH & CH \end{smallmatrix} S$, boiling at 134° (Krekeler, *B* 19, 3270)

Salts — Ag_2A'' — CaA'' — PbA'' $\frac{1}{2}aq$ Trimetric needles $a b c = 6331$ 1 6072

Anhydride $C_6H_8O_3$ [46°] (282°-284°) V D 4.61 (calc 4.43) Mass of slender prisms (from CS_2) V sl sol cold water, neutral to litmus, on boiling it dissolves, becoming strongly acid, changing to the hydrated acid V sol alcohol, ether, benzene, chloroform, or glacial acetic acid, v sl sol petroleum

δ Di methyl glutaric acid $C_8H_{14}O_4$ *rac*
 CO_2H $CHMe$ CH_2 $CHMe$ CO_2H [114°] (*Z*), [c 90°] (*D*), [101°] (*B*), [106°] (*A* & *J*) Formed by treating a cyano-propionic ether with methylene iodide and alcoholic NaOEt, heating the mixture on the water bath, and saponifying the product with HCl (Zelinsky, *B* 22, 2823) Formed also by heating pentane tetracarboxylic acid either by itself, or in aqueous solution, or by heating its ether with H_2SO_4 (Dressel, *A* 256, 184) Obtained from sodium methyl malonic ether and methylene iodide (Bischoff, *B* 23, 1464, 1951) Crystals resembling those of di methyl-succinic acid, v sol water, alcohol, and ether Not volatile with steam According to Zelinsky the product obtained by his method really consisted of two isomeric modifications, melting at 102°-104° and 128° respectively A solution of the ammonium salt gives white pps with $AgNO_3$ and $HgCl_2$, and a brown pp with $FeCl_3$ — Ag_2A''

Anhydride $CH_2 \begin{smallmatrix} < CMe & CO \\ & CMe & CO \end{smallmatrix} O$ [93°]

Formed by boiling the acid for five minutes or by warming it with $AcCl$ (Auwers & Jackson, *B* 23, 1611) Rectangular prisms (from hot ligroin)

Tri methyl-glutaric acid $C_9H_{16}O_4$ *rac*
 CO_2H $CHMe$ CH_2 $CHMe$ CO_2H [97°] S 22 at

11° Formed, together with the isomeric tetramethyl-succinic acid, by the action of finely divided silver on a bromo isobutyric acid (Hell & Wittekind, *B* 7, 820, Auwers & V Meyer, *B* 23, 300) Flat plates (from hot water), sol cold water, m sol CS_2 and ligroin, v sol other solvents Can be distilled in small quantities Not volatile with steam When the acid (6 g) is mixed in the cold with dry red phosphorus (8 g) and bromine (16 g) the product is bromo trimethyl glutaric anhydride [114°]

Anhydride $CH_2 \begin{smallmatrix} < CMe & CO \\ & CMe & CO \end{smallmatrix} O$ [96°]

(262°) Formed by boiling the acid for some time, or by heating it with excess of Ac_2O Coarse flat satiny needles (from hot ligroin)

Ethyl ether Et_2A'' (230°) SG 2 1012

METHYL-GLYCERAMINE $C_6H_{11}NO_2$ *rac*
 $CH_2(OH)$ $CH(OH)$ CH_2NMeH Formed, together with the following body, by heating the (a) chlorhydrin of glycerin with aqueous NMe, in sealed tubes at 100° (Hanriot, *A Ch* [5] 17, 62, cf vol 11 p 623)

Tri-methyl-glyceramine chloride

$CH_2(OH)$ $CH(OH)$ CH_2NMe_3Cl Formed by heating glycerin chlorhydrin with trimethyl amine on the water bath for 12 hours (Hanriot, *C R* 86, 1335) — $C_6H_{11}ONMe_3Cl$ syrup — ($C_6H_{11}ONMe_3Cl$), $PtCl_4$, orange tables, sol water, insol alcohol, not decomposed by boiling the aqueous solution

DI-METHYL-GLYCERIC ACID *v Di oxy VALERIC ACID*

α METHYL-GLYCIDIC ACID $C_5H_8O_3$ *rac*

$O \begin{smallmatrix} < CH_2 \\ & CMe & CO_2H \end{smallmatrix}$ *Propylene oxide carboxylic*

acid Formed by decomposing chloro oxy iso butyric acid (the product of the union of $HOCl$ with methacrylic acid) with alcoholic potash (Mélkoff, *Bl* [2] 41, 311, 43, 115) Thick liquid, v sol water and ether

Reactions — 1 When heated with water for half an hour it forms di oxy butyric acid CH_2 $CH(OH)$ $CH(OH)$ CO_2H [100°] — 2 Heated with ammonia it forms oxy amido isobutyric acid — 3 Conc HCl combines, forming chloro oxy isobutyric acid [107°] — 4 HBr forms bromo oxy iso butyric acid [101°]

Salts — $KA' \frac{1}{2}aq$ glittering plates, sl sol cold alcohol — AgA' thin needles (from hot water)

Ethyl ether EtA' (164°) SG $\frac{1}{2}$ 10546 From AgA' and EtI (Mélkoff, *B* 21, 2053) Oil

β Methyl-glycidic acid $O \begin{smallmatrix} < CHMe \\ & CH & CO_2H \end{smallmatrix}$ [84°]

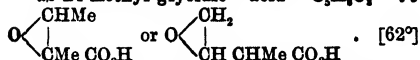
Formed by the action of alcoholic potash on chloro oxy butyric acid [63°] (the product of the union of $HOCl$ with crotonic acid) (Mélkoff, *Bl* [2] 43, 115) Trimetric crystals (from ether), v e sol water, m sol ether Volatile with steam

Reactions — 1 Combines with HCl forming chloro oxy butyric acid [86°] — 2 HBr yields bromo oxy butyric acid [90°] — 3 Ammonia forms oxy amido butyric acid — 4 When heated with water it slowly combines, forming di oxy-butyric acid [80°]

Salts — $KA' \frac{1}{2}aq$ powder, v sol water, sl sol cold alcohol — AgA' crystalline powder

Ethylether EtA' (174°) SG $\frac{1}{12}$ 1 0534 (Mélkoff a Zelnsky, B 21, 2052)

as Di-methyl-glycidic acid $C_3H_4O_2$ is



Formed by the action of alcoholic potash on chloro oxy valeric acid (the product of the union of HOCl on angelic acid) [45°] (Mélkoff, B 21, 47, 166, A 257, 118) Minute prisms, v sol water, alcohol, and ether Unites with HCl, forming chloro oxy valeric acid [75°] Water at 99° converts it into di oxy valeric acid [107°]

Salts—KA' $\frac{1}{2}$ aq prisms—AgA' thin scales

Ethylether EtA' (178°) SG $\frac{1}{12}$ 1 0250 From AgA' and F'I Oil

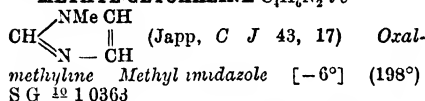
METHYL-GLYCOLLIC ACID v *Methyl derivative of GLYCOLLIC ACID*

DI-METHYL-GLYCOLURILE v *DI METHYL DIKETONE*

Tetra-methyl glycolurile v *GLYOXAL, Reaction 13*

METHYL-GLYOXAL v *PYRUVIC ALDEHYDE*
DI-METHYL-GLYOXAL v *DI METHYL DIKE*

METHYL-GLYOXALINE $C_4H_6N_2$ is

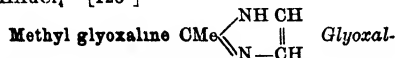


Formation—1 By the action of sodium amalgam on tri bromo methyl glyoxaline (g v) in alcoholic solution (Wyss, B 10, 1372)—2 By methylation of glyoxaline (Wallach, B 15, 644, A 214, 319)—3 From sulphhydro methyl glyox

aline $\text{HS} \begin{array}{c} \text{NMe CH} \\ \diagup \quad \diagdown \\ \text{N} - \text{CH} \end{array}$ by treatment with dilute nitric acid (Wohl a Marekwald, B 22, 1359)—4 By reducing chloro methyl glyoxaline with HIAq and P at 140° (Wallach, A 214, 310)

Properties—Liquid, sol water With aqueous HgCy₂, it gives a pp composed of slender needles [119°] Combines readily with MeI, forming a methylo iodide, whence B'Me₂PtCl₄ [206°] may be obtained

Salts—B'H PtCl₄ [191°] S 44 at 13° (W) 7.5 at 13° (W a M) Orange red prisms—B'H ZnCl₂ [128°–131°] Soluble crystals—B'HNO₃ Prisms—B'C₆H₅(NO₂), OH. [158°] Needles, sl sol alcohol and ether—B'HAuCl₄ [120°]



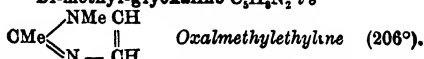
ethylene Para oxalmethylene [137°] (267°)

Formation—1 By saturating an aqueous solution of glyoxal (1 mol) and acetic aldehyde (1 mol) with ammonia (Radziszewsky, B 15, 2706, 16, 487)—2 By passing the preceding methyl glyoxaline [–6°] through a red hot tube (Wallach, B 16, 541)—3 By distilling the zinc salt of the preceding methyl glyoxaline or of chloro methyl ethyl glyoxaline with lime (Wallach, A 214, 296)

Properties—Long thin needles, v sol water, alcohol, and hot benzene, m sol cold benzene Decolourises bromine, forming C₂H₄Br₂ [258°] Yields oxamide on oxidation with H₂O₂ (Radzi-

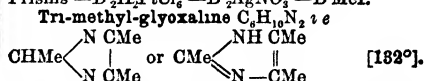
szewsky, B 17, 1290) Its solutions are ppd by tannin and by picric acid.

Salt—B'H₂PtCl₄ Trimetric needles
Di-methyl-glyoxaline $C_4H_6N_2$ is



SG $\frac{1}{12}$ 1 0051 Formed by the action of MeI on methyl-glyoxaline (Radziszewsky, B 16, 488). Liquid with narcotic odour, sol water, alcohol, and ether Its solution gives with CuSO₄ a blue pp, with AgNO₃ a crystalline pp, with HgCl₂ with tannin, and with phosphomolybdic acid white pps, and with picric acid a yellow pp All these pps are soluble in hot water—B'H₂PtCl₄ [138°]

Methylo iodide B'MeI Crystalline
Chloro di methyl - glyoxaline $C_4H_6ClN_2$ (213°) Formed by the action of PCl₅ on methyl-ethyl oxamide (Wallach, A 184, 71)—B'HCl Prisms—B'H₂PtCl₄—B'AgNO₃—B'MeI

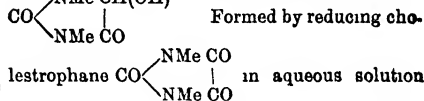


(271°) Formed by adding di methyl diketone CH₃COCOCH₃ to an ammoniacal solution of AgNO₃, and decomposing the resulting pp with dilute HCl (Fittig, A 249, 206) Formed also, with other bases, by heating di methyl diketone with conc NH₄Aq at 100° for an hour (Von Pechmann, B 21, 1411) Small white needles (from ether or ligron) Not affected by nitrous acid—B'HCl tufts of snow white needles

METHYL-GLYOXIM v *Oxim of PYRUVIC ALDEHYDE*

Di methyl glyoxim v *DI ACETYL and DI METHYL DIKETONE*

DI-METHYL-GLYOXYL-UREA $C_4H_8N_2O_3$ is



lestrophane $\begin{array}{c} \text{NMe CO} \\ \diagup \quad \diagdown \\ \text{CO} \quad \text{NMe CO} \end{array}$ in aqueous solution with zinc and H SO₄ (Andreassch, M 3, 436) Small needles, melting below 100° May be sublimed v e sol water and alcohol, insol ether Decomposed by boiling baryta water into CO₂, methylamine, and glyoxylic acid Readily re oxidised to cholestrophane

METHYL GUANACIL v *GUANIDINE, Reaction 11*

METHYL-GUANAMINE The name originally given by Nencki to *Acetoguanamine*, vol. II p 655

METHYL-DIGUANIDE $C_4H_6N_4$ Obtained, in combination with CuSO₄, by adding a 20 p c solution of methylamine to a mixture of di-cyan di amide C₂N₂(NH₂)₂, and copper sulphate, the crystalline (C₂H₅N₃)₂CuSO₄ 2 $\frac{1}{2}$ aq which separates is then decomposed by baryta (Reibenschuh, M 4, 388) Thick alkaline syrup, which absorbs CO₂ from the air

Salts—Colourless leaflets or prisms, v sol water—B'H₂SO₄ Prisms, obtained by decomposing B'₂CuSO₄ by H₂S—B'H₂SO₄ From B'H₂SO₄ and H₂SO₄ Crystalline powder v sl sol alcohol—Cu(C₂H₅N₃)₂ 3 $\frac{1}{2}$ aq Formed by shaking di cyan diamide with cupric oxide and methylamine Rose-red crystalline powder—

$B_2CuSO_4 \cdot 2\frac{1}{2}aq$ Pink needles — The chromate and picrate form yellow prisms

METHYL-GUANIDINE $C_2H_5N_3$, *re*
 $NH C(NH_2)(NHMe)$ *Methyl uramine*

Formation — 1 By boiling an aqueous solution of creatine with HgO or with PbO_2 and dilute H_2SO_4 (Dessaignes, *C R* 38, 839, 41, 1258, *A* 92, 407, 97, 340) — 2 By oxidising creatinin with $KMnO_4$ (Neubauer, *A* 119, 46) — 3 By heating cyanamide with methylamine hydrochloride in alcoholic solution at 100° (Erlenmeyer, *B* 3, 896) — 4 By the action of methyl cyanamide on NH_4Cl (Tavildaroff, *B* 5, 477)

Properties — Strongly alkaline, deliquescent, crystalline mass. Decomposed by KOH , giving off NH_3 and methylamine. When boiled with chloroacetic acid it yields glycolyl methylguanidine $C_4H_{11}N_3O_2$, which crystallises in tablets, *v* *e* sol water, and neutral in reaction, and forms the salts $C_4H_{11}NO_2 \cdot HCl$, and $C_4H_{11}N_3O_2 \cdot H_2PtCl_6$.

Salts — $B_2H_2PtCl_6$. Monoclinic crystals (Haushofer, *J* 1878, 351) *S* 143 at 19° — $B_2H_2AuCl_4$. Trimeric crystals, *v* sol ether, *m* sol water — $B_2H_2C_2O_4 \cdot 2aq$. Crystals, *v* sol water

Di-methyl-guanidine $C_3H_7N_3$, *re*
 $NH C(NH_2)(NMe)_2$ Formed by heating equimolecular quantities of cyanamide and dimethylamine hydrochloride at 110° (Tartaroff, *C R* 89, 608)

Salts — $B'HCl$. Trimeric crystals, *a b c* = 862 1 α (Haushofer, *Z K* 6, 130) — $B_2H_2PtCl_6$. Trichinic crystals, *a b c* = 941 1 678, $\alpha = 90^\circ 55'$, $\beta = 90^\circ 20'$, $\gamma = 90^\circ 4'$ — $B_2H_2AuCl_4$. Trimeric crystals (Haushofer, *J* 1882, 364)

s-Di-methyl-guanidine $NH C(NHMe)_2$. Formed by the action of methylamine on cyanogen iodide (Erlenmeyer, *B* 14, 1868)

Platinochloride $B_2H_2PtCl_6$. Trichinic crystals, *a b c* = 1212 1 1761, $\alpha = 90^\circ 22'$, $\beta = 110^\circ 20'$, $\gamma = 90^\circ 5'$

METHYL-GUANIDO-ACETIC ACID *v* **CREATINE**

(a) **METHYL-GUANIDO-BENZOIC ACID** $C_8H_{11}N_3O_2$, *re* $NH C(NH_2)NMe C_6H_4 CO_2H$

(a) *Benzcreatine*. Formed by methylation of benzglycocyamine (Griess, *B* 8, 324). Small acicular plates (containing $1\frac{1}{2}$ eq), *sl* sol hot water and alcohol. Decomposed by boiling baryta water into methyl amido benzoic acid and urea — $HA'HCl_{aq}$ plates, *m* sol water — $(HA')_2H_2PtCl_6 \cdot 2aq$

Anhydride *v* **BENZCREATININE**

(b) **Methyl-guanido-benzoic acid** $C_8H_{11}N_3O_2$, *re* $NH C(NHMe)NH C_6H_4 CO_2H$. Formed by the action of a cold concentrated solution of methylamine on the so called ethoxy carbumidamido benzoic acid (Griess, *B* 8, 325, vol 1 p 157). Plates, *sl* sol cold water and alcohol. Decomposed by baryta-water into *m* amido benzoic acid, CO_2 , and methylamine — $HA'HCl$ — $(HA')_2H_2PtCl_6 \cdot 2aq$

Anhydride *v* (b) **BENZCREATININE**

METHYL-GUANIDO-ETHANE SULPHONIC ACID $ON_2H_4Me(CH_2CH_2SO_3H)$ or probably $NH C(NHMe)NH CH_2CH_2SO_3H$. Formed by heating methyl amido ethane sulphonic acid (2 grms) and cyanamide (7 grms) with enough water to dissolve them, for 5 hours at 120° . Crystals of the new body are found in the tube (E. Dittrich, *J pr* [2] 18, 72). Transparent

monoclinic prisms. Crystallises with a molecule of water that goes off at 110° . *V* sol hot, *m* sol cold, water, *insol* alcohol and ether. Turns brown at 285° , giving off methylamine. Does not form salts.

METHYL-GUANIDO-VALERIC ACID *Anhydride* $C_6H_{11}N_3O_2$, *re*

$(CH_3)_2CH CH \langle \overset{NMe}{\underset{O}{C}} NH \rangle C NH$ *Isovaleric creatinine*. Formed by heating methyl-amido-isovaleric acid with cyanamide and ammonia (Duvillier, *Bl* [2] 39, 539). *Sol* alcohol

METHYL-HEMIPIC ACID *v* **Methyl ether of HEMIPIC ACID**

METHYL HENDECYL KETONE $C_{15}H_{32}O$, *re*. $CH_3 CO C_6H_{13}$, *Methyl undecyl ketone* [28°] (263°) *SG* (liquid) d_4^{25} 823. Formed by distilling a mixture of barium laurate and barium acetate (Krafft, *B* 12, 1667, 15, 1724). Yields acetic and hendecic acids on oxidation.

Isomeride $C_{15}H_{32}O$ (265° – 275°) *SG* d_4^{25} 887. One of the products of the action of CO on a mixture of $NaOAc$ and $NaOC_6H_{11}$ at 180° (Poetsch, *A* 218, 62). Liquid

METHYL HEPTADECYL KETONE $C_{17}H_{34}O$, *re*. $CH_3 CO C_6H_{13}$, [56°] (267° at 110 mm) *SG* (liquid) d_4^{25} 811. Prepared by distilling a mixture of barium stearate and barium acetate. On oxidation it gives margaric acid (Krafft, *B* 12, 1672, 15, 1724).

Methyl heptaacyl ketone $C_{15}H_{32}O$, *re*. $CH_3 CO CH(C_6H_{13})_2$. *Di octyl acetone* (325° – 330°). Formed by boiling di octyl acetoacetic ether with alcoholic potash (Guthzeit, *A* 204, 10). Liquid

METHYL HEPTINYL KETONE $C_{11}H_{20}O$, *re*. $CH_3 CO CH(C_6H_9)_2$. *Di allyl acetone* (175°). Formed by boiling di allyl acetoacetic ether with conc KOH_{aq} (Wolff, *A* 201, 48). *V* sol alcohol and ether

METHYL HEPTINYL OXIDE $C_9H_{16}O$, *re*. $CH_3 O CH(C_6H_9)_2$ (136°) *SG* d_4^{25} 8258. Formed from $NaOCH(C_6H_9)_2$ and MeI (Rabinin, *J pr* [2] 23, 270). Yields, on oxidation by cold $KMnO_4$, $CH_3 O CH(C_6H_9)_2$

METHYL HEPTYL KETONE $C_9H_{18}O$, *re*. $CH_3 CO CHPr_2$. *Di propyl acetone* (174°). Formed by heating di propyl acetoacetic ether with alcoholic potash (Burton, *Am* 3, 390). Liquid. Appears not to combine with $NaHSO_4$.

Methyl heptyl ketone $CH_3 CO C_6H_{13}$, (177°). Formed by the decomposition of *sec* hexyl acetoacetic ether by KOH (Lundahl, *B* 16, 789).

METHYL HEPTYL OXIDE $C_8H_{16}O$, *re*. $CH_3 O C_6H_{13}$, (161°) *SG* d_4^{25} 830 *VD* 42 (calc 45) *SV* 1946 (Lossen, *A* 254, 57). Formed by the action of MeI on $NaOC_6H_{13}$, derived from *canthol* (Wills, *C J* 6, 314). Mobile liquid, with strong odour, *insol* water, *v* sol alcohol and ether

METHYL-HESPERETIC ACID *v* **Dimethyl derivative of Caffeic acid**

METHYL-HEXADECYL-BENZENE *v* **HEXADECYL-TOLUENE**

Di-methyl-hexadecyl-benzene *v* **HEXADECYL-TYLENE**

Tri-methyl-hexadecyl-benzene *v* **HEXADECYL-MESITYLENE**

METHYL-HEXANE *v* **HEPTANE**

METHYL-HEXOSE $C_6H_{12}O_6$, *re*

$CH_2(OH)(CH_2OH)(CH_2OH)(CH_2OH)(CH_2OH)(CHO)CHO$

[181°] Formed by reducing the lactone of rhamnose carboxylic acid with sodium amalgam (Emil Fischer, *B* 23, 936) Crystallises from methyl alcohol Hydrogen cyanide yields $C_8H_{13}O_5N$, which on saponification forms methyl heptonic acid $C_8H_{15}O_6$, the lactone of which crystallises easily, and gives rise, on reduction, to methyl-heptose $CH_2(CH(OH)_2CHO$, of which the phenyl hydrazide is sl sol water

Phenyl-hydrazide V sol water

Oxazone [200° with decomposition]

METHYL HEXYL-CARBINOL *v* OCTYL ALCOHOL

DI METHYL HEXYLENE DIKETONE

$C_{16}H_{32}O_2$ *ie* $(CH_2CO)_2CH_2C_6H_{11}$ Iso amyl acetyl acetone (c 222°) Formed by heating $(CH_3CO)_2CHNa$ with isoamyl iodide at 180° (Combes, *A Ch* [6] 12, 249) Liquid Decomposed by caustic potash into acetic acid and $CH_3COCH_2C_6H_{11}$

Di methyl hexylene diketone $C_{16}H_{32}O_2$ *ie* $CH_3COCH_2CH_2CH_2CH_2CHMeCOCH_3$ a methyl *ae-di* acetyl pentane (232°-235°) Formed by decomposing its carboxylic ether with alcoholic potash (Kipping a Perkin, *C J* 55, 346) Colourless mobile oil Dissolves in a solution of $NaHSO_3$, and is reprecipitated by K_2CO_3

DI METHYL HEXYLENE DIKETONE

CARBOXYLIC ETHER $C_{16}H_{32}O_4$ *ie* $CH_3COCHMe(CO_2Et)CH_2CH_2CH_2CH_2COCH_3$ (255°-260°) Formed by the action of $NaOEt$ and $BrCH_2CH_2CH_2CH_2COCH_3$ on methyl acetoacetic ether (Kipping a Perkin, *C J* 55, 346) Colourless liquid, with faint odour, v sl sol water

METHYL-HEXYL-GLYOXALINE $C_{16}H_{32}N_2$

ie $C_6H_{13}CH \begin{matrix} \diagup NMech \\ | \\ \diagdown N=CH \end{matrix}$ (262°) SG 155 923

Formed by heating hexyl glyoxaline with a solution of MeI in methyl alcohol (Karez, *M* 8, 221) Colourless oil, insol water, sol alcohol and ether Gives the alkaloidal reactions — $B'H.PtCl_4$, yellow plates, sol water

Methyl iodide $B'MeI$ [124°] V sol alcohol and ether

METHYL *n* HEXYL KETONE $C_{16}H_{32}O$ *ie*

$CH_3COCH_2CH_2CH_2CH_2CH_2CH_2CO$ (172°) SG 29 8185 (Bruhl, *A* 203, 29), d_4^{20} 835 (C) SV 186 6 (R. Schiff, *A* 220, 103) $\mu_D = 1.4213$ R_{∞} 63 29 H C 1,209,510 (Lougumme, *Bl* [2] 41, 389)

Formation — 1 By distilling sodium ricinoleate with $NaOH$ (Lumprecht, *A* 93, 242, Bouas, *A* 97, 34) — 2 By distilling a mixture of calcium acetate and calcium *n* heptate (enanthe) (Stadeler, *J pr* 72, 246) — 3 From octoic aldehyde by treatment with PCl_5 , decomposing the resulting $C_8H_{17}CHCl_2$ by alcoholic KOH , and adding the octene C_8H_{16} , CCH so formed to cold H_2SO_4 , diluting and distilling with water (Béhal, *Bl* [2] 47, 39, *A Ch* [6] 15, 275) — 4 Together with ethyl amyl ketone by treating $C_8H_{17}CCH_2$ with H_2SO_4 and water successively (Béhal, *Bl* [2] 48, 704) — 5 'Oil of wine' contains a methyl hexyl ketone (164°), which yields hexoic and acetic acids on oxidation (Hartwig, Scholl, *J pr.* [2] 23, 449) — 6 By heating octyl alcohol with boric acid at 170°, and distilling *in vacuo* (Councler, *B* 11, 1108)

Properties — Colourless, mobile liquid, smelling like apples and tasting like camphor, sol alcohol and ether, insol water Combines with $NaHSO_3$, forming a compound sol alcohol, but decomposed by hot water Does not reduce ammoniacal $AgNO_3$ or Fehling's solution Chromic acid mixture oxidises it to acetic and hexoic acids (Béhal a Combes) Nitric acid oxidises it to heptonic acid (Petersen, *A* 118, 78) PCl_5 forms $C_8H_{17}Cl_2$ (190°-200°) (Dachauer, *A* 106, 271)

Combinations — $C_8H_{17}OKHSO_3$ $\frac{1}{2}$ aq — $C_8H_{17}ONH_2HSO_3$ V sol water (L)

Oxim $CH_3C(NO)H_2C_6H_{11}$ (214°) at 725 mm (*B* 21, 509, cf Béhal, *Bl* [2] 47, 163)

Methyl isohexyl ketone $CH_3COCH_2C_6H_{11}$ Isoamyl acetone (170°) Formed by the action of KOH upon di methyl hexylene diketone $(CH_3CO)_2CH_2C_6H_{11}$ (Combes, *A Ch* [6] 12, 249) Limpid liquid, with agreeable odour Combines with $NaHSO_3$

Methyl hexyl ketone (?) $MeCO C_6H_{13}$ (208°-210° cor) SG 15 843 Occurs in small quantity (40 g) among the products of the passage of CO over a mixture of sodic acetate (546 g) and sodic iso amylate (746 g) (Poetsch, *A* 218, 60)

Methyl hexyl diketone $C_8H_{17}O_2$ *ie* $CH_3COCH_2COCH_2CH_2Pr$ (165°) (Von Pechmann, *B* 21, 2140)

DI-METHYL-HEXYL-PYRIDINE $C_{15}H_{21}N$

ie $C_6H_5 \cdot C \begin{matrix} \diagup CHCMe \\ | \\ \diagdown CHCMe \end{matrix} N$ *n* Hexyl lutidine (250°) at 719 mm Obtained by distilling its dicarboxylic acid with lime (Jaekle, *A* 246, 41) Colourless liquid, with faint blue fluorescence — $B'H.PtCl_4$ [163°] Orange prisms — $B'AgNO_3$ [65°] White needles, sol hot water and alcohol

Hexahydrate $C_{15}H_{21}N \cdot 6H_2O$

$C_6H_5 \cdot CH \begin{matrix} \diagup CHMe \\ | \\ \diagdown CHNH \end{matrix}$ *Hexyl lupetidine*

(240°) at 715 mm Obtained by the action of sodium amalgam on an alcoholic solution of di methyl hexyl pyridine (J) Colourless oil, not exhibiting fluorescence Its dilute alcoholic solution gives a dark brown pp, with mercurous nitrate Its platinumchloride and dichromate could not be obtained in a crystalline state — $B'HCl$ long white prisms, v sl sol water

DI-METHYL-HEXYL-PYRIDINE DICARB-

OXYLIC ACID $C_{15}H_{21}NO_4$ *ie*

$C_6H_5 \cdot C \begin{matrix} \diagup C(CO_2H)CMe \\ | \\ \diagdown C(CO_2H)CMe \end{matrix} N$ *Hexyl lutidine dicarboxylic acid* The potassium salt is formed by saponifying its ether with boiling KOH aq (Jaekle, *A* 246, 40) The free acid has not been prepared — PbA'' $\frac{1}{2}$ aq

Di ethyl ether Et_2A'' Formed by oxidising its dihydride with nitrous acid Yellowish-brown oil, forming salts with mineral acids — $H_2A''H_2PtCl_4$ [141°] Orange plates

Dihydrate *Di ethyl ether*

$C_6H_5 \cdot CH \begin{matrix} \diagup C(CO_2Et)CMe \\ | \\ \diagdown C(CO_2Et)CMe \end{matrix} NH$ [54°] Formed by the condensation of heptonic aldehyde (enanthal) (1 mol) with acetoacetic ether (2 mols) and ammonia (1 mol) (J) Yellow prisms, crystallising with difficulty from a dilute alcoholic solution

DI-METHYL-HOMO-CAFFEIC ACID *v* *Di-methyl derivatives of* (4.3.1) **DI OXY-PHENYL-CROTONIC ACID**

METHYL-HOMO FERULIC ACID *v* *Di-methyl derivative of (431) Di-oxy phenyl-cro-tonic acid*

METHYL-HYDANTOIC ACID $C_4H_5N_2O_4$ *v* $NH_2 \cdot CO \cdot NMe \cdot CH_2 \cdot CO_2H$ *Methyl uramido acetic acid*

Formation—1 Occurs in the urine of a dog whose food is mixed with methyl amido acetic acid (Schultzen, *B* 5, 578)—2 By digesting an aqueous solution of sarcosine with potassium cyanate and ammonium sulphate for two days at 40° (Baumann a Hoppe Seyler, *B* 7, 35)—3 By boiling methyl amido acetic acid with urea and excess of baryta-water, or by digesting the same mixture for two days at 40° (*B* a *H*)—4 By the action of potassium cyanate and H_2SO_4 on methyl amido acetic acid (*E* Salkowsky, *B* 7, 116)

Properties—Transparent plates (from alcohol), m sol cold water and cold alcohol, v sol hot water and hot alcohol. Acid in reaction. When the concentrated aqueous solution is boiled it is partially converted into methyl-hydantoin, this dehydration is more rapidly effected by boiling with water and $PbCO_3$ or $BaCO_3$, only traces of the acid then passing into solution. When heated in a sealed tube with a saturated solution of baryta at 140° it yields NH_3 , methyl amido acetic acid, and CO_2 . Moist Ag_2O forms laminae of silver methyl hydantoin. Salts— BaA_2 Ppd by adding alcohol to its aqueous solution— CuA_2 . Amorphous.

Di methyl hydantoinic acid *v* **ACETONURAMIC ACID**

METHYL-HYDANTOIN $C_4H_5N_2O_4$ *v* $CO \begin{smallmatrix} \text{NMe} \cdot CH_2 \\ \text{NH} \cdot CO \end{smallmatrix}$ *Anhydride of methyl uramido acetic acid* Mol w 114 [182°] (*F* a *K*), [156°] (*E* Salkowsky, *B* 7, 119), [145°] (*N*)

Formation—1 By heating creatinin with baryta water at 100° in a sealed tube (Neubauer, *A* 137, 288)—2 By boiling methyl hydantoinic acid with water and $PbCO_3$ or $BaCO_3$ —3 By melting methyl-amido acetic acid with urea (Huppert, *B* 6, 1278)—4 By the action of cyanogen chloride on melted methyl amido-acetic acid (Traube, *B* 15, 2110)—5 By warming caffeic acid with baryta water (Fischer, *A* 215, 286)—6 By the reduction of methyl allantoin by HI (Hill, *B* 9, 1091)—7 By heating a mixture of hydantoin (3 pts), MeI (6 pts), methyl alcohol (16 pts), and KOH (2 pts), for three hours at 100°, and extracting the product with boiling alcohol (Franchimont a Klobbie, *R T C* 8, 289)

Properties—Short prisms, v sol boiling water, sl sol cold water, v sol alcohol, v sl sol ether. May be sublimed. It does not unite with baryta, but its hot solution dissolves Ag_2O , and the alkaline filtrate then deposits $C_4H_5AgN_2O_4$ as groups of thin laminae. The mercuric compound, obtained in like manner, forms nodular groups of minute needles, very soluble in water. Nitric acid (*S G* 15) converts methyl hydantoin into the nitramine

$CO \begin{smallmatrix} \text{NMe} \cdot CH_2 \\ \text{N}(NO_2) \cdot CO \end{smallmatrix}$ which crystallises from alcohol in scales [168°], v sl sol cold water, decomposed by boiling water

(a)-Methyl-hydantoin *v* **LACTYL-UREA.**

Di-methyl hydantoin $C_4H_5N_2O_4$ *v*

$CO \begin{smallmatrix} \text{NH} \cdot CMe_2 \\ \text{NH} \cdot CO \end{smallmatrix}$ [175°] Formed by slowly

adding HCl to commercial potassium cyanide (containing cyanate) covered by acetone. The liquid, after the action is completed, is left to evaporate, and the crystals which separate extracted with ether, and finally sublimed (Urech, *A* 164, 264). Large prisms, v sol water, alcohol, and ether. Slowly converted by boiling baryta water into acetouramic acid $NH_2 \cdot CO \cdot NH \cdot CMe_2 \cdot CO_2H$. By heating in a sealed tube with fuming hydrochloric acid at 160° it is resolved into CO_2 , ammonia, and a amido isobutyric acid $CMe_2(NH_2) \cdot CO_2H = C_4H_5AgN_2O_4$ crystalline powder, sl sol hot water— $C_4H_5N_2O_4 \cdot AgNO_3$ large prisms, v sol water

METHYL-HYDANTOIN CARBOXYLIC ACID $C_5H_5N_2O_5$ Formed by boiling caffeic acid with baryta water (*E* Fischer, *A* 215, 286). The free acid is decomposed into CO_2 and methyl-hydantoin when its aqueous solution is warmed

METHYL HYDRASTINE $C_7H_{11}NO_6$ *v* $CH_3O \cdot C \cdot C(OMe) \cdot C \cdot CO_2O \cdot CH \cdot C_6H_4(O_2CH_3) \cdot CH$
 $HC \cdot CH = C \cdot C = CH \cdot C \cdot C(C_6H_4NMe_2) \cdot CH$

[156°] Formed, together with hydrastine methylo hydroxide, by the action of alkalis on a solution of hydrastine methylo chloride. Prepared by adding potash to a hot aqueous solution of hydrastine methylo iodide until no further pptn takes place (Freund a Rosenberg, *B* 23, 406). Small yellow needles (from alcohol), almost insol water, sol chloroform, benzene, CS_2 , ether, and dilute alcohol. With sulphomolybdic acid (Froehde's reagent) it gives a violet colour, changing through blue to green. Forms sparingly soluble double salts with $SnCl_4$, zinc chloride, and $HgCl_2$. Ammonia converts it into the amide $C_{10}H_{15}N_2O_6$ [180°] which forms a crystalline hydrochloride $C_{10}H_{15}N_2O_6 \cdot HCl$ 2aq, and on boiling with dilute nitric acid yields hemipic acid (Freund a Heim, *B* 23, 2902). The formula of this amide may be written

$C_6H_2(OMe)_2CONH_2$

$CO \cdot CH_2 \cdot C_6H_2(O_2CH_3) \cdot CH_2 \cdot CH_2 \cdot NMe_2$

and it may also be got by heating hydrastine methylo iodide with alcoholic ammonia. The corresponding methylamide $C_{10}H_{15}N_2O_6$ [182°], ethylamide $C_{12}H_{19}N_2O_6$ [162°], allylamide $C_{12}H_{19}N_2O_6$ [158°], and amylamide $C_{14}H_{23}N_2O_6$ [171°] have been prepared

Salts— $B'HCl$ [241°] M sol hot water. Decomposed on fusion. A 1 p c solution is in active— $B'_2H_2PtCl_6$ sl sol water— $B'H \cdot SO_4$ [250°] Less soluble than the chloride— $B' \cdot HNO_3$ Decomposes at 250° V sl sol water

Methylo-iodide $C_{12}H_{19}NO_6 \cdot MeI$ Yellow needles, more soluble in water than in alcohol. Decomposes at 250° On warming with aqueous KOH it gives off trimethylamine

Methyl-hydrastimide $C_{12}H_{19}N_2O_6$ *v*

$C_6H_2(OMe)_2 \begin{smallmatrix} \text{CO} \cdot NH \\ \text{CO} \cdot NH \end{smallmatrix} \cdot C \cdot CH \cdot C_6H_2(O_2CH_3) \cdot CH_2 \cdot CH_2 \cdot NMe_2$ [192°] Formed by the action of dilute H_2SO_4 on the amide (Freund a Heim, *B* 23, 2899). Yellow needles, insol water, sl sol alcohol— $B'HCl$ [227°]— $B'HCl$ 2aq.— $B'_2H_2PtCl_6$ [205°].

— BH_2SO_4 , [218°] Yellow crystals (from alcohol) — $BHNO_3$ aq

Methylol-iodide $BMeI$ aq [245°]

Methyl hydrastine $C_{22}H_{27}NO_3$, [161°] Formed by warming hydrastine methyl-chloride with excess of KOH aq, or, better, by warming methyl-hydrastine with strong caustic potash (Freund a Rosenberg, *B* 23, 408) Colourless needles (containing aq) (from water), sl sol cold water, more sol hot water and alcohol Decomposed by long heating at 110° Very soluble in alkalis, but reprecipitated by CO_2 , insoluble in aqueous Na_2CO_3 It dissolves in aqueous NH_3 , but is precipitated on evaporation It also dissolves in HCl aq In a 5 p.c. hydrochloric acid solution it is inactive Tertiary base Coloured reddish brown by chlorine water and ammonia Iodine has no action Hot conc HBr aq reconverts it into methyl hydrastine Its salts are colourless — $BHCl$ aq [290°] Compact groups of crystals

Methyl hydrastinine *Methylol iodide* $C_{13}H_{19}NO$, I *re* CHO C_6H_5O , CH_3 , CH_2 , NMe , I [267°] Formed by heating hydrastinine with MeI (Freund, *B* 22, 2329) Pale yellow needles (from alcohol or water) $AgCl$ gives $C_{13}H_{19}NO_2Cl$ [213°], which forms white crystals (containing aq) — $(C_{13}H_{19}NO_2Cl)_2PtCl_2$, yellow crystalline pp

Oxime of the methylol iodide

$HO-N-CH(C_6H_5O)_2NMe$, I Yellowish needles, decomposed at 250° V sol alkalis and precipitated unchanged by acids Gives a dark reddish brown pp with platinum chloride

METHYL HYDRAZINE CN_2H_5 , *re* CH_3NH-NH (87° i.v.) at 745 mm Formed by treating methyl urea nitrate with $NaNO$, reducing the resulting nitroso methyl urea [124°] with zinc and dilute acetic acid, and decomposing the product with HCl (Brunner, *B* 21, 1810, *A* 253, 7) Mobile liquid with strong ammoniacal odour, soluble in water (heat being evolved), miscible with alcohol and ether Reduces Fehling's solution

Salts — BH_2SO_4 , [139.5°] White needles v sol water, sl sol alcohol — $BHCl$ v sol water and alcohol — $B_2C_6H_5(NO)_2O_2H$ [162°] Yellow needles, decomposed on fusion

Di benzoyl derivative $CH_3N_2HBz_2$, [c 143°] Colourless needles (from water), m sol hot water, v sol alcohol, sl sol ether Does not reduce Fehling's solution

Picryl derivative $CH_3N_2H_2C_6H_2(NO_2)_3$, [171°] Obtained from the hydrazine and picryl chloride (chloro tri nitro benzene) in diluted alcoholic solution Yellow plates, v sol alcohol and ether, m sol chloroform Decomposed on fusion

Oxalyl derivative $CH_3N_2H_2CO-CO-N_2H_2CH_3$, [221.5°] Formed by adding an aqueous solution of methyl hydrazine to oxalic ether White needles, v sol alcohol, m sol water, v sl sol ether Sublimes at about 160° in needles Reduces Fehling's solution Nitrous acid forms the nitrosamine $CH_3N_2H(NO)CO-CO-N_2H_2CH_3$, [147°]

Di methyl-hydrazine $C_2H_5N_2$, *re* NMe , NH_2 , (63°) at 720 mm SG Δ 801 VD ($H=1$) 30 Prepared by reducing di methyl nitrosamine $(CH_3)_2NNO$ with zinc dust and acetic acid (Fischer, *B* 9, 111, Renout, *B* 13, 2171) Formed also by reducing di methyl nitramine

$(CH_3)_2NNO$, (Franchimont, *R T C* 2, 123) Light volatile liquid with ammoniacal odour, v sol water, alcohol, and ether Its haloid salts volatilise without decomposition

Reactions — 1 CS_2 forms di methyl thiocarbazoic acid NMe , NH CS_2H — 2 *Phenyl thiocarbimide* (phenyl mustard oil) forms C_6H_5NH CS NH NMe — 3 *Oxalic ether* produces NMe , NH CO CO NH NMe — 4 *Alkyl iodides* unite, forming azonium iodides, such as NH , NMe , I — 5 $K_2S_2O_8$ forms NMe , NH SO_3K , which crystallises in white plates, v sol water It is split up by hot HCl aq into dimethyl hydrazine and H_2SO_4 — 6 HgO oxidises it, forming tetra methyl tetrazone NMe , N N NMe , a yellow oil (130°) which explodes when heated above its boiling point It forms a mirror with ammoniacal $AgNO_3$, even in the cold, and is decomposed by boiling dilute acids into formic aldehyde, NH_3 , Me , nitrogen, and $NHMe$ The tetrazone is a strong base — 7 *Nitrous acid* decomposes it into N_2O and dimethylamine — 8 *Acetophenone* at 100° forms NMe , N $CMepH$, a liquid (165° at 100 mm) It is split up again by acids into its generators (Reisenegger, *B* 16, 663)

Salts — $BHCl$ — BH_2Cl , hygroscopic crystals — $B_2H_2PtCl_6$, orange yellow prisms, v sol water, sl sol alcohol — $B_2H_2SO_4$, [105°] White needles — $B_2H_2CO_4$, colourless plates, v sol water and alcohol, sl sol ether

Ethyl chloride NH , NMe , $EtCl$ Crystallises with difficulty, v sol water The ethyl hydroxide is reduced by zinc dust and acetic acid to HCl , ammonia, and NMe , Et — $(NH_2NMeEtCl)_2PtCl_2$, Crystals

DI-METHYL-DI-HYDRO ANTHRACENE v **DI-METHYL ANTHRACENE DI-HYDRIDE**

METHYL HYDRO-ETHYL PYRIDINE v **METHYL FHTYL PYRIDINE HYDRIDE**

DI-METHYL-HYDRO-HOMO-CAFFEIC ACID v *Di methyl derivative of* (4 3 1) *Di oxy phenyl iso butyric acid*

METHYL-HYDRO-HOMO-FERULIC ACID v *Di methyl derivative of* (4 3 1) *Di oxy phenyl iso butyric acid*

METHYL DI-HYDRO PARVOLINE v *PFNTA* **METHYL PYRIDINE DIHYDRIDE**

METHYL HYDRO-PYRIDINES v **METHYL PYRIDINE HYDRIDES**

METHYL-HYDRO-QUINALDINE v (*Py* 3 4) *Di methyl quinoline tetra hydride*

METHYL HYDRO-QUINOLINES v **METHYL QUINOLINE HYDRIDES**

DI-METHYL HYDROQUINONE v *Di-methyl derivative of* **HYDROQUINONE**

DI-METHYL HYDROQUINONE TRI-METHYL AMMONIUM IODIDE v *Methylol iodide of the di methyl derivative of* **DI-METHYL-AMINO HYDROQUINONE**

METHYL HYDRO TOLUQUINONE v *Di-methyl derivative of* **HYDROTOLUQUINONE**

METHYL HYDROXYLAMINE v *Alkyl derivatives of* **HYDROXYLAMINE**

METHYL HYPOPHOSPHATE $Me_2P_2O_5$, SG, Δ 1109 From MeI and $Ag_2P_2O_5$ (Sanger, *A* 232, 11) Saponified by water

METHYL-HYPOPHOSPHATE of calcium $MeCaH_2P_2O_5$, 6aq Needles

METHYLIA. A name formerly used for **METHYLAMINE**.

METHYL-IMESATIN Described as *Imide* of *Methyl-ismatin* under ISATIN

METHYL-INDAZOLE A name employed by Wohl and Marckwald (*B* 22, 1859) to denote the substance usually called **METHYL-GLYOXALINE**
METHYL-INDAZYL MERCAPTAN *v*
SULPHYDRO-METHYL GLYOXALINE

METHYL-IMIDO-DI-METHYL-THIAZOLE $S \langle \text{C(NMe)} \rangle \text{NMe}$ [96°] Formed by the action of chloro acetone on di methyl urea (Traumann, *A* 249, 49) White needles (from water), with strong alkaline reaction Is identical with di methyl amido methyl thiazole

METHYL-IMIDO DI (β) NAPHTHYL SULPHIDE $\text{NMe} \langle \text{C}_{10}\text{H}_7 \rangle \text{S}$ [285°] Formed by heating imido di naphthyl sulphide (10 g) with MeI (5 g) and MeOH (12 c c) for five hours at 150° Formed also by heating di-(β) naphthyl-methyl amine with sulphur (Kym, *B* 23, 2459) Lemon-yellow plates or needles Gives a blue colour with H_2SO_4 containing HNO_3

METHYL-IMIDO PHENYL NAPHTHYL SULPHIDE $\text{NMe} \langle \text{C}_6\text{H}_5 \rangle \text{S}$ [133°] Formed by heating $\text{NH} \langle \text{C}_{10}\text{H}_7 \rangle \text{S}$ with MeI and MeOH at 150° (Kym, *B* 23, 2466) Light greenish-yellow needles, sl sol hot spirit, v e sol hot benzene

METHYL IMIDO DI PHENYL SULPHIDE $\text{C}_6\text{H}_5\text{NS} \text{ v e } S \langle \text{C}_6\text{H}_5 \rangle \text{NMe}$ [99.3°] (c 363°) Formed by heating imido di phenyl sulphide (thiodiphenylamine) with MeI and MeOH in sealed tubes at 110° (Bernthsen, *A* 230, 88, *B* 16, 2899) Long white prisms (from alcohol), insol water, v sol ether, benzene, and hot HOAc Its alcoholic solution is coloured brown by FeCl_3 Nitric acid forms a yellow nitro derivative which, unlike that of imido di phenyl sulphide, is insol NaOHAq The nitro derivative may be reduced to an amido compound which gives a deep bluish green colour with FeCl_3

Isomeride $\text{C}_6\text{H}_5\text{NS} \text{ v e } S \langle \text{C}_6\text{H}_5 \rangle_2\text{NMe}$ or $\text{C}_6\text{H}_5\text{N} \langle \text{C}_6\text{H}_5 \rangle \text{S} (?)$ [79°] Formed by the action of SCl_2 dissolved in petroleum on di phenyl methyl amine NMePh_2 (Holzmann, *B* 21, 2065) Thin yellow scales (from hot benzene alcohol), insol water, sl sol hot alcohol and ether, v sol hot benzene On heating with copper it yields di phenyl methyl amine

METHYL IMIDO-DI PHENYL SULPHONE $\text{C}_6\text{H}_5\text{NSO}_2 \text{ v e } \text{NMe} \langle \text{C}_6\text{H}_5 \rangle \text{SO}_2$ [222°] Formed by the action of KMnO_4 on methyl-imido di phenyl sulphide suspended in water (Bernthsen, *A* 230, 91) Chains of white needles, often slightly reddish (from alcohol) or small compact prisms (from glacial acetic acid) V sl sol cold alcohol, glacial acetic acid or ether Insol acids or alkalis, not affected by boiling KOH or HCl Boiled with conc H_2SO_4 it forms a splendid blue liquid, turned pale violet-brown by pouring into water Its nitro-derivative after reduction gives no colour with FeCl_3

METHYL INDAZINE $\text{C}_6\text{H}_7\text{N}_2 \text{ v e}$

$\text{C}_6\text{H}_7 \langle \text{CMe} \rangle \text{NH}$. Quinazole. *Methyl indazole*.

[113°]. (281° v) at 736 mm V D 4.37 (calc 4.53) Formed by slowly adding NaNO_2 to a paste composed of *o* amido acetophenone and HClAq The resulting diazo compound is soluble in water, and the liquid is poured into a solution of Na_2SO_3 The product at first contains $\text{CH}_3\text{CO C}_6\text{H}_4\text{NH NHSO}_3\text{Na}$, but it gradually loses its reducing power, and deposits needles of

$\text{C}_6\text{H}_7 \langle \text{CMe} \rangle \text{N SO}_3\text{Na}$ When these needles are

heated with HClAq they yield methyl indazine and H_2SO_4 (Fischer a Tafel, *A* 227, 303) Methyl indazine is also formed, together with carbonic acid gas, when indazyl acetic acid

$\text{C}_6\text{H}_7 \langle \text{C CH}_2\text{CO}_2\text{H} \rangle \text{NH}$ is heated

Properties — Colourless needles (from water), m sol hot water, v sol alcohol, ether, and chloroform, almost insol conc NaOHAq May be sublimed Does not reduce Fehling's solution Gives crystalline compounds with metallic salts

Salts — B'HCl [177°] Needles, v sol water and alcohol — Sulphate needles — Picrate yellow crystalline powder — Platino chloride yellow needles

Nitrosamine $\text{C}_6\text{H}_7\text{N}_3\text{O}$ v e

$\text{C}_6\text{H}_7 \langle \text{CMe} \rangle \text{NNO}$ [60°] From B'HCl and

cold aqueous NaNO_2 Yellow needles, v sol alcohol, ether, and chloroform

Di methyl indazine $\text{C}_6\text{H}_9\text{N}_2 \text{ v e}$

$\text{C}_6\text{H}_7 \langle \text{CMe} \rangle \text{NMe}$ [80°] Formed by methylation of the preceding Colourless plates, v sol alcohol, ether, benzene, and hot water

Di methyl ψ indazine $\text{C}_6\text{H}_7 \langle \text{CMe} \rangle \text{NMe} \text{N}$ [36°]

Formed from methyl *o* amido acetophenone by conversion into the nitrosamine

$\text{CH}_3\text{CO C}_6\text{H}_4\text{NMeNO}$ and reducing this body with zinc and HOAc (Fischer a Tafel, *A* 227, 336) Yellow oil, solidifying on cooling as nearly colourless plates Very volatile with steam It forms crystalline compounds with metallic salts — The chloride forms colourless needles — The sulphate crystallises in needles, and the picrate in rectangular tables

METHYL INDAZINE ψ -SULPHONIC ACID

$\text{C}_6\text{H}_7\text{N}_2\text{SO}_3 \text{ v e } \text{C}_6\text{H}_7 \langle \text{CMe} \rangle \text{N SO}_3\text{H}$ The

sodium salt of this acid is formed as described under methyl indazine It is m sol cold water, but ppd on addition of NaOH or NaCl It does not reduce HgO or Fehling's solution Boiling conc HClAq splits it up into NaHSO_4 and methyl-indazine

METHYL INDAZOLE ψ METHYL INDAZINE.

DI-METHYL-INDIGO ψ INDIGO

METHYL-ISO-INDILEUCINE ψ INDIGO.

(In. 1) METHYL INDOLE C_8H_7N i.e.

$C_8H_7\langle\begin{smallmatrix} CMe \\ NH \end{smallmatrix}\rangle CH$ Scatole Skatole [95°]
(266° i V) V D (H=1) 65.2 (calc 65.5)

Occurrence—The chief volatile constituent of human faeces, but not present in that of dogs (Brieger, *J pr* [2] 17, 129, *B* 12, 1986) It occurs among the products of the putrefaction of albuminous substances (Brieger, Nencki, *J pr* [2] 17, 98, *H* 4, 371, *E* a *H* Salkowsky, *B* 12, 651)

Formation—1 By fusing egg albumen with KOH (Nencki)—2 By reducing indigo with tin and HCl, and distilling the product with zinc dust A mixture of indole and scatole is thus obtained, and these are combined with picric acid When the picrates are distilled with conc NaOHAq the indole is destroyed, and the scatole passes over, and may be crystallised from water The yield is 3 p.c. Scatole prepared in this way has no faecal odour (Baeyer, *B* 13, 2339)—3 By heating aniline zinc chloride with glycerine (Fischer a German, *B* 16, 710)—4 By adding nitro cummic acid (6 pts) to amido cummic acid (obtained by reducing 4 pts of the nitro acid), mixing with baryta (10 pts), drying, and distilling (Fileti, *G* 13, 358, 378)—5 The phenyl hydrazide of propionic acid is warmed with $ZnCl_2$, and the product distilled with steam (*E* Fischer, *A* 236, 138)—6 In small quantity in distilling strychnine with lime (Stoehr, *B* 20, 1108, Lobisch a Malfatti, *M* 9, 629)—7 By heating its carboxylic acid (Arnold, *A* 246, 835)

Preparation from pancreas—2,300 grms pancreas and 500 grms flesh are freed from fat, cut up and put into a loosely covered pot containing 8 litres of water The whole is left for 5 months at the ordinary temperature At the end of the fourth month the odour of scatole appears The liquid is acidified with acetic acid and distilled The distillate is acidified with HCl and picric acid is added Red needles of skatepicric acid, $C_8H_7N \cdot C_6H_4(NO_2)_3OH$, separate This is decomposed by ammonia, and the scatole is distilled over with steam and crystallised from water (Nencki, *J pr* [2] 20, 467)

Properties—Glittering plates of powerful faecal odour, sl sol water It differs from indole in giving no colour when its solutions are treated with chlorine water Its solution gives with fuming HNO_3 , an opalescence, with KNO_3 and acetic acid a white pp of the nitrosamine, with CrO_3 a red amorphous pp in concentrated solutions, and with $FeCl_3$ no colour (Brieger, *J pr* [2] 17, 130) It is not attacked by warm dilute HNO_3 It colours pine wood moistened with HCl red, this is best seen by dropping pine wood saturated with an alcoholic solution of scatole into cold conc HClAq (Fischer, *A* 236, 138)

Reactions—1 Potash-fusion yields the corresponding indole carboxylic acid (Ciamician a Magnanini, *B* 21, 673)—2 By passing CO over a mixture of sodium and scatole, heated at 240° there is formed indole (In-1)-carboxylic acid (C a M)—3 With chloroform and NaOEt it yields a chloro methyl-quinoline C_9H_7ClN —4 Benzonic aldehyde and $ZnCl_2$ form, slowly at 100°, $C_8H_7 \cdot CH(C_6H_5 \cdot N)$, which crystallises from alcohol [142°] (Fischer, *B* 19, 2989)—5 When adminis-

tered to animals it appears in the urine in the form of the chromogen of a red pigment and as methyl indyl sulphuric acid $C_8H_7N \cdot SO_3H$ (?) (Brieger, *H* 4, 414, Mester, *H* 12, 130)

Salts— B^+HCl Ppd in needles when ether is added to its alcoholic solution (Wenzing, *A* 239, 239) [168°] Insol ether, v sl sol water, v sol alcohol—Picrate $B^+C_6H_4(NO_2)_3OH$ Red needles

Acetyl derivative so called v METHYL-INDYL METHYL KETONE

Dihydride $C_8H_{11}N$ i.e.

$C_8H_7\langle\begin{smallmatrix} CHMe \\ NH \end{smallmatrix}\rangle CH_2$ (232° i V) at 744 mm

Formed by reducing an alcoholic solution of scatole with zinc dust and HCl (Wenzing, *A* 239, 242) Colourless oil, resembling quinoline and piperidine in odour V sol alcohol, ether, and ligroin Its alcoholic solution stains pine wood, moistened with HClAq, orange It reduces $AgNO_3$ and $FeCl_3$ on warming It yields a nitrosamine which may be reduced to an oily hydrazine With phenyl thiocarbimide it forms a compound melting at 125°— B^+HCl sol. alcohol and water, insol ether—Oxalate [126°], insol ether— $B^+H_2PtCl_6$, yellow needles, sl sol water, decomposed by hot water—Picrate [150°], yellow granular crystals (from benzene)

(In 2) Methyl-indole C_8H_7N i.e.

$C_8H_7\langle\begin{smallmatrix} CH \\ NH \end{smallmatrix}\rangle CMe$ Methyl ketole [60°] (272° i V) at 750 mm V D 4.75 (calc 4.54) (Treadwell, *B* 14, 1466)

Formation—1 By nitrating benzyl methyl ketone with fuming HNO_3 , reducing the resulting $[2]C_8H_7(NO_2)CH_2COCH_3$ with zinc dust and ammonia, and distilling with steam (Baeyer a Jackson, *B* 13, 187, 14, 879)—2 Obtained by heating the phenyl hydrazide of acetone $(CH_3)_2C=N \cdot NHC_6H_5$ (1 pt) with $ZnCl_2$ (5 pts) for half an hour at 100° and then for some minutes at 180° (*E* Fischer, *B* 19, 1564, *A* 236, 124)

Properties—Needles or plates (from ligroin), al. sol hot water, v sol alcohol and ether Smells like indole V sol cold HClAq, but decomposed on boiling with conc HClAq HNO_3 colours its solution yellow, and soon gives a yellow amorphous pp which does not give Liebermann's reaction Colours pine wood, moistened with HClAq, red

Reactions—1 $KMnO_4$ oxidises it to acetyl o amido benzoic acid—2 Potash fusion yields indole (In 2) carboxylic acid (Ciamician a Magnanini, *B* 21, 673)—3 By heating with sodium in a stream of CO_2 at 240° there is formed methyl indole carboxylic acid (Ciamician a Magnanini, *B* 21, 671)—4 Reduced by tin and HCl to a hydride, although sodium amalgam does not act upon it—5 With chloroform and NaOEt it yields chloro methyl quinoline [71°] Bromoform and NaOEt yield the corresponding bromo methyl quinoline (Magnanini, *G* 17, 246)—6 On heating with conc HClAq for 7 hours at 225° there is formed aniline and a liquid base C_9H_9N (250°) which smells like quinoline and forms the salts $B^+H_2PtCl_6$ and B^+HAuCl_4 (Magnanini, *B* 20, 2609)—7 Benzonic aldehyde at 100° forms $C_8H_7 \cdot CH(C_6H_5 \cdot N)$, which separates from acetone in colourless shining crystals

[247°], insol water, v. sl sol hot alcohol and ether. On boiling with FeCl_3 in HOAc it is converted into di-methyl rosindole (Fischer, *A* 242, 378, *B* 19, 2988) — 8 *Nitro-benzoic aldehyde* heated with (*In* 2) methyl indole on the water bath forms the corresponding $[\text{3.1}]\text{O}_2\text{C}_6\text{H}_4(\text{NO}_2)\text{CH}(\text{C}_6\text{H}_5)_2$, small crystals (from acetone), sl sol alcohol, ether, and HOAc . This body melts at 263° , and is reduced by zinc dust and ammonia to $\text{C}_6\text{H}_4(\text{NH}_2)\text{CH}(\text{C}_6\text{H}_5)_2$, a yellow crystallisable oil (Fischer, *A* 242, 375) — 9 By fusing with *phthalic anhydride* and a little ZnCl_2 , there is formed an acid $\text{C}_6\text{H}_4\text{NO}_2$, probably $\text{C}_6\text{H}_4\text{N CO C}_6\text{H}_4\text{CO}_2\text{H}$ which crystallises from alcohol in colourless prisms, insol water, l sol ether, v sol hot alcohol and HOAc (Fischer) — 10 On heating with *benzoyl chloride* and a little ZnCl_2 , the products are di-methyl-rosindole $\text{C}_6\text{H}_4\text{N}_2$ and benzoyl methyl indole. — 1 On heating with *zinc chloride* alone, a small quantity of quinoline is formed (Fischer a Steche, *B* 20, 819) — 12 *Diazo benzene chloride* and NaOAc form compact red crystals (from petroleum ether) of $\text{C}_6\text{H}_4\text{N N C} \begin{smallmatrix} \text{C}_6\text{H}_5 \\ \text{CMe} \end{smallmatrix} \text{NH}$, 116° , insol water, v sol alcohol, ether, and benzene, m sol petroleum ether. This body is split up again, by reduction, into aniline and amido methyl indole $\text{C}_6\text{H}_4 \begin{smallmatrix} \text{C}(\text{NH}_2) \\ \text{NH} \end{smallmatrix} \text{CMe}$

[113°] (Fischer, *A* 242, 384) — 13 When (*In* 2)-methyl indole (1 pt) is heated with MeI (2½ pts) and a little methyl alcohol at 100° for 15 hours, here is produced di-methyl quinoline dihydride (243°) (Fischer a Steche, *B* 20, 818, 2199)

Salts — B^+HI^- Formed by passing dry HI into a solution of the methyl indole in ether (Wagner, *A* 242, 388). Flocculent pp, very easily decomposed by water and by moist air — $\text{B}^+\text{H}_2\text{PtCl}_6$ 3aq yellow needles, decomposed by water

Acetyl derivative $\text{C}_6\text{H}_4 \begin{smallmatrix} \text{CH} \\ \text{NAc} \end{smallmatrix} \text{CMe}$ (200° – 210° at 40 mm) Formed in small quantity, together with methyl indyl methyl ketone (*q v*), by heating (*In* 2) methyl indole with Ac_2O and NaOAc . The product is extracted with chloroform, and the extract distilled *in vacuo* (Magnanini, *G* 18, 95). Pale yellow liquid. Decomposed by boiling aqueous KOH into KOAc and methyl indole. Yields indole carboxylic acid on fusion with potash. KMnO_4 oxidises it to acetyl o-amido benzoic acid (Ciamician a Marinum, *B* 21, 673)

(8) *Acetyl derivative v METHYL INDOL METHYL KETONE*

Benzoyl derivative $\text{C}_6\text{H}_4\text{NBz}$ [82°] Formed, together with di-methyl rosindole, by heating (*In* 2) methyl indole with BzCl and a little ZnCl_2 on the water bath (Fischer a Wagner, *B* 20, 817). Glittering plates (from alcohol), v sl sol hot water, m sol alcohol and ether

Dihydride $\text{C}_6\text{H}_4\text{N} \begin{smallmatrix} \text{CH}_2 \\ \text{NH} \end{smallmatrix} \text{CMe}$

Hydromethylketone (228° 1 V) at 742 mm Prepared by reduction of (*In* 2)-methyl indole with tin and HCl (Jackson, *B* 14, 883, Wenzig, *A* 239, 244). Colourless oil with powerful odour. Heavier than water. Strong base. With phenyl-thiocarbamide it forms a compound $\text{C}_6\text{H}_4\text{N}_2\text{S}$, which crystallises from ether in prisms

[101°] — $\text{B}^+\text{H}_2\text{PtCl}_6$, orange yellow needles, decomposed by water — Oxalate [130°], crystalline — Picrate [151°], crystalline

Acetyl derivative $\text{C}_6\text{H}_4\text{NAc}$ [56°] From the dihydride and Ac_2O . White needles, insol water, sol most other solvents

Nitrosamine $\text{C}_6\text{H}_4\text{N NO}$ [55°] Formed by adding NaNO_2 to a solution of the hydrochloride of (*In* 2) methyl indole dihydride (Jackson). Yellow crystals (from ligroin), v sol alcohol, ether, and boiling ligroin. On treatment with tin and HCl , methyl indole dihydride is regenerated. On reduction with zinc dust and HOAc it yields the hydrazine $\text{C}_6\text{H}_4\text{N NH}_2$, which crystallises from ligroin in prisms [41°], and yields a crystalline sulphate and hydrochloride

(*In* 3)-Methyl indole $\text{C}_6\text{H}_4 \begin{smallmatrix} \text{CH} \\ \text{NMe} \end{smallmatrix} \text{CH}$ (240° 1 V) at 720 mm SG η 1.0707. Obtained by long heating at about 205° from its carboxylic acid [212°], which is formed by the action of HCl on the phenyl methyl hydrazide of pyruvic acid (Fischer a Hess, *B* 17, 562). Formed also by heating ω chloro methyl o amido styrene $\text{C}_6\text{H}_4(\text{NHMe})\text{CH CHCl}$ with NaOEt at 135° (Lapp, *B* 17, 2510)

Properties — Yellowish liquid, nearly insol water, v sol alcohol, ether, and benzene. Volatile with steam. A chip of pine wood, dipped in HClAq , is coloured violet red by its vapour or solution. It dissolves in conc HClAq and is reprecipitated by addition of water. Fuming nitric acid added to (*In* 3) methyl indole suspended in water gives a deep red colour and finally a red pp

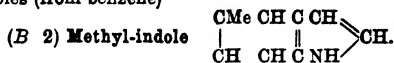
Reactions — 1 An alkaline solution of bromine (NaOBr) converts it into di-bromo methyl oxindole $\text{C}_6\text{H}_3\text{Br}_2\text{NO}$, which crystallises in transparent tables, melting at 204° . It is converted by heating with alcoholic potash into methyl isatin, which is ppd, after boiling off the alcohol, by adding HCl — 2 (*In* 3) methyl indole (2 mols) heated with benzoic aldehyde (1 mol) and ZnCl_2 for 2 hours on the water bath forms $\text{C}_6\text{H}_4\text{CH}(\text{C} \begin{smallmatrix} \text{C}_6\text{H}_5 \\ \text{CH} \end{smallmatrix} \text{NMe})_2$, which crystallises in colourless prisms [197°], insol water, sl sol alcohol and ether, v sol hot acetone and HOAc . It yields a red dye on heating with FeCl_3 in HOAc (Fischer, *A* 242, 377, *B* 19, 2988) — 3 On fusing equal weights of *phthalic anhydride* and (*In* 3) methyl indole with a little ZnCl_2 , at 100° there is formed $\text{C}_6\text{H}_4\text{C}_2\text{O}_2(\text{C}_6\text{H}_5\text{N})$, which crystallises from acetone in colourless prisms [300°]. It is insol water and alkalis, v sl sol ether and alcohol, but v sol hot acetone (Fischer, *A* 242, 382, *B* 19, 2989) — 4 Scarcely attacked by MeI at 100° , but at 120° it appears to yield a methyl quinoline dihydride (Fischer a Steche, *B* 20, 2199)

Picrate $\text{C}_6\text{H}_4\text{N} \begin{smallmatrix} \text{CH} \\ \text{NO}_2 \end{smallmatrix} \text{OH}$ [150°] Long dark red prisms or needles, v sol benzene, sl sol ether. Decomposed by alcohol

Chloro derivative v Di-chloro methyl-indole.

Dihydride $\text{C}_6\text{H}_4 \begin{smallmatrix} \text{CH}_2 \\ \text{NMe} \end{smallmatrix} \text{CH}_2$ (216° 1 V) at 728 mm Prepared by reducing (*In* 3) methyl indole with zinc dust and conc HClAq (Wenzig, *A* 239, 246). Liquid, sl sol water, v sol alcohol and ether, volatile with steam. Most of its salts dissolve in alcohol and water — $\text{B}^+\text{H}_2\text{PtCl}_6$:

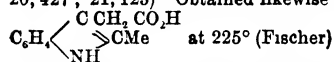
yellow needles, decomposed by boiling water — Oxalate [105°] — Picrate [155°], yellow tabs (from benzene)



Tolindole [58 5°] Formed by heating at 240° its carboxylic acid, which is obtained from the *p* tolyl hydrazide of pyruvic ether (Raschen, *A* 239, 226) Needles (from water), m sol hot water, v sol alcohol, ether, benzene, and ligroin Volatile with steam Reacts like indole with pine wood and with nitrous acid — Picrate $\text{B}'\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OH}$ [151°] Red needles (from water)

(In 1,2) Di methyl indole $\text{C}_{10}\text{H}_{11}\text{N}$ *z* *e*

$\text{C}_6\text{H}_5 \begin{array}{c} \text{CMe} \\ \diagup \quad \diagdown \\ \text{NH} \end{array} \text{CMe.}$ [108°] (285°; ∇) Formed by heating the phenyl hydrazide of methyl ethyl ketone with ZnCl_2 at 180° (E Fischer, *B* 19, 1565, *A* 236, 126) Formed also by heating bromo acetyl propionic acid $\text{CH}_3\text{COCHBrCH}_2\text{CO}_2\text{H}$ (1 pt) with aniline (3 pts) at 100° (Wolff, *B* 20, 427, 21, 123) Obtained likewise by heating



Properties — White plates (from dilute alcohol) Smells like indole V sl sol hot water, v e sol alcohol and ether, sl sol cold ligroin Dissolves in conc HClAq but is reprecipitated on dilution with water Does not colour pine wood

Reactions — 1 NaNO_2 added to its solution in HOAc forms a nitrosamine [63°], crystallising in yellow needles, v sl sol water, v sol alcohol By zinc dust and HCl it is reconverted into di methyl indole It exhibits Liebermann's reaction — 2 MeI in MeOH converts it into tri methyl quinoline dihydride

Picrate $\text{B}'\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OH}$ [157°] Brown needles (from alcohol)

Dihydride $\text{C}_6\text{H}_5 \begin{array}{c} \text{CHMe} \\ \diagup \quad \diagdown \\ \text{NH} \end{array} \text{CHMe}$ (231°) at 750 mm Formed by reducing di methyl indole with zinc dust and HClAq (Steche, *A* 242, 371)

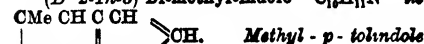
(In 2,3) Di methyl indole $\text{C}_{10}\text{H}_{11}\text{N}$ *z* *e*

$\text{C}_6\text{H}_5 \begin{array}{c} \text{CH} \\ \diagup \quad \diagdown \\ \text{NMe} \end{array} \text{CMe}$ [56°] Formed by heating the phenyl methyl hydrazide of acetone (1 pt) with zinc chloride (5 pts) for 3 hours at 130° (Fischer, *B* 19, 1565, Degen, *A* 236, 153) Formed also by heating its carboxylic acid

$\text{C}_6\text{H}_5 \begin{array}{c} \text{C}(\text{CO}_2\text{H}) \\ \diagup \quad \diagdown \\ \text{N}(\text{CH}_3) \end{array} \text{CMe}$ at 200° (D) White needles, v sol alcohol, ether, benzene, and ligroin, v sl sol water, v sol conc HClAq May be distilled without decomposition It gives the pine wood test very distinctly Nitrous acid forms a complicated product The picrate crystallises in dark red needles The hydride is v sol weak acids

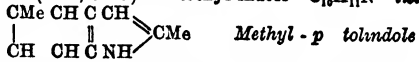
(In 1,3) Di methyl indole $\text{C}_6\text{H}_5 \begin{array}{c} \text{CMe} \\ \diagup \quad \diagdown \\ \text{NMe} \end{array} \text{CH}$ (230°–255°) Formed by heating the phenyl-methyl hydrazide of *n* propionic aldehyde (1 pt) with ZnCl_2 (5 pts) at 135° (Degen, *A* 236, 163) Oil

(B 2-In-3) Di-methyl-indole $\text{C}_{10}\text{H}_{11}\text{N}$ *z* *e*



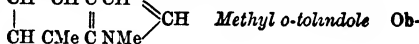
(242°–245°) Formed by heating its carboxylic acid at 225° (Hegel, *A* 232, 216) Liquid, volatile with steam, v sol. alcohol, ether, and benzene Dyes pine wood, moistened with HCl , red, fuming HNO_3 gives a red colour and, finally, a pp The picrate is crystalline

(B 2, In 2) Di methyl indole $\text{C}_{10}\text{H}_{11}\text{N}$ *z* *e*



[114°–117°] Formed by heating the *p*-tolyl-hydrazide of acetone with ZnCl_2 (Raschen, *A* 239, 227) May be distilled without decomposition Almost insol hot water, v sol hot alcohol, ether, and benzene — $\text{B}'\text{C}_6\text{H}_4(\text{NO}_2)_2(\text{OH})$ [155°] Dark red needles (from benzene)

(B 4, In 2) Di methyl indole $\text{C}_{10}\text{H}_{11}\text{N}$ *z* *e*



tained by the action of heat on its carboxylic acid, which is derived from the *o* tolyl methyl-hydrazide of pyruvic acid (Hegel, *A* 232, 220) Liquid, smelling like indole, volatile with steam Dyes pine wood, acidified by HCl , violet red. Behaves like indole towards nitrous acid

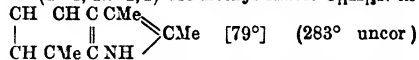
Di-methyl indole $\text{C}_{10}\text{H}_{11}\text{N}$ (275°) Formed by allowing a solution of the hydrochloride of (a) di methyl dipyrrole in dilute H_2SO_4 to stand for some time in the cold (Dennstedt, *B* 21, 3439) Liquid, volatile with steam — Picrate $\text{B}'\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OH}$ [156°] Dark red silky needles (from benzene)

Di methyl indole $\text{C}_{10}\text{H}_{11}\text{N}$ (c 270°) Formed from (B) di methyl dipyrrole in the same way as the preceding isomeride (D) Liquid Smells like scatole — Picrate $\text{B}'\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OH}$ [149°] Red silky needles

(In 1,2,3) Tri methyl indole $\text{C}_{11}\text{H}_{13}\text{N}$ *z* *e*

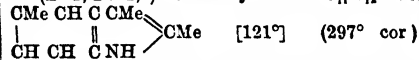
$\text{C}_6\text{H}_5 \begin{array}{c} \text{CMe} \\ \diagup \quad \diagdown \\ \text{NMe} \end{array} \text{CMe}$ (280°) Formed by heating di methyl indyl acetic acid for several hours at 210° (Degen, *A* 236, 160) Formed also by heating the phenyl methyl hydrazide of methyl ethyl ketone with ZnCl_2 at 180° (D) Yellow oil with characteristic odour Miscible with alcohol and ether Gives no colour to pine wood Picrate $\text{B}'\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OH}$ [150°] Dark red needles (from hot benzene)

(B 4, In 1,2) Tri methyl indole $\text{C}_{11}\text{H}_{13}\text{N}$ *z* *e*



Formed by heating bromo acetyl propionic (bromo levulic) acid $\text{CH}_3\text{COCHBrCH}_2\text{CO}_2\text{H}$ with *o* toluidine (Wolff, *B* 21, 3362) White plates, v sol alcohol, chloroform, and petroleum ether, v sl sol water Volatile with steam — Picrate $\text{B}'\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OH}$ [152°] Purple red scales

(B 2, In 1,2) Tri-methyl indole $\text{C}_{11}\text{H}_{13}\text{N}$ *z* *e*



Formed by warming β bromo acetyl propionic acid (1 pt) with *p* toluidine (3 pts), and finally heating the mixture to boiling (Wolff, *B* 21, 3361) Plates, v sol alcohol, chloroform, and petroleum ether, v sl sol water Does not give the pine wood reaction The solution in HOAc , mixed with FeCl_3 , becomes green on boiling, and

finally blue Picrate $B/C_6H_3(NO_2)_3OH$ [189°] Brownish-red needles, v sol benzene and alcohol

Nitrosamine $C_8H_7<\frac{CMe}{N(NO)}>CMe$ [73°] Golden-yellow needles, v e sol alcohol and HOAc, sl sol water

Tetra-methyl-indole $C_{12}H_{15}N$ (235°) Light-yellow oil, with characteristic odour (Dennstedt, B 22, 1924) Picrate $B/C_6H_3(NO_2)_3OH$ [100°] Shining red needles

METHYL-INDOLE ACETIC ACID v **METHYL-INDYL-ACETIC ACID**

(In 1)-**METHYL-INDOLE** (In 2)-**CARBOXYLIC ACID** $C_{10}H_9NO_2$ \pm e

$C_8H_7<\frac{CMe}{NH}>C CO_2H$ *Scatole* (β)-carboxylic acid [165°] Formed by saponifying with alcoholic potash its ether which is obtained by boiling with alcoholic hydrogen sulphate (10 p c), the phenyl hydrazide of ethyl glyoxylic ether $CH_3CH_2C(N,HPH)CO_2H$ (Wilschusen a Arnold, B 20, 3395, A 246, 335) Formed also by heating scatole with sodium in a current of CO_2 at 240° (Ciamician a Magnanini, G 13, 61, B 21, 672, 1927, *Rend Accad Linc* [4] 4, 740) Thin white needles (from boiling water), v sol alcohol and benzene, v sl sol water Its alcoholic solution is coloured deep red by $FeCl_3$. On heating above 165° it is split up into CO_2 and scatole [95°] — AgA' white powder, insol water

Ethyl ether EtA' [134°] Needles (from alcohol), insol water, v e sol benzene and ether

(In 1)-**Methyl-indole carboxylic acid** $C_{10}H_9NO_2$ *Scatole* (α) carboxylic acid [164°] (Possibly identical with the preceding acid) Occurs among the products of the putrefaction of serum albumen, and of muscular tissue (H a E Salkowsky, B 13, 191, 2217, H 8, 23, 9, 8) Small plates (from benzene) Decomposed on heating above its melting point into CO_2 and scatole v sl sol cold water, v sol alcohol and ether, m sol benzene $FeCl_3$ colours its dilute solution (containing HCl) violet on boiling Nitrous acid colours its dilute solution cherry red, and presently gives a pp Bleaching powder colours a dilute solution, acidified by HCl , purple, this reaction, and that with nitrous acid, are not exhibited by (In 1)-indole (In 2) carboxylic acid — AgA' Sparingly soluble pp

(In 2) **Methyl-indole** (In 1)-carboxylic acid $C_8H_7<\frac{C CO_2H}{NH}>CMe$

Methyl ketole carboxylic acid [172°] or [183°] Prepared by heating (In 2)-methyl indole (10 g) mixed with sodium (3 g) in a current of dry CO_2 first at 235° and finally at 315°, the unaltered methyl-indole is removed by steam distillation, and the acid re-crystallised from acetone (Ciamician a Magnanini, G 13, 60, B 21, 672, *Rend Accad Linc* [4] 4, 740) White crystalline powder, completely decomposed at its melting point into CO_2 and methyl indole (methyl ketole), this decomposition is partially effected by merely boiling the aqueous solution Sl sol water and benzene, v sol alcohol and acetone It gives white pps with lead and mercuric salts, and a

green pp with a cupric salt — AgA' white oryztalline pp

(In 3)-**Methyl-indole** (In 1) carboxylic acid $C_8H_7<\frac{C(CO_2H)}{N(CH_3)}>CH$ [212°] Obtained by

heating the phenyl methyl hydrazide of pyruvic acid (1 pt) with 10 p c aqueous HCl (15 pts) on the water bath (E Fischer a Hess, B 17, 559) White needles, v sol hot alcohol, ether, and benzene, sl sol hot water, nearly insol cold water Its solution in H_2SO_4 is red By prolonged heating at its melting point it is split up into CO_2 and (In 3)-methyl-indole Readily oxidised by $KMnO_4$. An alkaline solution of Br or Cl oxidises it to methyl- ψ isatin and CO_2

(B 4)-**Methyl-indole** (In 2) carboxylic acid $CH CH C CH > C CO_2H$ [171°] Formed by

saponifying with alcoholic potash its ethyl ether which is obtained by heating the o-tolyl hydrazide of pyruvic ether (1 pt) with $ZnCl_2$ (1 pt) at 220° (Raschen, A 239, 228) Needles (from water), v sol alcohol, ether, and HOAc

(B 2)-**Methyl-indole** (In 2) carboxylic acid $CMe CH C CH > C CO_2H$ [228°] Formed from

its ethyl ether which is obtained by heating the p-tolyl hydrazide of pyruvic ether with $ZnCl_2$ at 220° (Raschen, A 239, 228) Needles (from water), m sol hot water, v sol alcohol, ether, chloroform, and HOAc Decomposed on fusion into CO_2 and (B 2) methyl indole

Ethyl ether EtA' [160°] Colourless needles or plates

(In 2, 3) **Di-methyl-indole** (In 1)-carboxylic acid $C_{11}H_{11}NO_2$ \pm e $C_8H_7<\frac{C(CO_2H)}{NMe}>CMe$

[185°] Formed by saponifying its ether which is obtained by heating the phenyl methyl hydrazide of acetoacetic ether with $ZnCl_2$ (Fischer, B 19, 1569, Degen, A 236, 157) Six-sided plates, m sol hot alcohol and chloroform, sl sol water, ether, benzene, and ligroin Splits up at 200° into CO_2 and (In 2, 3) di-methyl indole The Na salt is insol conc $NaOHAq$ The Ag salt is insol NH_4Aq Colours pine wood

Ethyl ether EtA' [95°] Colourless needles (from alcohol-ligroin), v sol alcohol ether, benzene, and chloroform, sl sol ligroin

(B 4, In 3) **Di-methyl indole** (In 2)-carboxylic acid $C_{11}H_{11}NO_2$ \pm e

$CH CH C CH > C CO_2H$ [210°] Formed by

warming the o-tolyl methyl-hydrazide of pyruvic acid (1 pt) with phosphoric acid (20 pts of S G 1 17) on the water bath (Hegel, A 262, 220) Colourless needles (from benzene), v e sol alcohol Splits up on heating into CO_2 and the corresponding di-methyl indole When its alkaline solution is warmed with $NaOCl$ and the product heated with water there is formed di-methyl ψ isatin $C_8H_7Me<\frac{CO}{NMe}>CO$ [157°]

(B 2, In 3)-**Di-methyl-indole** (In 2) carboxylic acid $CMe CH C CH > C CO_2H$ [221°].

Formed by warming the p-tolyl methyl hydra-

sides of pyruvic acid (1 pt) with HCl (20 pts of 10 p.c.), the acid being pptd (Hegel, *A* 232, 216). Needles (from hot alcohol), sl sol ether, sol hot benzene, chloroform, and glacial HOAc, insol in groin. Forms the corresponding di-methyl-indole on heating. A neutral solution of its Na salt gives with NaOCl a yellow crystalline ppt [135°], which on boiling with water is changed to di-methyl-ψ isatin $C_8H_7Me \begin{smallmatrix} <CO \\ <NMe \end{smallmatrix} > CO$ crystals, crystallising in red needles [148°].

METHYL INDONAPHTHENE $C_{11}H_{10}$, *ψ* $C_8H_7 \begin{smallmatrix} <CMe \\ <CH_2 \end{smallmatrix} > CH$ Methyl-indene (206°)

(Roser), (201°) (Von Pechmann, *B* 16, 516). Formed by distilling methyl indonaphthene carboxylic acid with soda lime (Roser, *A* 247, 159). Transparent highly refractive liquid, smelling somewhat like naphthalene. Absorbs oxygen from the air, becoming gummy. On heating with conc HClAq it also becomes gummy, and H₂SO₄ has a like effect. By heating with HIAq at 180° it is converted into a solid body, fusible under water, v sol ether, sl sol alcohol.

Picric acid compound
 $C_8H_7C_6H_2(NO_2)_3OH$ [76°] Unstable orange powder (Von Miller, *B* 23, 1882).

METHYL INDONAPHTHENE CARBOXYLIC ACID $C_{11}H_{10}O_2$, *ψ* $C_8H_7 \begin{smallmatrix} <CMe \\ <CH_2 \end{smallmatrix} > CCO_2H$

'Dihydronaphthoic' acid [200°] Formed by gently warming benzyl aceto acetic ether (1 pt) with conc H₂SO₄ (7 pts), and pouring the product into water (Von Pechmann, *B* 16, 516, Roser, *B* 20, 1574, *A* 247, 158). White needles (from alcohol), almost insol water, m sol hot alcohol and ether. Crystallises from glacial acetic acid in prisms (containing HOAc). May be distilled with slight decomposition, but when boiled for a long time it splits up into CO₂ and methyl indonaphthene. On oxidation it yields phthalic acid.

Methyl ether *MeA' [78°] Small needles, v sol alcohol and ether.

Di-bromide $C_8H_7 \begin{smallmatrix} <CMeBr \\ <CH \end{smallmatrix} > CBr CO_2H$

[215°] Formed by exposing the acid to bromine-vapour. White crusts (from ether). On warming with HOAc it gives off HBr and leaves $C_8H_7 \begin{smallmatrix} <CMeBr \\ <CH \end{smallmatrix} > C CO_2H$ [245°], which forms a methyl ether melting at 100°.

Methyl ether $C_8H_7BrCO_2Me$. [157°] White needles, sl sol methyl alcohol.

METHYL INDONAPHTHENE DIHYDRIDE CARBOXYLIC ACID $C_{11}H_{12}O_2$, *ψ* $C_8H_9 \begin{smallmatrix} <CHMe \\ <CH_2 \end{smallmatrix} > CH CO_2H$ Methyl-hydrindonaphthene carboxylic acid [80°] (300°-310°)

Formed by the action of sodium amalgam on an alkaline solution of methyl indonaphthene carboxylic acid (Roser, *B* 20, 1574, *A* 247, 165). Small needles (from hot water), v sol alcohol and ether, scarcely volatile with steam. On continued heating above 310° it becomes viscid, but the crystalline acid is reproduced on treating this viscid mass with alkalis—AgA'—BaA', 2aq needles (from alcohol), v sol water.

METHYL INDONAPHTHOQUINONE

$C_{11}H_8O$, *ψ* $C_8H_7 \begin{smallmatrix} <CO \\ <CO \end{smallmatrix} > CHMe$. Methyl-di-keto-

hydrindene [85°] (c 150° at 18 mm) Formed by the action of sodium on a mixture of ethyl propionate and phthalate (Wishlencus a Kötze, *A* 252, 80). Blunt pyramids (from alcohol) or needles (from light petroleum), sol ether and hot water. The sodium derivative $C_{11}H_7NaO_2$ crystallises in small dark-red prisms.

Di-oxim $C_8H_7 \begin{smallmatrix} <C(OH) \\ <C(OH) \end{smallmatrix} > CHMe$ [117°].

Needles, sol alcohol, ether, alkalis, and HOAc.

Mono-phenyl-hydrazide

$C_8H_7 \begin{smallmatrix} <C(N_2HPh) \\ <CO \end{smallmatrix} > CHMe$ [164°] Crystals, v sol ether and benzene. Forms a green solution in HOAc.

Di-methyl-indonaphthoquinone $C_{11}H_8O_2$, *ψ* $C_8H_7 \begin{smallmatrix} <CO \\ <CO \end{smallmatrix} > CMe$ [108°] (c 250°) Formed

by heating methyl indonaphthoquinone with MeI and MeOH at 100° (W a K). V sol alcohol, ether, and benzene.

Phenyl hydrazide

$C_8H_7 \begin{smallmatrix} <C(N_2HPh) \\ <C(N_2HPh) \end{smallmatrix} > CMe_2$ [184°-187°] Slender yellowish prisms (from alcohol), sol ether and HOAc. FeCl₃ colours its solution in H₂SO₄ intensely dark green.

METHYL INDOPHENINE *ψ* **INDOPHENINE**
METHYL-INDYL ACETIC ACID $C_{11}H_{11}NO_2$, *ψ* $C_8H_9 \begin{smallmatrix} <C \\ <CH_2CO_2H \end{smallmatrix} > NH$ [195°-200°]

Formed by warming the phenyl hydrazide of acetyl propionic (levulic) acid with ZnCl₂ at 125° (E Fischer, *B* 19, 1565, *A* 236, 149). Colourless plates, sl sol hot water and chloroform, m sol ether, v sol hot alcohol, v e sol acetone and hot HOAc. Nitrous acid yields a nitrosamine. At 225° it is split up, slowly into CO₂ and di-methyl indole. Its picric acid compound crystallises in slender dark red needles.

Di-methyl-indyl-acetic acid $C_{12}H_{13}NO_2$, *ψ* $C_8H_9 \begin{smallmatrix} <C \\ <CH_2CO_2H \end{smallmatrix} > NH$

$C_8H_9 \begin{smallmatrix} <C \\ <NMeCMe \end{smallmatrix} >$ [188°] Formed by

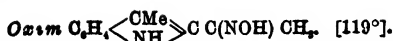
saponification (by alcoholic KOH) of its ether, which is obtained by warming the phenyl methyl-hydrazide of acetyl propionic ether with ZnCl₂ (Fischer, *B* 19, 1568, Degen, *A* 236, 158). Colourless plates, sl sol water, ether, and benzene, v sol alcohol and chloroform. Its alkaline salts are v e sol water, but pptd on addition of caustic alkalis. At 210° it is split up into CO₂ and (In-1,2,3) tri-methyl-indole. Does not give the pine-wood reaction.

METHYL INDYL KETONE *ψ* (β) **Acetyl-INDOLE**

(In 1) **METHYL INDYL METHYL KETONE**

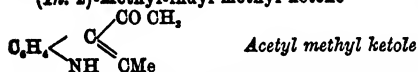
$C_{11}H_{11}NO$, *ψ* $C_8H_9 \begin{smallmatrix} <CMe \\ <NH \end{smallmatrix} > CCOCH_3$ Acetyl-scatole [148°] Formed by heating (In-1)-methyl indole (scatole) (1 g) with AcCl (10 g) and ZnCl₂ (½ g) (Magnanini, *G* 18, 99, *B* 21, 1938). Long needles (from dilute alcohol), insol cold water, v sol hot alcohol and acetone, m sol ether. Volatile with steam. Not affected by KOHAq, but conc HCl forms scatole. Hot conc H₂SO₄ gives a purple solution.

Picrate [156°] Yellow needles, sl sol cold, v sol hot, benzene.



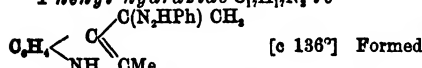
Small needles, decomposed by acids and alkalis, reproducing the ketone

(*In* 2)-Methyl-indyl methyl ketone



[196°]. Formed by heating (*In* 2) methyl indole (methyl ketole) (1 pt) with Ac_2O (5 pts) and NaOAc (1 pt) for six hours with inverted condenser (Jackson, *B* 14, 879, Fischer, *A* 242, 379). Colourless needles (from acetone), sl sol water, v sol alcohol and hot benzene. Dissolves in HClAq . Not decomposed by boiling NaOHAq , but boiling conc HClAq regenerates the ketone. KMnO_4 oxidises it to acetyl amido benzoic acid (Magnanini, *G* 18, 97).

Phenyl hydrazide $\text{C}_{11}\text{H}_9\text{N}_3$, s



by heating the ketone with phenyl hydrazine hydrochloride and NaOAc . Colourless plates, m sol hot benzene, v sl sol petroleum ether.

DI-METHYL INOSITE v DAMBONITE

METHYL-IODAMINE v METHYLAMINE

METHYL IODIDE CH_3I Iodo methane Mol

w 142 (42.8°) (Dobner, *A* 243, 23), (42.3°) (Perkin, *O* J 45, 459) S G 2 3346 (D), 2 2852, 2 2529 (P) S 008 at 15° (Bardy a Bordet, *A* Ch [5] 16, 569) CE (0°–10°) 00118 (D) S V 64.1 (Lossen, *A* 254, 69), 64.8 (Ramsay), M M 9 009 at 15° H F p (gas) +3420 H F v (gas) 2840 (Thomsen, *Th*). Obtained by distilling phosphorus (1 pt) with iodine (8 pts), dissolved in moist methyl alcohol (14 pts), or phosphorus (60 g) with iodine (1000 g) and methyl alcohol (500 g) (Dumas a Pélignot, *A* 15, 30, Pierre, *A* 56, 147, Landolt, *A* 84, 44, Hofmann, *C* J 13, 69). Prepared also from red phosphorus (10 pts), MeOH (35 pts), and I (100 pts) (Personne, *J* 1861, 607, Butlerow, *B* 6, 561), and from KI, MeOH , and gaseous HCl (De Vri, *J* 1857, 441). The distillate is washed with water and rectified over CaCl_2 and lead oxide. Colourless, slightly combustible liquid. Not attacked by gaseous HCl .

Reactions—1 Chlorine converts it into MeCl —2 When heated with zinc it forms I ZnMe . The moist copper zinc couple forms methane (Gladstone a Tribe, *C* J 26, 682)—3 When heated with alloys of K or Na with As or Sb it yields methides of arsenic or antimony—4 With Mg and Al it forms methides—5 When heated in a sealed tube with aqueous NH_3 it forms mono, di, and tri methylamine and NMe_3 —6 With EtOH at 125° it forms EtI and MeOEt (Busse a Kraut, *A* 177, 272)—7 Heated with water (15 pts) at 100° it is converted into methyl alcohol (Niederist, *A* 196, 350)—8 With H_2S and water it forms $\text{MeI}(\text{H}_2\text{S})$, 28aq (Forcrand, *A* Ch [5] 28, 21)—9 Sulphur at 160° to 190° forms SMe_2 and other products (Klinger, *B* 10, 1880).

Hydrate (MeI), aq [–4°] (F), [4.8°] (Villard, *C* R 111, 185). Formed by passing a current of moist air through the iodide (Forcrand, *C* R 90, 1491).

METHYL-iodoform v TRI IODO ETHANE

METHYL-ISATIC ACID v ISATIC ACID

METHYL-ISATIN v ISATIN

METHYL-ISATOIC ACID v ISATOIC ACID

METHYL-ISATOID v ISATOID

METHYL-ITACONIC ACID $\text{C}_6\text{H}_4\text{O}_4$, [166°].

A product of the distillation of methyl paraconic acid (Fränkel, *A* 255, 37). Formed also by heating a solution of methyl citraconic acid at 150° in sealed tubes. Prisms, sol water, insol chloroform. When distilled it partially carbonises, and is partially converted into the isomeric methyl citraconic acid. Nitric acid appears to convert it into methyl mesaconic acid [195°]. Sodium amalgam reduces it to ethyl succinic acid— $\text{BaA}''\frac{1}{2}\text{aq}$ — $\text{CaA}''\text{aq}$ — AgA'' .

TETRA METHYLUM HYDROXIDE v

Methyl hydroxide of TRIMETHYLAMINE

DI METHYL-KETINE 13 TETRA METHYL

PYRAZINE

DI-METHYL-KETOL v METHYL OXYETHYL KETONE

METHYL-KETOLE v (*In* 2) METHYL

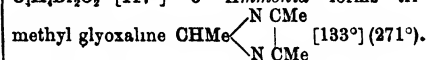
INDOLE

DI METHYL-KETONE is ACETONE

Di methyl di ketone $\text{C}_4\text{H}_8\text{O}_2$, s

$\text{CH}_3\text{COCOCH}_3$, Di acetyl (88°) S G 27 9734 S 25 at 15° Formed by heating its dicarboxylic acid (ketipic acid) either by itself or with dilute H_2SO_4 (Fittig, *B* 20, 3179, *A* 249, 200). Formed also from its mono oxim (nitroso methyl ethyl ketone) by successive treatment with NaHSO_3 and dilute acid $\text{CH}_3\text{CO Me NOH} + \text{H}_2\text{SO}_4 + \text{H}_2\text{O} = \text{CH}_3\text{CO COMe} + \text{NH}_4\text{SO}_4$ (Von Pechmann, *B* 20, 3162). Most easily prepared by saponifying methyl aceto acetic ether with dilute (3 p c) alkali, treating the product with NaNO_2 and H_2SO_4 , removing alcohol by distillation, adding dilute H_2SO_4 (20 vols of 15 p c), and distilling with steam (Von Pechmann, *B* 21, 1411). Yellow mobile liquid, smelling like acetone and quinone, miscible with alcohol and ether. Decomposed by alkalis or hot alkaline carbonates. Forms metallic derivatives of trimethyl glyoxaline with ammoniacal solutions of silver nitrate and of cuprous chloride. With alcohol it forms an unstable compound $\text{C}_4\text{H}_8\text{O}_2\cdot\text{HOEt}$, boiling at 75°. With water it forms a crystalline hydrate ($\text{C}_4\text{H}_8\text{O}_2$), 2aq, insol water, alcohol, and ether. Yields a very unstable compound with SO_2 . Combines with NaHSO_3 .

Reactions—1 With HCy it forms the nitrile of di oxy di methyl succinic acid—2 Bromine dissolved in CS_2 forms a di bromo derivative $\text{C}_4\text{H}_6\text{Br}_2\text{O}_2$ [117°]—3 Ammonia forms tri



4 When warmed with o tolylene diamine acetate it is converted into tri methyl quinoxaline $\text{C}_6\text{H}_3\text{Me}_3$, which crystallises from light petroleum in prisms [91°] (270°)—5 Aniline forms the anilide PhN Me Me NPh , which crystallises from alcohol in lustrous sulphur-yellow scales [133°], v sol ether, m sol alcohol, insol water—6 On adding dilute aqueous NaOH or Na_2CO_3 to an aqueous solution of the diketone until it becomes colourless, then at once acidifying with dilute H_2SO_4 , and extract-

ing with ether, the product is di methyl quino-
gen $\text{CH}_2\text{CO CMe CH CO CO CH}_2$, a yellowish
bitter syrup, which forms a phenyl-hydrazide
 $\text{C}_6\text{H}_5\text{N}(\text{N}_2\text{HPh})$, [205°], and is converted by excess
of alkali into *p*-xyloquinone—7 Heated with
an aqueous solution of *urea* it forms di methyl
glycolurate $\text{C}_6\text{H}_5\text{N}_2\text{O}_2$ (Franchi nont a Klobbie,
R T C 7, 251)—8 Reduced in alkaline solu-
tion to a product $(\text{CH}_2\text{CO CH(OH) CH}_2)_2$, which
instantly reduces Fehling's solution in the cold

Mono-oxim $\text{CH}_2\text{CO C(NOH) CH}_2$ *Iso*
nitroso methyl ethyl ketone [74°] (186°)
VD 351 (calc 349) Formed by treating
methyl aceto acetic ether (1 mol) with aqueous
KOH (3 mols), followed by nitrous acid (V
Meyer a Züblin, *B 11*, 322) Prisms (from
chloroform) or plates (from water), *v e* sol
alcohol, ether, and chloroform Decomposed
by heating with dilute HClAq at 140° into am-
monia and acetic acid Boiling conc HClAq
yields HOAc, hydroxylamine, and a little of the
dioxim (Schramm, *B 16*, 177) By boiling with
dilute H_2SO_4 it is split up into hydroxylamine
and the diketone (Von Pechmann, *B 20*, 3213)
Treatment with aqueous NaHSO_3 and dilute
 H_2SO_4 also forms di methyl diketone (Von Pech-
mann, *B 20*, 3162) Alkaline K_2FeCy_4 oxidises
it, even in the cold, to acetic and nitrous acids
(Gutknecht, *B 12*, 2290) On reduction by
 SnCl_2 and HCl , followed by removal of tin by
 H_2S and addition of alkali, tetra methyl pyrazine
is obtained But if the solution is allowed to stand
when acid another base is got (Braun, *B 22*,
559)

Methyl ether of the mono oxim
 $\text{CH}_2\text{CO C(OMe) CH}_2$ (125° cereasol) Colour-
less oil, lighter than water (Cereasole, *B 16*,
834)

Di oxim $\text{CH}_2\text{C(NOH) C(NOH) CH}_2$ *Me*
thyl ethyl acetoximic acid [234°] Formed
by the action of hydroxylamine on the mono
oxim (Schramm, *B 16*, 179, Auwers a V Meyer,
B 21, 3527) or on the diketone (Fittig, *A 249*,
204) Small colourless needles, insol water, *v*
sol alcohol and ether May be sublimed

Oxim phenyl hydrazide
 $\text{CH}_2\text{C(NOH) C(N}_2\text{HPh) CH}_2$ [158°] Formed
from the oxim and phenyl hydrazine (Von Pech-
mann a Wehsarg, *B 21*, 2997) Large crystals
(from alcohol) Its solution in H_2SO_4 is coloured
bluish violet by FeCl_3

Phenyl-hydrazide
 $\text{CH}_2\text{CO C(N}_2\text{HPh) CH}_2$ [138°] Formed by
running a solution of diazobenzene chloride
into one of methyl acetoacetic acid, the reaction
being completed by adding a solution of NaOAc
(Japp a Klingemann, *B 21*, 549, *A 247*, 218,
Von Pechmann, *B 21*, 1411) Yellow tables
(from benzene) Forms a yellow solution in
conc H_2SO_4

Di-phenyl di hydrazide
 $\text{CH}_2\text{C(N}_2\text{HPh) C(N}_2\text{HPh) CH}_2$ [243°] (Japp),
[239°] (Fittig) Formed by treating the mono
phenyl hydrazide with phenyl hydrazine (Japp,
A 247, 218), or by the action of phenyl-hydra-
zine on an ethereal solution of di methyl di
ketone (Fittig, *A 249*, 203, Von Pechmann,
B 20, 3164) Formed also by heating the
phenyl hydrazide of pyruvic acid (Japp a
Klingemann, *B 21*, 550) Yellowish needles,
almost insol water, alcohol, and ether, *sl* sol

chloroform, *m* sol acetone and hot benzene
Dissolves in cold conc H_2SO_4 forming a brown
solution, which after some time becomes dirty
wine red, appearing green in thin layers $\text{K}_2\text{Cr}_2\text{O}_7$
in HOAc oxidises it to the 'osotetrazone'
 $\text{CH}_2\text{C N NPh}$

$\text{CH}_2\text{C N NPh}$, which crystallises in matted red
needles [169°], insol water, sol chloroform and
benzene, *m* sol ether, *sl* sol acetone and alco-
hol, almost insol HOAc This 'osotetrazone'
is reduced to the original diphenyl dihydrazide
by warming with phenyl hydrazine HCl con-
verts the 'osotetrazone' into an 'osotriazone'
 $\text{CH}_2\text{C N}$

$\text{CH}_2\text{C N}$ > NPh, which is a very weak base,
[35°], (*c* 255°), insol water, sol alcohol and
ether (Von Pechmann, *B 21*, 2759)

DI METHYL DI-KETONE DICARBOXYLIC
 $\text{ACID C}_6\text{H}_5\text{O}_4$ *ie* $\text{CO}_2\text{H CH}_2\text{CO CO CH}_2\text{CO}_2\text{H}$ *Di-keto*
adipic acid Ketipic acid Ketipic acid Obtained by warm-
ing its ethyl ether with cold conc HClAq (Fittig
a Daumler, *B 20*, 203, *A 249*, 183) White
amorphous powder, insol or *v sl* sol cold water,
alcohol, chloroform, CS_2 , benzene, and petroleum
ether On heating alone or with dilute H_2SO_4 it
is converted into dimethyl diketone

Ethyl ether $\text{Et}_2\text{A}''$ [77°] Prepared by
warming oxalic ether with chloro-acetic ether
and amalgamated granulated zinc at 80° for
three days The mass is extracted with water
and the insoluble portion treated with dilute
 H_2SO_4 , and the solution so obtained extracted
with ether The ether is distilled off, and the
ketipic ether left recrystallised from alcohol
(Fittig a Daumler) Formed also by the action
of NaOEt and oxalic ether on acetic ether (Wis-
licenus, *B 20*, 589, *A 246*, 323) Colourless
plates or prisms, insol cold water, *v* sol ether
and chloroform, *m* sol hot alcohol, benzene,
and CS_2 Decomposed by boiling water FeCl_3
colours its alcoholic solution deep red

Reactions—1 Bromine added to a warm
solution of the ether in CS_2 forms
 $\text{CO}_2\text{Et CBr}_2\text{CO CO CBr}_2\text{CO}_2\text{Et}$ [119°], which is
converted by ammonia into oxamide and di
bromo acetamide [156°] Bromine added to a
solution of ketipic ether in cooled CS_2 forms
 $\text{CO}_2\text{Et CHBr CO CO CHBr CO}_2\text{H}$, which is de-
posited in colourless plates [70°]—2 Chlorine
passed into a boiling solution of the ether in
chloroform forms $\text{CO}_2\text{Et CCl}_2\text{CO CO CCl}_2\text{CO}_2\text{Et}$
[93°], which is also produced by chlorinating
di oxy quinone dicarboxylic ether (Böninger, *B*
22, 1285)—3 Phenyl hydrazine forms a phenyl
hydrazide

$\text{CO}_2\text{Et CH}_2\text{C(N}_2\text{HPh) C(N}_2\text{HPh) CH}_2\text{CO}_2\text{Et}$
crystallising from chloroform in light yellow
needles [160°-180°]

DI-METHYL-KETOXIM v ACETOXIM

METHYL-LEPIDONE v OXY DIMETHYL-
QUINOLINE

METHYL-PARA-LEUCANILINES v METHYL-
TRI AMIDO TRI PHENYL-METHANES

METHYL LUTIDONE v OXY TRI METHYL-
PYRIDINE

METHYL-PSEUDO-LUTIDO-STYRIL v OXY
TRI METHYL PYRIDINE

DI-METHYL-MALEIC ACID $C_4H_4O_4$, &c.**CO₂H.COMe OMe.CO₂H.** *Pyrocinchoninic acid**Di-methyl-fumaric acid.* *Di-methyl-ethylene**di-carboxylic acid* *Butylene dicarboxylic acid*

Formation—1 Its anhydride is formed by the dry distillation of cinchoninic acid $C_4H_4O_4$ (Weidel & Schmidt, *B* 12, 1151; Weidel & Brix, *M* 8, 608)—2 The anhydride is formed by distilling the mother liquor from the preparation of terebic acid by oxidation of turpentine with HNO_3 (Roser, *B* 15, 1818)—3 By heating the lactone $CH_2.C(CO_2H).CH(CO_2H).CH_2.CO$ at 180°

(Rach, *A* 234, 39)—4 Together with di-chloro di-methyl succinic acid $CO_2H.CClMe.CClMe.CO_2H$ by the action of reduced silver upon di- α -chloro propionic acid (Otto & Beckurts, *B* 18, 825)—5 By the action of reduced silver upon di-chloro di methyl succinic acid (O & B)—6 The anhydride is formed by treating either 'anti' or 'para' di-methyl-succinic acid in the fused state with bromine (Bischoff & Voit, *B* 23, 646)

Properties—The free acid is unstable, on adding an acid to its salts the anhydride is at once pptd. The sodium salt gives a dark red colouration with $FeCl_3$, and crystalline pps with $Pb(NO_3)_2$ and $HgCl_2$.

Salts— Na_2A'' aq white crystalline solid— Na_2A'' l₂ aq— CaA'' small needles, less sol hot water than cold (Roser)— CaA'' aq small white plates (O & B)— BaA'' white glistening plates, more sol cold water than hot— Ag_2A'' sparingly soluble pp, decomposed on heating into Ag_2O and the anhydride

Methyl ether Me_2A'' Colourless oil

Ethyl ether Et_2A'' (240°) Oil Formed by treating the anhydride with alcohol and HCl

Anhydride $\begin{matrix} \text{CMe.CO} \\ \parallel \\ \text{CMe.CO} \end{matrix} O$ [96°] (223°)

VD 42 (calc 43) Formed as above. Crystallises in glistening white leaflets or trimetric tables (from water), $a, b, c = 626, 1152, 1$ or $201, 392, 1$ V sol alcohol, ether, and benzene, sl sol cold water. The aqueous solution is acid and has a sweet but burning taste. Volatile with steam. Readily sublimes. Not attacked by HNO_3 . Chromic acid mixture oxidises it to acetic acid and CO_2 . Reduced by sodium-amalgam to *s*-di-methyl-succinic acid [194°] and two isomeric acids [241°] and [120°], the last acid being methyl-ethyl-malonic acid (Otto & Rössing, *B* 20, 2736). Conc $HIAq$ at 220° , and zinc filings in water or ammonia also reduce it to *s*-di methyl succinic acid. The anhydride is not attacked by PCl_5 or $AcCl$. It is dissolved by aqueous alkalis, forming the sodium salt of the acid. Potash fusion yields oxalic acid. By heating with bromine and water at 100° there is formed di-bromo-acetic acid. When it is dissolved in benzene and shaken with phenyl-hydrazine in the cold the anhydride forms the compound $CH_2.C(COON_2H_5Ph).CMe.CO.ON_2H_5Ph$, which, when shaken with $HClAq$, regenerates the anhydride, but when heated at 115° yields CO_2 .

phenyl-hydrazine, and $\begin{matrix} \text{CMe.CO.NH} \\ \parallel \\ \text{CMe.CO.NH} \end{matrix}$ [129°], which crystallises from dilute alcohol in yellow monoclinic prisms, and yields di methyl maleic acid

when warmed with $KOHAq$ or $HClAq$ (Otto & Holst, *J pr* [2] 42, 67)

Chloride $\begin{matrix} \text{CMe.COCl} \\ \parallel \\ \text{CMe.COCl} \end{matrix}$ (220°–230°) Formed

by the action of PCl_5 on the anhydride (O & H). Yellow oil. When acted upon by the Na salt it yields the anhydride. NH_3 converts it into the imide. Phenyl-hydrazine forms the compound $\begin{matrix} \text{CMe.CO} \\ \parallel \\ \text{CMe.CO} \end{matrix} N.NHPh$ [187°] isomeric with the body melting at 129° (v *supra*)

Imide $\begin{matrix} \text{CMe.CO} \\ \parallel \\ \text{CMe.CO} \end{matrix} NH$ [118°] Formed by

heating the anhydride with alcoholic NH_3 . Triclinic plates, m sol hot water, v sol alcohol. May be sublimed— $B''_2H_2PtCl_4$.

Anilide $\begin{matrix} \text{CMe.CO} \\ \parallel \\ \text{CMe.CO} \end{matrix} NPh$ [96°] Formed

by heating the anhydride with aniline at 180° . Prisms (from alcohol)

METHYL-MALONIC ACID $C_4H_6O_4$, &c.

$CH_2.CH(CO_2H)_2$ *Iso succinic acid* Mol w 118 [130°] $H C v$ 365, 100 $H C p$ 364, 800 $H F$ 218, 200 (Stohmann, Kleber, & Langbein, *J pr* [2] 40, 207) $S H$ (0°–50°) 3372 (Hess *P* [2] 35, 410)

Formation—1 By decomposing a cyano propionic acid with potash (Wichelhaus, *Z* 1867, 247, Byk, *J pr* [2] 1, 19)—2 From sodium malonic ether and MeI (Zublin, *B* 12, 1112)

Preparation—By acting on potassium a bromo propionate with pure aqueous KCy , and saponifying the product by boiling with aqueous KOH (Cohn, *A* 251, 335)

Properties—Long prisms or tables (by sublimation). Decomposed on distillation into CO_2 and propionic acid. $FeCl_3$ gives no pp in neutral solutions. Fuming HNO_3 decomposes it into tri nitro ethane, CO_2 and acetic acid (Franchimont, *R T C* 5, 281). Electrolysis of a concentrated solution of its potassium salt yields hydrogen, CO_2 , and oxygen.

Salts— $NaHA''$ aq— Na_2A'' 2aq— KHA'' — K_2A'' aq— CaA'' aq— CaA'' aq Solubility Miczynsky (*M* 7, 269)— BaA'' 2aq— ZnA'' 3aq— PbA'' aq— Ag_2A'' heavy granular pp gradually becoming crystalline

Methyl ether Me_2A'' (179°) SG 18 1107. When treated with HNO_3 (SG 15) it gives a small quantity of $CH_3.C(NO_2)(CO_2Me)_2$.

Ethyl ether Et_2A'' (196.5° cor) (Krestownikoff, *B* 10, 409), (199°) (Perkin, *C J* 45, 510) SG $\frac{1}{2}$ 1021 (Conrad & Bischoff, *A* 24, 146), $\frac{1}{2}$ 10213, $\frac{1}{2}$ 10130 (P)

Di-amide $CH_2.CH(CO NH_2)_2$ [206°]. Formed, together with methyl carbonate, by treating the compound $CH_2.C(NO_2)(CO_2Me)_2$ (v *supra*) with ammonia (Franchimont, *R T. C.* 8, 286) V sl sol alcohol

Di-methyl-di-amide $CH_2.CH(CO NHMe)_2$ [154°] Formed by the action of methylamine on the ether (Franchimont, *R T C* 4, 204). Small needles (from benzene), v sol water and alcohol, sl sol ether and benzene. Fuming HNO_3 decomposes it, forming CO_2 and N_2O

Di-methyl-malonic acid $\text{CMe}_2(\text{CO}_2\text{H})_2$. *Isopyrotartaric acid* Mol w 132 [186°] H C 515,300 H F 230,700 (Stohmann, Kleber, a Langbein, *J pr* [2] 40, 208) S H (0°-50°) 310 (Hess, *P* [2] 35, 410)

Formation—1 By heating bromo isobutyric acid with KCy and decomposing the resulting nitrile with potash or HClAq (Markownikoff, *B* 6, 1440, *A* 182, 324)—2 By boiling di methyl-barbituric acid with potash (Conrad a Guthzeit, *B* 14, 1644)—3 From methyl malonic acid, NaOEt, and MeI (Thorne, *C J* 39, 543)—4 By oxidising β acetyl di a methyl propionic acid with nitric acid (3 pts of S G 14 and 1 pt water) (Anschutz, *A* 247, 105)

Properties—Transparent four sided monoclinic prisms, sl sol alcohol, v sol water and ether Sublimes in white needles, even at 100° Split up on melting into CO and isobutyric acid Not attacked by boiling dilute HNO_3 , and scarcely at all by boiling chromic acid mixture

Salts—*NaA" small efflorescent needles, sl sol water—*BaA" stellate groups of thin needles—*CaA" nodules, m sol cold water, deposited on warming its solution—*MgA" crystalline, v sol water—*PbA" aq insoluble pp, changing to shining scales on boiling—ZnA" aq S 68 at 24° Monoclinic pyramids (Thorne)—ZnA" 3aq S 94 (Markownikoff)—AgA" small needles, insol water

Ethyl ether EtA" (194.5°) (Thorne), (196.5° cor) (Peikin, *C J* 45, 511) S G $\frac{1}{2}$ 9965 (T), $\frac{1}{15}$ 10015, $\frac{2}{25}$ 9936 (P) M M 9 268 at 14.4°

Amide $(\text{CH}_3)_2\text{C}(\text{CO NH})_2$ [197°] From the ether and alcoholic NH_3 at 120° (Thorne)

Amic acid $\text{CO}_2\text{H CMe}_2\text{CO NH}$ [85°] A product of the oxidation of mesitylic acid

$\text{CMe} \begin{cases} \text{CH}_3 \text{ CMe CO}_2\text{H} \\ \text{CO NH} \end{cases}$ by KMnO_4 and H_2SO_4

(Hanner, *B* 15, 580) At 130° it decomposes, giving off CO_2 Boiling KOHAq forms NH_3 , and di methyl malonic acid The potassium salt KA' 2aq crystallises in prisms, v e sol water, m sol alcohol

Di methyl di-amide $\text{CMe}(\text{CO NHMe})_2$ [123°] Long needles (from benzene), v e sol water and alcohol, sl sol ether, m sol benzene (Franchimont, *R T C* 4, 206) Fuming HNO_3 attacks it, forming N_2O , methyl nitrate, and dimethylmalonic acid

Tetra methyl di amide $\text{CMe}_2(\text{CO NMe}_2)_2$ [80°] (276°) Long prisms (from ligroin), v e sol water, alcohol, and benzene Fuming HNO_3 forms di methyl malonic acid and di methyl nitramine

DIMETHYL-MALONYL-UREA v DI METHYL-BARBITURIC ACID

METHYL-MANDELIC ACID v Methyl derivative of MANDELIC ACID and OXY-TOLYL ACETIC ACID

Tetra methyl mandelic acid v OXY DURYL ACETIC ACID

METHYL-MELAMINE v Methyl cyanuramide in the article CYANIC ACIDS

METHYL MERCAPTAN CH_3S v CH_3SH Methyl sulphhydrate (5.8°) at 752 mm (Klason, *B* 20, 3407), (20°) (Gregory, *A* 15, 239), (21°) (Obermeyer, *B* 20, 2919) H F p 5,950 H F v 6,370 (Thomsen, *Th*) Occurs in human

excrement (Nencki, *M* 10, 863) Obtained by distilling KMeSO_4 with KHS Prepared by diluting with ice a cold mixture of MeOH (500 cc) and H_2SO_4 (750 cc), adding (2.75 kilos of) Na_2CO_3 10aq, and evaporating until most of the Na_2SO_4 has separated The mother liquor is mixed with a solution of potash (500 g) in water (1,000 cc) previously saturated with H_2S The mixture is distilled from a water bath, and the gases evolved are passed first through a concentrated aqueous solution of KOH (50 g) and then into a solution of KOH (350 g) in water (700 cc) A small quantity of lead acetate is added to the last solution to ppt H_2S , and the methyl mercaptan is then liberated by HCl, dried, and distilled The yield is fair (200 g of MeSH and 40 g of Me_2S) (Klason, *B* 20, 3407)

Properties—Thin, colourless, highly refractive liquid with very repulsive odour Yields a crystalline hydrate

Salts— $\text{Hg}(\text{SMe})_2$ [175°] Obtained by treating HgO with MeSH, or by passing the gas through an aqueous solution of HgCy_2 , V sl sol water— $\text{Pb}(\text{SMe})_2$ minutetables— $\text{Bi}(\text{SMe})_3$ minute yellow needles— AgSMe yellow crystalline pp

Reference—PER CHLORO METHYL MERCAPTAN **DI-METHYL-MESIDINE** $\text{C}_{11}\text{H}_{11}\text{N}$ v $\text{C}_6\text{H}_5\text{NMe}_2$ (214°) S G 908 Formed by methylation of mesidine (Hofmann, *B* 5, 718, Klobbie, *R T C* 6, 33) By treatment with H_2SO_4 and HNO_3 (S G 1.5) it is converted into $\text{C}_6\text{Me}_5(\text{NO})$, NMeNO [138°]— $\text{B}'_2\text{H}_2\text{PtCl}_4$

METHYL METHANE v ETHANE
Di methyl methane v PROPANE
Tri methyl methane v ISO BUTANE
Tetra methyl-methane v PENTANE

METHYL PENTAMETHENYL TRIHYDRIDE DICARBOXYLIC ACID $\text{C}_8\text{H}_{10}\text{O}_4$ v $\text{CO H CH} < \begin{matrix} \text{C}(\text{CH}_3) \\ \text{CH}_2\text{CH}_2 \end{matrix} > \text{CO}_2\text{H}$ [188°] Ob-

tained by saponifying its ether which is among the products resulting from the distillation of di methyl butylene diketone dicarboxylic (di acetyl adipic) ether with KOH (Perkin, jun, *C J* 57, 227, 233) Glistening needles or groups of plates, v sol hot water, alcohol, and acetone, sl sol cold light petroleum, benzene, CS_2 , and chloroform

Reactions—1 Its aqueous solution decolourises bromine in the cold and on evaporating to a syrup it gives off HBr—2 HBrAq in sealed tubes at 110° forms $\text{C}_8\text{H}_{11}\text{Br}$, a light brown oil, probably bromo methylpentamethylene—3 Sodium-amalgam has no action

Salts—*(NH_4)A" gelatinous— AgA'' : heavy white pp which darkens in daylight— AgHA'' slender needles, v sol hot water

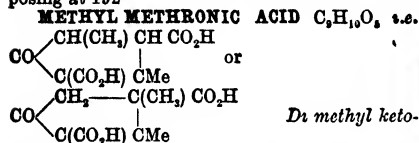
METHYL PENTAMETHENYLYL METHYL KETONE TRIHYDRIDE $\text{C}_8\text{H}_{10}\text{O}$ v

$\text{CH}_3 \begin{cases} \text{CH}_2\text{CMe} \\ \text{CH}_2\text{C CO CH}_3 \end{cases}$ Methyl dihydropentene

methyl ketone (191°) Formed by the action of boiling alcoholic potash on di-methyl butylene diketone dicarboxylic (di acetyl adipic) ether (Marshall a Perkin, jun, *C J* 57, 232, 244) Colourless mobile oil, smelling like peppermint Lighter than water V sol alcohol and ether Readily reacts with phenyl hydrazine. May be

reduced to the alcohol $C_6H_{13}O$, whence HI at 250° yields C_6H_{13} .

Oxim $C_6H_{12}(NOH)$ [85°] Formed by the action of hydroxylamine hydrochloride on the ketone dissolved in methyl alcohol. Thick colourless prisms, v sol alcohol, ether, acids, and alkalis. May be distilled with slight decomposition — $(C_6H_{13}NO)_2H_2PtCl_6$ tufts of orange crystals, v sol boiling alcohol, decomposing at 192° .



pentene dicarboxylic acid [198°] Formed by heating acetoacetic ether, sodium pyruvate, and Ac_2O at 140° , and saponifying the resulting ether with baryta or NaOH (Fittig, *A* 250, 195). White needles, v sol HOAc, ether, hot alcohol, and hot water, sl sol cold water, CS_2 , and benzene, almost insol petroleum ether. At 250° it is split up into CO_2 and methyl uvic acid $C_5H_{10}O_4$ [98°], which on prolonged boiling yields $\begin{array}{c} \text{CH}(\text{CH}_3) \text{CH}_2 \\ \text{CO} \begin{array}{l} \text{CH} = \text{CMe} \\ \text{CH} = \text{CMe} \end{array} \end{array}$ or $\begin{array}{c} \text{CH} \text{CH CH}_2 \\ \text{CO} \begin{array}{l} \text{CH} = \text{CMe} \\ \text{CH} = \text{CMe} \end{array} \end{array}$, an oil, boiling at 119° .

Salts — BaA' 2aq small needles, v e sol water — CaA' 3aq small prisms — AgA' silky nodules, sl sol water.

Mono-ethyl ether EtHA" Obtained by boiling the diethyl ether with alcoholic potash. Yellow syrup, sl sol water — Ba(EtA')₂ aq yellow gummy pp, v e sol water and alcohol — Ca(EtA')₂ 2aq needles, m sol water, v sol hot alcohol — AgEtA" flocculent pp, sl sol water, v sol alcohol and ether.

Di-ethyl ether EtA" (280° uncor) Heavy oil, miscible with alcohol and ether.

METHYL TETRAMETHYLENE C_6H_{10} *v.e.*
 $\text{CH}_2 \begin{array}{c} \text{CH}_2 \\ \text{CH}_2 \end{array} \text{CHMe}$ (c 40°) Formed by the action of sodium on α di bromo pentane dissolved in toluene (Colman & Perkin, *C J* 53, 201). Very volatile oil. Does not combine with conc HIAq.

METHYL PENTAMETHYLENE C_6H_{12} *v.e.*
 $\begin{array}{c} \text{CH}_2 \text{CH}_2 \\ \text{CH}_2 \text{CH}_2 \end{array} \text{CHMe}$? Formed by the action of finely divided sodium on α di bromo hexane dissolved in toluene (Perkin, *C J* 53, 214). Oil, not attacked by HI.

METHYL-PENTAMETHYLENE CARBOXYLIC ACID $C_6H_{12}O_4$ *v.e.*
 $\text{CH}_2 \begin{array}{c} \text{CH}_2 \text{CHMe} \\ \text{CH}_2 \text{CH}_2 \end{array}$ (220°) SG $\frac{1}{12}$ 1.0205,

$\frac{2}{3}$ 1.0174, $\frac{2}{3}$ 1.0144 MM 6.914 Formed by heating methyl pentamethylene dicarboxylic acid a little above its melting point (Colman & Perkin, jun, *C J* 53, 194). Colourless oil. Bromine at 110° attacks it with evolution of HBr — BaA' syrup — AgA' white amorphous pp.

Methyl-pentamethylene dicarboxylic acid
 $C_6H_{12}O_4$ *v.e.* $\text{CH}_2 \begin{array}{c} \text{CH}_2 \text{CHMe} \\ \text{CH}_2 \text{C}(\text{CO}_2\text{H})_2 \end{array}$ [175°] Obtained by boiling its ether with alcoholic potash

(C & P) Prisms (from ether), or plates (from water), v sol alcohol, ether, and hot water, m. sol cold water — AgA' white pp.

Ethyl ether EtA" (244°) Obtained from $\text{CH}_3\text{CHBrCH}_2\text{CH}_2\text{CH}_2\text{Br}$, malonic ether and NaOEt. Thick oil.

METHYL-HEXAMETHYLENE CARBOXYLIC ACID $C_6H_{14}O_4$ *v.e.*

$\text{CH}_2 \begin{array}{c} \text{CH}_2 \text{CHMe} \\ \text{CH}_2 \text{CH}_2 \end{array} \text{CH CO}_2\text{H}$ *o Toluic acid hexahydrate?* (236°) SG $\frac{1}{4}$ 1.0079, $\frac{1}{2}$ 1.0033, $\frac{2}{3}$ 9966 MM 7.975 Formed by decomposing the dicarboxylic acid by heat (Perkin, jun, *C J* 53, 208, 213). It is also one of the products of the electrolysis of methyl hexamethylene methyl ketone carboxylic ether by alcoholic potash. Colourless oil — AgA' white pp.

Methyl hexamethylene dicarboxylic acid
 $C_6H_{14}O_4$ *v.e.* $\text{CH}_2 \begin{array}{c} \text{CH}_2 \text{CHMe} \\ \text{CH}_2 \text{CH}_2 \end{array} \text{C}(\text{CO}_2\text{H})_2$ [147°].

Obtained by hydrolysing its ether with alcoholic potash (Perkin, *C J* 53, 207). Crystalline powder, v sol ether, alcohol, and hot water, sl sol cold water — AgA' white amorphous pp.

Ethyl ether EtA" (c 263°) Obtained from $\text{CH}_3\text{CHBrCH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{Br}$, malonic ether, and NaOEt (Perkin, jun, *C J* 53, 206). Thick oil, with unpleasant odour.

METHYL TETRAMETHYLENE KETONE *v.* **TETRAMETHYLENYL METHYL KETONE**

METHYL-TRI METHYLENE KETONE CARBOXYLIC ACID *v.* **TRI METHYLENYL METHYL KETONE CARBOXYLIC ACID**

METHYL PENTAMETHYLENE METHYL KETONE *v.* **METHYL PENTAMETHYLENYL METHYL KETONE**

TETRA-METHYL- ν -TRIMETHYLENE-DI-PYRROLE $C_{15}H_{22}N_2$ *v.e.*

$\text{HC CMe} \begin{array}{c} \text{N C}_2\text{H}_4 \text{N} \\ \text{CMe CH} \end{array} \text{CMe CH}$ [77°] Formed by heating acetonyl acetone (2 mols) and trimethylene diamine (1 mol) in conc alcoholic solution at 120° . Crystalline solid. Sol alcohol and ether, insol water (Paal & Schneider, *B* 19, 3157).

DI METHYL-TRI METHYLENE TRI SULPHONE $C_6H_4(\text{CH}_3)_2\text{S}_3O_6$ [330° - 340°] Two bodies of this formula are produced when 1 g tri methylene tri sulphone in 10 cc of 5 p.c. caustic soda is heated with an equal volume of methyl iodide. These two substances crystallise together in glittering needles. By further methylation the hexamethyl derivative is formed (E. Baumann & R. Camps, *B* 23, 72).

Tetra methyl di methylene disulphone
 $\text{CMe}_2 \begin{array}{c} \text{SO}_2 \\ \text{SO}_2 \end{array} \text{CMe}_2$ *Di isopropylidene di sulphone* [220° - 225°] Obtained from acetone (1 pt) by heating it with P_2S_5 (1 pt) at 125° for 7 hours, distilling with steam, and oxidising the resulting 'duplo'-thio acetone $\text{S}_2(\text{CMe}_2)_2$ with KMnO_4 (Autenrieth, *B* 20, 373). Slender white needles, v sol alcohol and ether, sl sol hot water. Not attacked by HNO_3 and H_2SO_4 , nor by boiling dilute aqueous KOH.

METHYL PENTAMETHYLENYL METHYL-CARBINOL $C_6H_{14}O$ *v.e.*
 $\text{CH}_2 \text{CHMe} \begin{array}{c} \text{CH CH}(\text{OH}) \text{CH}_2 \\ \text{CH}_2 \text{CH}_2 \end{array}$ (180°) Obtained by boiling its ether with alcoholic potash

tained by reducing the corresponding ketone in ethereal solution with sodium (Marshall a Perkin, jun, *C J* 57, 247) Colourless liquid, with strong odour of menthol, sl sol water, v sol alcohol and ether. On boiling with Ac_2O it yields a strongly refracting acetyl derivative with agreeable odour. Excess of solution of hydriodic acid (S G 196) forms a heavy oily iodide CH_2CHMe

$\text{CH}_2\text{CH}_2 \begin{matrix} \diagup \\ \text{CH} \end{matrix} \begin{matrix} \diagdown \\ \text{CH} \end{matrix} \text{CH}_2$ (155°–160° at 90 mm), which smells like *sec* hexyl iodide, and is slightly decomposed on distillation

METHYL-HEXAMETHYLENYL METHYL-CARBINOL $\text{C}_8\text{H}_{18}\text{O}$ *ze*

$\text{CH}_2 \begin{matrix} \diagup \\ \text{CH}_2\text{CHMe} \\ \diagdown \\ \text{CH}_2\text{CH}_2 \end{matrix} \begin{matrix} \diagup \\ \text{CH} \end{matrix} \begin{matrix} \diagdown \\ \text{CH} \end{matrix} \text{CH}(\text{OH})\text{CH}_3$ *Hexahydride of o tolyl methyl carbinol* (195°–200°) Formed by reducing tetrahydro tolyl methyl ketone in ethereal solution with sodium (Kipping a Perkin, jun, *C J* 57, 22) Thick, colourless liquid, smelling like menthol, sl sol water, miscible in alcohol and ether. When mixed with conc HIAq it dissolves with evolution of heat and separation of the corresponding iodide

METHYL-PENTAMETHYLENYL METHYL

KETONE $\text{C}_8\text{H}_{16}\text{O}$ *ze* $\text{CH}_2 \begin{matrix} \diagup \\ \text{CH}_2\text{CHMe} \\ \diagdown \\ \text{CH}_2\text{CH CO CH}_3 \end{matrix}$

(171°) S G $\frac{1}{4}$ 9222, $\frac{1}{10}$ 9174, $\frac{1}{15}$ 9136, $\frac{2}{35}$ 9070 M M 8019 Obtained as one of the products of the hydrolysis of its carboxylic ether by alcoholic potash (Colman a Perkin, *C J* 53, 198) Colourless oil, smelling of peppermint. Readily combines with phenyl hydrazine and with hydroxylamine. Dissolves with difficulty in a solution of NaHSO_3 .

Methyl-hexamethylenyl methyl ketone

$\text{CH}_2 \begin{matrix} \diagup \\ \text{CH}_2\text{CHMe} \\ \diagdown \\ \text{CH}_2\text{CH}_2 \end{matrix} \begin{matrix} \diagup \\ \text{CH} \end{matrix} \begin{matrix} \diagdown \\ \text{CH} \end{matrix} \text{CO CH}_3$ *o Tolyl methyl ketone hexahydride* (198°) Obtained as one of the products of the hydrolysis of its carboxylic ether by alcoholic potash (Perkin, *C J* 53, 213) Colourless oil, smelling, when in small quantities like new mown hay

METHYL TRIMETHYLENYL METHYL

KETONE CARBOXYLIC ACID $\text{C}_8\text{H}_{16}\text{O}_2$ *ze*

$\text{CHMe} \begin{matrix} \diagup \\ \text{C}(\text{CO}_2\text{H}) \end{matrix} \begin{matrix} \diagdown \\ \text{CO CH}_3 \end{matrix}$ *Propylene acetoacetic acid* Obtained by saponifying its ether, which is produced by treating acetoacetic ether with NaOEt and propylene bromide (Perkin, jun, *B* 17, 1443) Thick oil, splitting off CO_2 when heated—AgA' amorphous powder, sl sol water

Ethyl ether EtA' (210°–215°) at 720 mm

Methyl-pentamethylenyl methyl ketone

carboxylic ether $\text{CH}_2 \begin{matrix} \diagup \\ \text{CH}_2\text{CHMe} \\ \diagdown \\ \text{CH}_2\text{C}(\text{CO}_2\text{Et}) \end{matrix} \begin{matrix} \diagup \\ \text{CH} \end{matrix} \begin{matrix} \diagdown \\ \text{CH} \end{matrix} \text{CO CH}_3$

(238°) Obtained by the action of *ad* di bromo pentane on sodium acetoacetic ether (Colman a Perkin, jun, *C J* 53, 197) Thick, colourless oil. On hydrolysis with alcoholic potash it yields the corresponding ketone as well as the acid

Methyl-hexamethylenyl methyl ketone carboxylic ether

$\text{CH}_2 \begin{matrix} \diagup \\ \text{CH}_2\text{CHMe} \\ \diagdown \\ \text{CH}_2\text{CH}_2 \end{matrix} \begin{matrix} \diagup \\ \text{C}(\text{CO}_2\text{Et}) \end{matrix} \begin{matrix} \diagdown \\ \text{CO CH}_3 \end{matrix}$ *Hexa*

hydro-o tolyl methyl ketone carboxylic ether (257°) Formed by the action of di bromo-hexane $\text{CH}_3\text{CHBrCH}_2\text{CH}_2\text{CH}_2\text{CH}_3$ and NaOEt on acetoacetic ether (Perkin, *C J* 53, 212) Colourless oil with disagreeable odour. Yields on hydrolysis methyl hexamethylenyl methyl ketone as well as its carboxylic acid

METHYL-MORPHINE v CODEINE

(a) **METHYL-NAPHTHALENE** $\text{C}_{11}\text{H}_{10}$ *ze*

$\text{C}_6\text{H}_5 \begin{matrix} \diagup \\ \text{CMe CH} \\ \diagdown \\ \text{CH CH} \end{matrix}$ Mol w 142 (242° i V)

(S), (232°) (F a R) S G $\frac{1}{15}$ 1029 Occurs, together with (B) methyl naphthalene, in the fraction of coal-tar oil boiling between 200° and 300° (Schulze, *B* 17, 844)

Formation—1 By the action of sodium on a mixture of MeI and (a) bromo naphthalene (Fittig a Remsen, *A* 155, 114)—2 By distilling colophony, gum benzoin, or aldehyde resin with zinc-dust (Ciamician, *B* 11, 269, *M* 1, 193)—3 By distilling (a)-naphthyl acetic acid with lime (Boessneck, *B* 16, 1546)

Properties—Colourless liquid, not solid at -18° Boiling conc HNO_3 oxidises it to (a) naphthoic acid

Picric acid compound

$\text{C}_{11}\text{H}_9\text{C}_6\text{H}_3(\text{NO}_2)_3\text{OH}$ [117°] Long, slender orange needles (from alcohol)

(B) - **Methyl-naphthalene** C_{10}H (CH_3) *ze*

$\text{C}_6\text{H}_5 \begin{matrix} \diagup \\ \text{CH CMe} \\ \diagdown \\ \text{CH CH} \end{matrix}$ [33°] (242° i V) Occurs in

the fraction of coal tar oils boiling at 200°–300° The oil is shaken with NaOH and with H_2SO_4 to remove phenols and bases, and fractionally distilled. By cooling the fraction 239°–242° to 0° the (B) methyl naphthalene separates out, leaving the liquid (a) methyl naphthalene (Schulze, *B* 17, 842, 1203, Reingruber, *A* 206, 367) Could not be obtained from (B) bromo naphthalene MeI and Na (Brunel, *B* 17, 1179) White plates, insol water, sol alcohol, ether, and benzene, volatile with steam. Completely decomposed on oxidation. Yields, on chlorination at 245°, $\text{C}_{10}\text{H CH}_2\text{Cl}$ [47°]

Picric acid compound [115°], yellow needles

Di-methyl-naphthalene $\text{C}_{12}\text{H}_{12}$ + e $\text{C}_{10}\text{H}_8\text{Me}_2$ (110° at 6 mm), (c 265°) at 760 mm S G $\frac{1}{10}$ 10283, $\frac{1}{15}$ 10199 (C a C), $\frac{2}{20}$ 10176 (G), $\frac{1}{14}$ 10180 (Nasini, *G* 15, 84) R₂₀ 87 25 (N)

Formation—1 From di bromo naphthalene [81°], sodium, and MeI (Mono, *B* 13, 1517)—2 By heating santolin, or the dimethyl naphthol derived therefrom, to redness with zinc dust (Cannizzaro a Carnelutti, *G* 12, 410)—3 By heating dimethyl naphtholhydride with sulphide of phosphorus (Cannizzaro, *G* 13, 393)

Preparation—Di bromo-naphthalene dissolved in toluene is heated with MeI and sodium, the product is heated to 150° to remove MeI and toluene, and the residue extracted with ether. The extract is fractionally distilled, and the fraction 260°–270°, after rectification under 12 mm pressure, is heated with sodium at 100° to remove any unaltered di bromo naphthalene. The di methyl naphthalene is finally obtained pure by means of its compound with picric acid (Giovannozzi, *G* 12, 147)

Properties — Colourless, highly refractive oil, not solid at -18° . With bromine it appears to form $C_{12}H_8Br_2$ and $C_{12}H_7Br$.

Picric acid compound [139°] Orange needles, sol alcohol and ether

Hexahydride $C_{12}H_{18}$ SG 194 922 R_∞ 86 14 (Nasim & Bornheimer, G 15, 93) Formed by treating the di methyl naphthalene with HIAq and phosphorus (Zucco, G 15, 81)

Di-methyl-naphthalene $C_{12}H_{12}$ (265°) Occurs in coal tar (Emmert & Reingruber, A 211, 365) Not solid above -18°

Picric acid compound
 $C_{12}H_{12}C_6H_2(NO_2)_3OH$ [118°] Orange prisms

Trimethyl-naphthalene $C_{12}H_9$, i.e. $C_{10}H_7Me_2$ (275°) Obtained (according to Maschke, C C 1886, 824) by heating di methyl-(β) naphthylamine methylo iodide at 320° , and distilling the resinous product with steam

Picric acid compound [133°]

References — BROMO- and CHLORO METHYL-NAPHTHALENES

(α)-**METHYL-NAPHTHALENE SULPHONIC ACID** $C_{10}H_8MeSO_3H$ Obtained by sulphonating (α) methyl naphthalene (Fittig & Remsen, A 155, 115) — BaA_2 , sl sol water

(β)-**Methyl-naphthalene sulphonic acid** $C_{10}H_8MeSO_3H$ Obtained by sulphonating (β) methyl naphthalene (Reingruber, A 206, 377) Syrup — BaA_2 , amorphous mass

Di-methyl-naphthalene sulphonic acid $C_{12}H_8Me_2SO_3H$ Obtained by the action of H_2SO_4 at 120° on the di methyl naphthalene derived from di bromo naphthalene [81°] (Giovannozzi, G 12, 147) Deliquescent scales — KA_2aq iridescent laminae

Di-methyl-naphthalene sulphonic acid $C_{10}H_8Me_2SO_3H$ Obtained by heating the di-methyl naphthalene of coal tar with fuming H_2SO_4 (Emmert & Reingruber, A 211, 365) — BaA_2 , sl sol water

Di-methyl-naphthalene di-sulphonic acid $C_{10}H_6Me_2(SO_3H)_2$ Formed at the same time as the preceding, from which it differs in having an easily soluble barium salt (E & R)

METHYL-NAPHTHAQUINOLINE v **METHYL-NAPHTHOQUINOLINE**

(In 2) **METHYL (α)-NAPHTHINDOLE**

$C_{13}H_{11}N$ i.e. $C_{10}H_7\langle\begin{smallmatrix} CH \\ NH \end{smallmatrix}\rangle CMe$ [132°] Obtained by heating (1 pt of) the (α) naphthyl hydrazide of acetone with $ZnCl_2$ (2 pts) at 180° (Schlieper, A 239, 237) Slender needles (from water), v sol alcohol and ether Colours pine wood, acidified by HCl, bluish violet $FeCl_3$ imparts a cherry red colour to its solution in HOAc, and on adding water a pp is formed

Picrate [168°], dark red needles (from hot benzene)

(In 2) **Methyl-(β)-naphthindole**

$C_{10}H_7\langle\begin{smallmatrix} CH \\ NH \end{smallmatrix}\rangle CMe$ (314° – 320° at 223 mm) Prepared by heating the (β) naphthyl hydrazide of acetone with $ZnCl_2$ at 175° , dissolving the fused mass in water, and extracting with ether (Schlieper, A 236, 182) Liquid, v sl sol water, v sol alcohol, ether, and benzene Colours acidified pine-wood violet $NaNO_2$ added to the acetic acid solution gives a brown pp

Picrate [176°]; reddish brown needles (from hot benzene).

Dihydride $C_{12}H_{12}$, i.e. $C_{10}H_7\langle\begin{smallmatrix} CH \\ NH \end{smallmatrix}\rangle CHMe$.

(190° – 200° at 20 mm) Obtained by treating methyl (β) naphthindole with zinc dust and HClAq, until it ceases to give the pine wood reaction, then adding NaOH and extracting with ether Yellow oil, which in an ethereal solution exhibits blue fluorescence Forms soluble crystalline salts with mineral acids Reduces $AgNO_3$ on warming Gives an oily nitrosamine

(In 1, 2)-**Di-methyl (α) naphthindole** $C_{13}H_{13}N$ i.e. $C_{10}H_7\langle\begin{smallmatrix} NH \\ CMe \end{smallmatrix}\rangle CMe$ [150°] Formed by heat

ing bromo acetyl propionic (bromo levulic) acid $CH_2COCHBrCH_2CO_2H$ (2 pts) with (α) naphthylamine (7 pts) (Wolff, B 21, 3365) White granules or prisms (from alcohol), v sol ether and benzene, m sol cold alcohol and HOAc, insol water Its solution in conc HClAq is pptd by water $FeCl_3$ colours its solution in HOAc cherry red, whilst $K_2Cr_2O_7$ give a deep blue colour

(In 1, 2) **Di-methyl-(β)-naphthindole**

$C_{10}H_7\langle\begin{smallmatrix} CH \\ CH \end{smallmatrix}\rangle\begin{smallmatrix} C \\ C \\ NH \end{smallmatrix}\begin{smallmatrix} CMe \\ CMe \\ CMe \end{smallmatrix}$ or $C_{10}H_7\langle\begin{smallmatrix} C \\ CH \end{smallmatrix}\rangle\begin{smallmatrix} C \\ CH \\ NH \end{smallmatrix}\begin{smallmatrix} CMe \\ CMe \\ CMe \end{smallmatrix}$ [132°] (above

360°) Prepared by the action of bromo acetyl propionic acid on (β) naphthylamine (Wolff, B 21, 3363) Plates (from alcohol), v sol ether and benzene, m sol cold alcohol and HOAc, insol water $FeCl_3$ colours its solution in boiling HOAc green The concentrated alcoholic solution exhibits violet fluorescence

Picrate [175°], dark brown needles

(In 1, 2)-**Di-methyl-(β)-naphthindole**

$C_{10}H_7\langle\begin{smallmatrix} CMe \\ NH \end{smallmatrix}\rangle CMe$ [126°] Possibly identical with the preceding body Formed by heating (In 1, 2) methyl (β) naphthindyl acetic acid at 210° (E Fischer & Steche, A 242, 370) Six sided plates, insol water, v sol alcohol and HOAc $FeCl_3$ gives a blue colouration to its solution in HOAc Yields a crystalline nitrosamine Reacts with MeI forming di methyl (β)-naphtho quinoline dihydride

Picrate dark red crystals

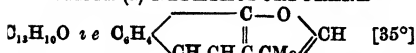
Dihydride $C_{11}H_{13}N$ i.e.

$C_{10}H_7\langle\begin{smallmatrix} CHMe \\ NH \end{smallmatrix}\rangle CMe$ Obtained by reducing the di methyl (β) naphthindole with zinc-dust and HCl oil, turning red on oxidation Forms a crystalline platinumchloride, decomposed by boiling water

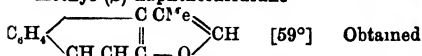
(In 1, 2)-**METHYL-(β) NAPHTHINDYL ACETIC ACID** $C_{11}H_{13}NO_2$, i.e.

$C_{10}H_7\langle\begin{smallmatrix} CMe \\ NH \end{smallmatrix}\rangle CCH_2CO_2H$ [210°] **Di methyl naphthindole α-carboxylic acid** Formed by heating the (β) naphthyl hydrazide of β acetyl-propionic ether with $ZnCl_2$ at 135° (Steche, A 242, 368) Small crystals containing $\frac{1}{2}C_6H_6O$ (from acetone), v sl sol water, benzene, and chloroform, v sol alcohol, ether, acetone, and HOAc Loses CO_2 on heating — AgA^+ white pp which yields a silver mirror on boiling with water

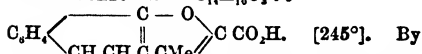
METHYL-NAPHTHOCUMARIN v **Anhydride of OXY-NAPHTHYL CROTONIC ACID**

METHYL-(α) NAPHTHOFURFURANE

[298°] Obtained by distilling its carboxylic acid alone or with KOH (Hantzsch a Pfeiffer, *B* 19, 1304) Very volatile with steam Reduces $AgNO_3$ on boiling Its solution in H_2SO_4 becomes green, and finally violet, on warming, on dilution with water it becomes green again

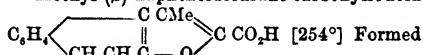
Methyl (β) naphthofurfurane

by distilling its carboxylic acid alone or with soda lime (H a P) Resembles the preceding isomeride

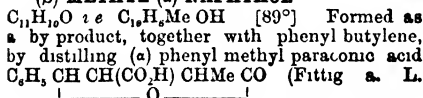
METHYL - (α) - NAPHTHOFURFURANE**CARBOXYLIC ACID $C_{11}H_{10}O_2$ i.e.**

the action of chloro aceto acetic ether on sodium (α) - naphthol there is formed $C_{11}H_9O \cdot CHAC \cdot CO_2Et$, and this ether is then mixed with H_2SO_4 , and the product poured into water The pnd ether is then saponified by alcoholic potash (Hantzsch a Pfeiffer, *B* 19, 1301) Needles (from HOAc), which may be sublimed, v sl sol alcohol and ether, almost insol water

Ethyl ether EtA' [108°] Flat needles (from alcohol), v sol ether and hot alcohol

Methyl-(β)-naphthofurfurane carboxylic acid

in the same manner as the preceding isomeride, which it greatly resembles, by using sodium (β) - naphthol (H a P) —NaA' 4aq blue fluorescent needles

(β) METHYL (α) NAPHTHOL

Liebmann, *A* 255, 263) Long slender elastic needles, v sl sol water $FeCl_3$ gives a white pp Bleaching powder produces a green pp which becomes yellow On distillation with zinc dust (β) methyl naphthalene is formed

(β)-Methyl-(α) naphthol $C_{11}H_9Me \cdot OH$ [92°] Formed by distilling (β)-phenyl methyl paracomic acid (Fittig a Liebmann, *A* 255, 272) Long elastic white needles (from boiling water), volatile with steam $FeCl_3$ gives a white pp which becomes yellow on standing Bleaching powder forms a green pp, quickly becoming yellow On distillation with zinc-dust it yields (β) methyl naphthalene

Di methyl naphthol $C_{12}H_{12}Me_2 \cdot OH$ [136°]

Obtained by heating santonic or isosantonous acid with $Ba(OH)_2$ above 360° The product is dissolved in water, and on passing CO_2 through the solution di-methyl naphthol is pnd together with $BaCO_3$, which is removed by $HClAq$ (Cannizzaro a Carnelutti, *G* 12, 406) Glistening needles (from alcohol), v sl sol water, sol alcohol and ether Begins to sublime at 100°

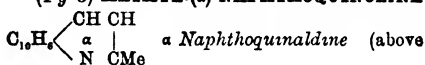
Chromic acid in HOAc oxidises it to $C_{12}H_{10}O_2$, crystallising in tables [105°], which may be reduced to di methyl naphthol by HI and phosphorus On heating to redness with zinc dust it yields di methyl naphthalene identical with that obtained from di bromo naphthalene [81°]

Methyl ether $C_{12}H_{11}OMe$ [68°] Prisms, sol alcohol and ether

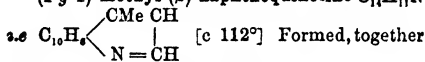
Ethyl ether $C_{12}H_{11}OEt$ Viscid liquid

Acetyl derivative $C_{12}H_{11}OAc$ [78°]. Laminae

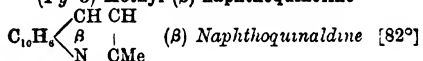
Dihydrate $C_{12}H_{11}O$ [113°] Obtained by saponification of its propionyl derivative, which is one of the products of the dry distillation of santonic acid Separated from accompanying di methyl naphthol by frequent crystallisation from light petroleum (Cannizzaro, *G* 13, 390) Glistening white needles, v sol ether and alcohol, insol water Volatile with steam P_2S_5 converts it into di methyl naphthalene

(Py 3) METHYL-(α) NAPHTHOQUINOLINE

$300^\circ)$ Heavy liquid Formed by heating (α) naphthylamine with paraldehyde and HCl Its salts have a blue fluorescence in dilute solution — $B'_2H_2Cl \cdot PtCl_2 \cdot 2aq$ concentric needles — $B'_2H_2Cr \cdot O$, yellow crystals (Doebner a Miller, *B* 17, 1711)

(Py 1) Methyl (β) naphthoquinoline $C_{11}H_{11}N$ 

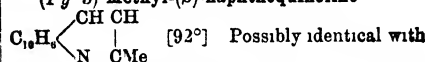
with (β) naphtho acridine and a base $C_{12}H_{13}N$, by the action of a mixture of methylal, acetone, and HCl upon (β) naphthylamine (Reed, *J pr* [2] 35, 316) —Picrate $B'_2C_6H_4(NO_2)_3OH$

(Py 3) Methyl-(β)-naphthoquinoline

(above 300°) Formed by heating (β) naphthylamine with paraldehyde and HCl (Doebner a Miller, *B* 17, 1711, Seitz, *B* 22, 254) Large colourless needles V sol alcohol and ether, sl sol water With chloral it forms a crystalline compound $C_{11}H_9N \cdot CH_2 \cdot CH(OH) \cdot CCl_2$, [185°]

Salts — $B'_2H_2Cl \cdot PtCl_2 \cdot 2aq$ yellow, sparingly soluble needles — $B'_2H_2Cr \cdot O$ small yellow needles, sl sol hot water — $B'HCl \cdot 2aq$ slender needles, sl sol cold water — $B'HNO_3$ slender needles, becoming rose coloured in air — $B'_2H_2SO_4 \cdot 2aq$ very slender needles, v e sol hot water — $B'_2C_6H_4(NO_2)_3OH$ [221°] Minute needles, v sl sol boiling water, v sol HOAc

Methyl iodide $B'MeI$ [241°–247°]. Straw coloured needles, v sol boiling water, sl sol alcohol

(Py 3) Methyl-(β)-naphthoquinoline

the preceding isomeride Formed by heating (Py 1, 3) oxy methyl naphthoquinoline to redness with zinc-dust (Knorr, *B* 17, 544) Crystalline May be distilled Its acid solutions fluoresce blue when dilute, green when concentrated — $B'_2H_2PtCl_2$ sl sol hot dilute $HClAq$.

(Py 1, 3) **Di-methyl-(α)-naphthoquinoline**

$$\text{C}_{10}\text{H}_7 \begin{array}{c} \text{CMe CH} \\ \diagup \quad \diagdown \\ \text{N}=\text{CMe} \quad \text{CH CH CMe CH} \\ \diagdown \quad \diagup \\ \text{C N}=\text{CMe} \end{array} \text{ v. e. } \text{C}_6\text{H}_5$$

 [44°] (361°) Formed by heating at 100° a mixture of (α)-naphthylamine with ethylidene-acetone, derived from acetone and paraldehyde (Reed, *J pr* [2] 35, 312) Formed also by heating (α)-naphthylamine (1 mol) with acetyl-acetone (1 mol), and heating the product with H_2SO_4 at 100° (Combes, *C R* 106, 1536) Needles (from petroleum-ether), v e sol ether, insol 90 p c alcohol. Somewhat volatile with steam. Its solution in H_2SO_4 is coloured purple by $\text{K}_2\text{Cr}_2\text{O}_7$. The solutions of its salts fluoresce violet (C) The hydrated platinochloride is violet, after drying *in vacuo* it melts at 260° (C) — $\text{B}'\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OH}$ [223°] Needles

(Py 1, 3)-Di methyl (β) naphthoquinoline

$$\text{C}_{10}\text{H}_7 \begin{array}{c} \text{CMe CH} \\ \diagup \quad \diagdown \\ \text{N}=\text{CMe} \quad \text{CH CH CMe CH} \\ \diagdown \quad \diagup \\ \text{C N}=\text{CMe} \end{array} \text{ v. e. } \text{C}_6\text{H}_5$$

$$\text{C}_6\text{H}_5 \begin{array}{c} \text{CMe CH} \\ \diagup \quad \diagdown \\ \text{N}=\text{CMe} \quad \text{CH CH CMe CH} \\ \diagdown \quad \diagup \\ \text{C N}=\text{CMe} \end{array} \text{ [127°] (above 300°)}$$

Formed from (β) naphthylamine hydrochloride by heating at 100° with ethylidene acetone, the product of the condensation of paraldehyde with acetone in presence of hydrochloric acid, as follows $\text{C}_{10}\text{H}_7\text{NH}_2 + \text{Me CO CH CHMe} = \text{C}_{10}\text{H}_7\text{N} + \text{H}_2\text{O} + \text{H}_2$ (Reed, *J pr* [2] 35, 299) Flat needles (from ether), v sl sol boiling water, hardly volatile with steam KMnO_4 oxidises it to (β) di methyl phenyl pyridine dicarboxylic acid. It does not yield a nitrosamine

Salts—Picrate $\text{B}'\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OH}$ [215°] — $\text{B}'\text{H}_2\text{Cr}_2\text{O}_7$ [c 115°] — $\text{B}'\text{H}_2\text{PtCl}_6 \cdot 2\text{aq}$ — $\text{B}'\text{H}_2\text{SO}_4$ — $\text{B}'\text{HNO}_3$ [181°] — $\text{B}'\text{HBr}$ [207°] — $\text{B}'\text{HBr} \cdot 2\text{aq}$

Methyl iodide $\text{B}'\text{MeI}$ needles

Sulphonic acid $\text{C}_{10}\text{H}_7(\text{SO}_3\text{H})\text{N} \cdot \frac{1}{2}\text{aq}$ minute needles, insol water and alcohol

Disulphonic acid $\text{C}_{10}\text{H}_7(\text{SO}_3\text{H})_2\text{N} \cdot 4\frac{1}{2}\text{aq}$ needles, v e sol water and alcohol — $\text{CuH}_2\text{A} \cdot 5\text{aq}$ — $\text{BaA}'' \cdot 7\text{aq}$ Potash fusion produces $\text{C}_{10}\text{H}_{11}(\text{OH})(\text{SO}_3\text{H})\text{N}$

Di-methyl (β) naphthoquinoline $\text{C}_{10}\text{H}_7\text{N}$ [67°] (380°) Obtained by heating (β) naphthylamine with acetyl acetone and H_2SO_4 at 100° (Combes, *C R* 106, 1537) Its yellow sulphonic acid $\text{C}_{10}\text{H}_7\text{NSO}_3$ is also pptd on adding ammonia to the aqueous extract of the product. The base forms a greenish yellow platinochloride, decomposing at 220° without melting

DI-METHYL-(β)-NAPHTHOQUINOLINEDI-

HYDRIDE $\text{C}_{10}\text{H}_7\text{N} \cdot \text{v. e. } \text{C}_6\text{H}_5$

$$\begin{array}{c} \text{CMe CMe} \\ \diagup \quad \diagdown \\ \text{NH CH} \quad \text{CH CH} \\ \diagdown \quad \diagup \\ \text{C N}=\text{CMe} \end{array} \text{ [115°]}$$

Formed by heating (*In* 1, 2) di-methyl (β) naphthindole with MeI in sealed tubes for 15 hours at 100° (Fischer a Steche, *B* 20, 820, *A* 242, 364) Plates, v sl sol water, v sol ether, alcohol, and mineral acids, volatile with steam. With nitrous acid it yields a crystalline nitrosamine — $\text{B}'\text{HI}$ Needles (from water), sl sol water and alcohol —The platinochloride is sl sol water and alcohol

METHYL NAPHTHYLAMINE v NAPHTHYLAMINE

Methyl di naphthyl-amine v. Di-naphthylamine.

DI-METHYL-NAPHTHYLENE DIAMINE v. NAPHTHYLENE DI-METHYL DI AMINE

METHYL-NARCEINE v NARCEINE

METHYL-NARINGENIC ACID v Methyl derivative of p COUMARIC ACID

METHYL-NICOTINE v NICOTINE

METHYL-NICOTINIC ACID v METHYL-

PYRIDINE CARBOXYLIC ACID

METHYL-NITRAMINE v METHYLAMINE

METHYL NITRATE $\text{CH}_3\text{NO}_2 \cdot \text{v. e. } \text{CH}_3\text{ONO}_2$
 Mol w 77 (65°) SG $\frac{1}{4}$ 1.2167, $\frac{3}{4}$ 1.2032
 M M 2 057 (Perkin, *C J* 55, 682) SV 69.3 (Lossen, *A* 254, 73) Obtained, together with methyl nitrite, by distilling wood spirit with NaNO_3 and H_2SO_4 (Dumas a Pélégot, *A Ch* [2] 58, 37) Prepared by distilling methyl alcohol (200 cc) with urea nitrate (40 g) and HNO_3 (150 cc of SG 1.31) free from nitrous acid. When two thirds have passed over, an additional quantity of MeOH (170 cc) and HNO_3 (110 cc) may be added and the distillation continued (Carey Lea, *Am S* [2] 33, 227) The process may also be conducted in a continuous manner (v LTHYL NITRATE)

Properties —Colourless liquid, exploding when struck or when its vapour is heated. With solid KOH it yields Me_2O (Berthelot, *A* 113, 80) When diluted with MeOH ($\frac{1}{2}$ vol) and treated with a current of gaseous NH_3 it gives methylamine nitrate and NMe_2NO , with only traces of di and tri methylamine. Aqueous ammonia forms NMeH_2 (13 pts), NMe_2NO (10 pts), and a small quantity (1 pt) of the bases NMe_2H and NMe_3 (Duvillier a Malbot, *A Ch* [6] 10, 284)

METHYL NITRITE $\text{CH}_3\text{NO}_2 \cdot \text{v. e. } \text{CH}_3\text{ONO}$
 Mol w 61 (−12°) SG (liquid) 1.991
 Produced by treating methyl alcohol with nitric acid and copper turnings or arsenious acid (Strecker, *C R* 39, 53, *A* 91, 82) The product is passed through a receiver at 0° and the gas after purification by passing through potash, a solution of ferrous sulphate, and dry CaCl_2 is condensed at −40°. It may also be prepared from methyl alcohol, NaNO_2 and H_2SO_4

Properties —Gas, smelling like nitrous ether. Burns with a green edged flame

METHYL NITRO-AMIDE v METHYLAMINE

METHYL-NITRO-ANILINE v NITRO

METHYL ANILINE

METHYL NITRO-BENZAMIDE v Amide of NITRO TOLUIC ACID

METHYL-NITROLIC ACID $\text{CHN O}_2 \cdot \text{v. e. } \text{CH}(\text{NO}_2)\text{NOH}$ or $\text{CH}_2(\text{NO}_2)(\text{NO})$ Mol w 90 [64°] Prepared by dissolving nitro methane (15 g) in water, adding a solution of KNO_2 (8 g), cooling to 0°, and adding a very dilute ice cold solution of H_2SO_4 (4 g) Caustic potash solution is added till the liquid turns red, and then more dilute H_2SO_4 . The liquid is then shaken with a little CaCO_3 and extracted with ether (Tscherniak, *B* 8, 114, *A* 180, 166) The result is very uncertain (V Meyer a Constam, *A* 214, 335)

Properties —Long needles (from ether). Decomposes slowly in the cold, rapidly at 64°, into formic acid, NO_2 , and nitrogen. By boiling dilute H_2SO_4 it is resolved into formic acid and N_2O . Sodium amalgam reduces it to methyl-azauric acid $\text{CH}(\text{NOH})\text{N N CH NOH}$ an amorphous powder which is violently gasified above 100°.

DI-METHYL-NITROSAMINE *v* **DI-METHYL-AMINE****METHYL NITROSO-ETHYL KETONE** *v*.*Mono oxim* of **DI-METHYL DIKETONE****TETRA METHYL-NITROSO-PHENYLENE-DIAMINE** *v* **NITROSO-PHENYLENE TETRA METHYL DIAMINE****METHYL NONYL KETONE** *v* **METHYL ENNYL KETONE****METHYL-NOROPIANIC ACID** *v* **OPIANIC ACID**

METHYL *n* OCTYL KETONE $C_{10}H_{20}O$ *z* $CH_3 \cdot CO \cdot CH_2 \cdot CH_2 \cdot CH_2 \cdot CH_2 \cdot CH_2 \cdot CH_2 \cdot CH_2 \cdot CH_3$ [4°] (214°) (J), (211°) (K) SG $\frac{17.6}{17.5}$ 8294, $\frac{1}{4}$ 838, $\frac{2}{3}$ 825 (K) Formed by boiling heptyl-aceto acetic ether with dilute alkalis (Jourdain, A 200, 106) Formed also by distilling a mixture of barium acetate and barium ennoate (Krafft, B 15, 1695) Liquid, with pleasant odour, insol water, solidifying in a freezing mixture Forms a crystalline compound with $NaHSO_4$.

Methyl octyl ketone $C_{10}H_{20}O$ *z*

$CH_3 \cdot CO \cdot CH_2 \cdot CHMe \cdot C_5H_{11}$ *Heptyl acetone* (197°) Formed by the action of baryta water on *n* sec heptyl aceto acetic ether (Venable, B 13, 1651) Colourless liquid, lighter than water

METHYL OCTYL OXIDE $C_8H_{16}O_2$ *z*

$CH_3 \cdot O \cdot C_7H_{15}$, (173°) SG $\frac{8}{8}$ 8014 SV 219 8 CE (0°-10°) 00101 (Dobner, A 243, 3)

METHYL OCTYL PHENYL AMINE *v* **OCTYL TOLYL AMINE****METHYL-OCTYL-THIOPHENE** $C_{13}H_{10}S$ *z*

$S < \begin{matrix} CMe \\ C(C_7H_5) \end{matrix} = CH >$ [10°] (272°) Formed by the action of sodium upon an ethereal solution either of MeI and iodo octyl thiophene or of octyl biomide and iodo methyl thiophene (Schweinitz, B 19, 648) Bromine forms $C_{12}H_9BrS$ [20°]

METHYL OXALATE *v* **Methyl ether of OXALIC ACID**

METHYL-OXALACETIC ETHER $C_5H_8O_4$ *z* $CO \cdot Et \cdot CO \cdot CHMe \cdot CO \cdot Et$ *Oxaloxy propionic ether* (138° at 23 mm) Formed by the action of $NaOEt$ on an ethereal solution of oxalic and propionic ethers (Wilschusen & Arnold, B 20 3394, A 246, 329) Formed also from sodium oxalacetic ether and MeI at 100° Colourless oil, *v* sol alcohol and ether Its alcoholic solution gives a red colouration with $FeCl_3$ Split up by boiling alcoholic potash into oxalic and propionic acids Boiling dilute H_2SO_4 yields ethyl glyoxylic acid

Salt $-CO_2Et \cdot CO \cdot CNaMe \cdot CO_2Et$ Does not crystallise from alcohol

Phenyl hydrazide

$CO \cdot Et \cdot C(N_2HPh) \cdot CHMe \cdot CO \cdot Et$ [100°] Small plates, *v* sol ether and benzene Its solution in conc H_2SO_4 is coloured reddish violet by $FeCl_3$ At 120° it gives off alcohol, forming a derivative of pyrazole

METHYL-OXALYL UREA *v* **PARABANIC ACID****METHYL-OXAMIC ACID** *v* **OXALIC ACID****METHYL-OXAMIDE** *v* **OXALIC ACID****METHYL-OXANTHRANOL** *v* **OXANTHRANOL****METHYL-OXAZOLE DIHYDRIDE** C_4H_6NO

$CH_2 \cdot O > CMe$. Formed, in small quantity, *i.e.* $CH_2 \cdot N >$

by the action of Ac_2O and $NaOAc$ on bromoethylamine $Br \cdot CH_2 \cdot CH \cdot NH_2$ (Gabriel, B 22, 2221, 23, 2502) Oil, with sweet smell, somewhat like quinaldine $-B \cdot C_6H_4 \cdot (NO_2)_2 \cdot OH$ [149°]. Yellow plates.

METHYL-OXETHYL- *v* **METHYL-OXYETHYL-DI-METHYL-OXETONE** *v* **Anhydride of DI- γ -OXY-DI-BUTYL DIKETONE**

DI-METHYL OXIDE C_2H_6O *z* $(CH_3)_2O$ *Methyl ether* Mol w 46 (-24°) (Regnault, J 1863, 70) VD 1 617 S (gas) 37 at 18° HF p 49,640 (Thomsen), 56,800 (Berthelot, A Ch [5] 23, 185) HF v 48,190 (T) HC p 344,200 (B) Formed by heating $MeOH$ with H_2SO_4 (Dumas & Péligot, A 15, 12, Kane, A 19, 166) or with B_2O_3 (Ebelmen, A 57, 328) Formed also, together with NMe_4Cl and NMe_3HCl , by heating NH_4Cl with excess of $MeOH$ (Weith, B 8, 458) Prepared by heating $MeOH$ (13 pts) with H_2SO_4 (20 pts) at 140° The gas is passed through aqueous KOH to remove SO_2 , CO_2 , and $MeOH$, and then into conc H_2SO_4 , which absorbs 600 vols It is obtained by dropping the solution in H_2SO_4 into an equal volume of warm boiled water, and is dried by passing through tubes containing $CaCl_2$ (Erlenmeyer & Kriechbaumer, B 7, 699, cf Teller, Ar Ph [3] 10, 57)

Properties — Gas Combines with HCl forming $MeOHCl$ which boils at 2°

References — **CHLORO** and **DI IODO DI METHYL OXIDE**

METHYL - OXINDOLE C_8H_7NO *z*

$C_6H_5 < \begin{matrix} CH \\ NMe \end{matrix} > CO$ *Oxy methyl indole* [88°]

Obtained from methyl indole carboxylic acid by the action of $NaOBr$, the resulting di bromo methyl oxindole being suspended in alcohol and reduced by sodium amalgam (Colman, C J 55, 7, A 248, 120) White needles, sl sol cold water and light petroleum, *v* sol alcohol, ether, acetone and benzene Dissolves in hot alkalis without change Partially decomposes when heated much above its melting point Does not react with phenyl hydrazine Bromine water gives a crystalline pp Nitrous acid converts it into $C_6H_5 < \begin{matrix} C(OH) \\ NMe \end{matrix} > CO$, the oxim of methyl ψ isatin

Di-chloro methyl oxindole $C_8H_5 < \begin{matrix} CCl \\ NMe \end{matrix} > CO$.

[147°] Formed by adding a solution of sodium methyl indole carboxylate to a cold solution of $NaOCl$ (Colman) Colourless needles, *v* sol hot alcohol and acetone, *m* sol ether Not decomposed at 210°

Bromo methyl oxindole $C_8H_5 < \begin{matrix} CHBr \\ NMe \end{matrix} > CO$

[134°] Formed, together with methyl oxindole, by reducing di bromo methyl oxindole with sodium amalgam Lustrous white plates, *v* sl sol cold water, *v* sol hot alcohol Not decomposed by boiling aqueous KOH

Di-bromo methyl oxindole $C_8H_5 < \begin{matrix} CBr_2 \\ NMe \end{matrix} > CO$.

[204°] Formed by the action of $NaOBr$ on methyl indole carboxylic acid (Fischer, B 17, 564) Yellowish white tables, *v* sol alcohol, insol cold water Melts at 204° when quickly heated, but 180° when slowly heated Converted by boiling water into methyl ψ isatin Phenyl-

hydrazine gives the phenyl hydrazide of methyl- ψ -isatin

Di- methyl - oxindole $C_{10}H_7NO$ ν e
 $[5 \ 3 \ 1] C_6H_5(CH_2)_2 \begin{smallmatrix} CH \\ \diagup \quad \diagdown \\ NH \end{smallmatrix} > CO$ *Carbomesyl Anhydride of amido di methyl phenyl acetic acid* [282°] Prepared by reduction of (2 5 3 1) nitro di-methyl-phenyl acetic acid with tin and HCl (Wispeh, B 16, 1580) Sublimable White needles Sol hot alcohol and hot benzene sl sol hot water and cold alcohol and ether, insol cold water

Methyl di-oxindole $C_8H_7 \begin{smallmatrix} CH(OH) \\ \diagup \quad \diagdown \\ NMe \end{smallmatrix} > CO$

Di-oxy methyl indole [151°] Formed by reducing methyl ψ isatin with sodium amalgam or with zinc and HCl (Colman, C J 55, 8, A 248, 121) Needles or prisms (from benzene), m sol water, alcohol, and benzene Oxidising agents reconvert it into methyl- ψ isatin

METHYL-OXY-BENZOYL-GLYCOCOLL ν
ANISURIC ACID

METHYL OXYBUTYLENE KETONE DI-CARBOXYLIC ACID ANHYDRIDE $C_8H_6O_4$, ν c
 $CH_3COCH \begin{smallmatrix} C(CO_2H) \\ \diagup \quad \diagdown \\ CO \end{smallmatrix} > CMe$ *Isocarboxypyrrotaric acid* Obtained from its ether by boiling with 20 p c aqueous NaOH, and ppg with dilute H_2SO_4 (Knorr, B 22, 163) Possesses great reducing power Decomposes at 200°-209°, leaving an oil $C_8H_6O_4$, which solidifies on cooling This is composed of two bodies, one being an acid v sl sol most solvents and melting at 175°, the other crystallising from ether in long prisms (60°), and giving an acid solution in water Water forms acetyl acetone

Ethyl ether EtA' [110°] (c 280° at 15 mm) Obtained by heating di acetyl succinic ether at 170°-180°, pyrotritaric and carboxypyrrotaric ethers being also formed (Knorr, B 22, 159) Slender needles (from hot water), v sl sol water and dilute acids, v sol alkalis, ether, chloroform, and hot alcohol Possesses great reducing power $FeCl_3$ gives a fine blue colour to its solution With phenyl-hydrazine it yields di oxy-di phenyl di methyl dipyrazyl N Ph CO CO N Ph N

$\begin{array}{c} \parallel \quad \parallel \quad \parallel \\ C(CH_3) \quad CH-CH- \quad CMe \\ \parallel \quad \parallel \quad \parallel \\ N \quad O \quad CO \quad CO \quad O \quad N \end{array}$ **Hydroxylamine ap**
 pairs to form the corresponding di oxy di methyl-dioxazyl $\begin{array}{c} \parallel \quad \parallel \quad \parallel \\ CMe \quad CH \quad CH- \quad CMe \end{array}$

DI-METHYL-DI-OXY-BUTYLENE DIKETONE $CH_3CO CMe(OH) CMe(OH) CO OH$, [96°] Formed by reducing di methyl diketone with zinc dust and dilute H_2SO_4 (Von Pechmann, B 21, 1411) Needles (from ligroin) Reduces Fehling's solution in the cold $FeCl_3$ re converts it into Me CO CO Me

METHYL OXY-BUTYL KETONE $C_8H_{12}O_4$, ν e
 $CH_3COCH_2CH_2CH_2CH_2OH$ *Acetyl - butyl-alcohol* (227°) SG $\frac{1}{4}$ 997, $\frac{1}{2}$ 989, $\frac{3}{4}$ 982 M M 6 502 at 18.5° Obtained by boiling the anhydride of its carboxylic acid with water, CO_2 being evolved (Perkin, jun, C J 51, 717) Thick oil with sweet burning taste Does not reduce Fehling's solution or ammoniacal $AgNO_3$ in the cold, but reduces both these reagents on warming Yields an oily phenyl hydrazide Warm conc HNO_3 gives a dark pink colour

Iodine and KOH give iodoform Chromic acid mixture oxidises it to acetic and succinic acids

METHYL OXY BUTYL KETONE ANHYDRIDE $C_8H_{10}O_4$, ν e $O \begin{smallmatrix} CH_2CH \\ \diagup \quad \diagdown \\ CMeCH \end{smallmatrix} > CH_2$ *Methyl-furandane dihydride 'Acetyl tetramethylene' 'Tetramethylene methyl ketone' Anhydride of aceto butyl alcohol* (109°) SG $\frac{1}{4}$ 9227, $\frac{1}{2}$ 9127, $\frac{3}{4}$ 9050 M M 6 074 at 22.5° Formed by distilling its carboxylic acid at 150° (Perkin, jun, C J 51, 723, B 19, 2558) Colourless mobile liquid with ethereal odour On prolonged standing in presence of water it is converted into methyl oxy butyl ketone

Methyl oxy butyl ketone anhydride carboxylic acid $C_8H_{10}O_5$, ν e $O \begin{smallmatrix} CH_2CH \\ \diagup \quad \diagdown \\ CMeC(CO_2H) \end{smallmatrix} > [119°]$

Obtained by saponifying its ether with alcoholic potash (Perkin, jun, C J 51, 715) Colourless needles, v sol hot benzene, alcohol, chloroform, and light petroleum, v e sol water Decomposes above 119° into CO and the preceding body When heated with aqueous NH_3 at 200° it yields CO_2 , methyl oxy butyl ketone, and a volatile base (probably a derivative of pyridine) Bromine vapour gives $C_8H_7BrO_4$, a thick syrup Conc HBr aq yields methyl bromo butyl ketone

Salts— NH_4A' white crystalline solid, v sol water— CuA' aq light green powder, v sl sol water, decomposed at 90°-100°— AgA' white crystalline solid, v sol water, decomposed on boiling with water or exposure to light

Ethyl ether EtA' [9°] (227°) V D 6 21 M M 10 069 at 23.7° SG $\frac{1}{4}$ 1 069, $\frac{1}{2}$ 1 0626, μ 1.4697 μ_D 1.4772 Formed by heating a mixture of NaOEt with aceto acetic ether and trimethylene bromide (Perkin, jun, C J 51, 709) Colourless oil with a disagreeable camphor like odour Does not react with phenyl hydrazine Does not contain an atom of hydrogen displaceable by sodium With PCl_5 it forms $C_8H_{10}O_2Cl_2$, a colourless oil (212°-215°)

Methyl oxy butyl ketone anhydride dicarboxylic acid $C_8H_{10}O_6$, ν e

$O \begin{smallmatrix} CH_2CHCH_2 \\ \diagup \quad \diagdown \\ C(CH_2CO_2H)C(CO_2H) \end{smallmatrix} > [185°-190°] Ob$

tained by saponification of its ethers by boiling alcoholic potash (Perkin, jun, C J 51, 744) Colourless hexagonal plates, v sol hot water and hot alcohol, sl sol acetic ether, benzene, and ether Decomposes on fusion, giving off CO_2 , and leaving a red liquid

Mono ethyl ether
 $O \begin{smallmatrix} CH_2CH_2CH_2 \\ \diagup \quad \diagdown \\ C(CH_2CO_2H)C(CO_2Et) \end{smallmatrix} > [114°] Obtained$

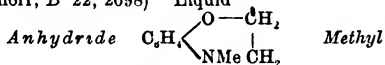
by treating the di ethyl ether with conc alcoholic potash in the cold Transparent four sided tri clinic needles, $a b c = 774 \ 1 \ 337$, $\alpha = 89^\circ 40'$, $\beta = 98^\circ 18'$, $\gamma = 89^\circ 50'$ V sol alcohol and ether, sl sol hot water— $AgEtA'$ white curdy pp, sl sol water

Di ethyl ether Et₂A' (239° at 150 mm) Obtained by the action of NaOEt on a mixture of trimethylene bromide and acetone di-carboxylic ether $CO_2EtCH_2COCH_2CO_2Et$ (Perkin, jun, C J 51, 739) Thick, colourless oil, with disagreeable odour

METHYL-OXY-GONINE & CONTINUED.

METHYL-OXY-ETHYL-AMIDO-PHENOL

Methyl ether $C_{10}H_{11}NO$, *ie* $[2] C_6H_4(OMe) NMe CH_2OH$ *Methyl-oxy ethyl anisidine* (290°) Formed by the action of $C_6H_4(OMe) NHMe$ on ethylene chlorhydrin (Knorr, *B* 22, 2098) Liquid



quinazoline dihydride (261°) Formed by boiling the methyl ether with caustic soda (Knorr, *B* 22, 2098) Liquid, with irritating smell, volatile with steam, very easily oxidised by contact with air $B'HCl$ [162°] Short four cornered plates (from alcohol)

METHYL OXY ETHYL AMINE C_3H_7NO *ie* $CH_3NHCH_2CH_2OH$ (130°-140°) Formed from ethylene chlorhydrin (1 mol) and methyl amine (1 mol) (Knorr, *B* 22, 2088) Oil, with strong ammoniacal smell, ν sol water, alcohol, and ether *Aurochloride* [110°-120°] Anhydrous prisms, ν sol water

Methyl di oxy di ethyl amine C_3H_7NO *ie* $CH_3N(CH_2CH_2OH)_2$ (250°-255°) Formed by heating ethylene chlorhydrin with excess of aqueous methylamine at 100° (Morley, *B* 13, 222) Formed also by the action of ethylene chlorhydrin on the preceding body (Knorr, *B* 22, 2081) Viscid oil, ν sol water, not volatile with steam— $B'HCl$ thick syrup— $B'_2H_2PtCl_6$ orange red prisms—*Aurochloride* ν sol water

Methyle chloride C_6H_5NOCl *ie* $CINMe(CH_2CH_2OH)_2$ Formed by heating ethylene chlorhydrin with a solution of dimethyl amine at 100° (Morley *B* 13, 223) Formed also from di methyl oxy ethyl amine and ethylene chlorhydrin (Knorr *B* 22, 2089) Thick syrup— $(C_6H_5NOCl) PtCl_6$ aq [218°] Small yellow crystals, ν sol water, sl sol alcohol—*Aurochloride* [233°], small plates, ν sol hot water

Anhydride $CH_3N \begin{array}{c} \diagup CH_2 \\ \diagdown CH_2 \end{array} O$ *Methyl pyrazine tetrahydride* (117°) Formed by heating methyl di oxy di ethyl amine with SO_2 (h) Mixes with water, alcohol and ether Readily combines with MeI —*Hydrochloride* [205°], long hygroscopic prisms—*Platino chloride* needles (from dilute alcohol), ν sol water—*Aurochloride* [143°], small needles m sol water

Methyle iodide of the anhydride Long needles With AgO it yields a methyle hydroxide, which is split up by heat into aldehyde and di methyl oxy ethyl amine

Methyle chloride of the anhydride Crystalline mass Forms a platinochloride crystallising from alcohol in needles, and an aurochloride, both being ν sol water

Di methyl oxy ethyl amine C_3H_7NO *ie* NMe, CH_3, CH_2OH *Di methyl ethyl alkane* (150°) Prepared from dimethylamine and ethylene chlorhydrin (Ladenburg, *B* 14, 2408) Formed also by heating 'methyl morphine' (*v* MORPHINE) with Ac_2O at 160°-190° (Knorr, *B* 22, 1114) Liquid— $B'_2H_2AuCl_6$ needles, sol hot water— $B'_2H_2PtCl_6$ easily soluble prisms

Acetyl derivative C_3H_7AcNO Forms an aurochloride $B'HauCl_6$ crystallising in plates
Vol III

Methyle hydroxide v NEURINE

TRI-METHYL-OXYETHYL-AMMONIUM

HYDROXIDE v NEURINE

METHYL OXYETHYL ANILINE

C_6H_5NO *ie* $C_6H_5N(CH_2)(C_6H_5OH)$ *Methyl phenyl ethyl alkane* (218° at 110 mm) $SG \frac{2}{2}$ 1 0806 Obtained by heating methyl aniline with ethylene chlorhydrin at 100° (Laun, *B* 17, 675) Colourless oil Weak base By exposure to air it is oxidised to a thick blue syrup

Methyle iodide $B'MeI$ colourless plates or tables

Methyle-periodide $B'MeI$, [87°], greenish glistening plates

METHYL OXY ETHYL KETONE $C_3H_7O_2$

ie $CH_3COCH(OH)CH_3$ (142°) *Ketoxyl butane* Formed by reducing di methyl diketone with zinc and H_2SO_4 (Von Pechmann, *B* 22, 2214, 23, 2421) Colourless liquid, miscible with water Reduces Fehling's solution Reacts with phenyl hydrazine, with formation of the compounds $CH_3C(NHPh)CH(OH)CH_3$ [84°] and $CH_3C(NH_2)CH(NHPh)CH_3$

METHYL OXY ETHYL PYRIDINE *ie* THORIN**METHYL OXY ETHYL p TOLUIDINE**

$C_{10}H_{13}NO$ *ie* $CH_3OHCH_2NMeC_6H_4Me$ (290°-300°) Formed from oxy ethyl toluidine and MeI at 60° (Demole, *A* 173, 129) Liquid— $B'HI$ — $B'HI$ — $B'HI$

Methyle iodide $B'MeI$ Liquid Yields $B'_2Me.PtCl_6$ and $B'MeAuCl_6$

METHYL OXY GLUTARIC ACID v OXY

METHYL CLUTARIC ACID

TRI METHYL TRI OXY HYDROBENZ

AMIDE *ie* ANISYDRAMIDE

TETRA METHYL OXY TRIMETHYLENE-

DIAMINE $C_4H_{13}N_2O$ *ie* $HOCH(CH_2NMe)_2$ *Tetra methyl oxy propylene diamine* *Tetra methyl allyl alkane* (170°-185°) Formed by heating ν dichlorhydrin $CH_2ClCH(OH)CH_2Cl$ with dimethylamine in a sealed tube at 60° (Berend, *B* 17, 510) The liquid is rendered alkaline by KOH and extracted with chloroform Liquid, ν e sol water— $B'_2H_2PtCl_6$ yellow plates, m sol water

DI METHYL TRI OXY PHENYL BENZYL-KETONE *ie* ANISON**DI METHYL OXY PROPYL AMINE**

C_4H_9NO (126°) Prepared by the action of dimethylamine on propylene chlorhydrin at 100° (Morley, *C* R 91, 333, Ladenburg, *B* 14, 2407) Liquid— $B'_2H_2PtCl_6$ prisms ν sol water

Methyle chloride $B'MeCl$ Prepared by heating aqueous trimethylamine with propylene chlorhydrin at 100° (Morley, *C* R 91, 333, *B* 13, 1805) Colourless, very hygroscopic crystals, turning brown in light Form a platinochloride $B'_2MePtCl_6$ crystallising in yellow feathery plates, m sol alcohol and ether, ν sol water

Methyle hydroxide $B'MeOH$ From the chloride and moist AgO Alkaline syrup Decomposed on distillation into NMe_3 and propylene glycol (Morley, *C* J 41, 889)

Di-methyl dioxypropyl amine $C_3H_7NO_2$ *ie* $NMe, CH_2, CH(OH)CH_2(OH)$ *Di methyl propyl-glycolline* (217°) Formed by heating di methyl amine with glycerin chlorhydrin (Roth, *B* 15, 1153) Colourless oil Sol water, alcohol, and ether— $B'_2H_2Cl_6PtCl_6$

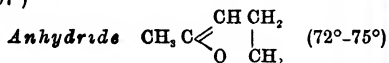
Methylo chloride $C_2H_5NO_2Cl$ \approx $NMe_2ClCH_2CH(OH)CH_2(OH)$ From glycerin chlorhydrin and NMe_2 at 100° (V. Meyer, *B* 2, 186, Hanriot, *A Ch* [5] 17, 99). Crystals, v e sol water — $(C_2H_5NO_2Cl)_2PtCl_6$ tables — $C_2H_5NO_2AnCl$, [190°] Orange crystals

METHYL OXYPROPYL-ANILINE $C_{10}H_{11}NO$ \approx $C_6H_5N(CH_3)C_2H_4OH$ **Methyl phenyl propyl-alkine** (262°) Colourless liquid Formed by heating methyl aniline with propylene chlorhydrin (Laun, *B* 17, 678)

METHYL β OXYPROPYL KETONE $C_5H_8O_2$ \approx $CH_3COCH_2CH_2CH_2OH$ **Acetyl propyl alcohol** (145°) at 100 mm, (209°) at 729 mm V D 2.25 (calc 3.58) SG $\frac{4}{5}$ 1.0051, $\frac{25}{4}$ 9990 M M 5.544 at 26° Formed by boiling trimethylene methyl ketone carboxylic acid (derived from aceto acetic ether, sodium, and ethylene bromide) with water, CO_2 being evolved (Perkin, jun, *C J* 51, 829) Formed also by boiling bromo ethyl aceto acetic ether with aqueous HCl Thick syrup, miscible with water, v sol alcohol and ether Decomposed by heat into water and its anhydride, which reunite in the cold, does not reduce cold ammoniacal silver nitrate, but reduces it on warming Gives with phenyl hydrazine solution an oily precipitate of $C_{11}H_{14}N_2$ or $CH_3CH(CMe)CH_2NC_6H_5$ Oxidised by $K_2Cr_2O_7$ and H_2SO_4 to acetyl propionic (levulic) acid (Colman a Perkin, *C J* 53, 189, 55, 352, Lipp, *B* 22, 1196) Reduced by sodium amalgam to $CH_3CH(OH)CH_2CH_2CH_2OH$ (Perkin, jun, a Freer, *B* 19, 2566) With $NaHSO_3$ it forms $CH_3C(C_2H_4OH)(OH)SO_3Na$ $\frac{1}{2}$ aq needles, v sol water and alcohol

Acetyl derivative $CH_3CO C_2H_4OAc$ (214° v) at 728 mm SG $\frac{8}{5}$ 1.0356 V D 5.02 (calc 4.98) Liquid, m sol water, v e sol alcohol and ether

Benzoyl derivative $CH_3CO C_2H_4OBz$ (297°)



Formed by slow distillation of the ketone Water converts it into the flocculent isomeride

$CH_3CH_2CH_2C \begin{array}{c} \swarrow O \\ \searrow \quad | \\ \quad CH_2 \end{array} \quad (111^\circ)$ at 718 mm, which reddens pine wood moistened by HCl v **METHYL ENEFURFURANE TRIHYDRIDE**

Methyl α -oxy-propyl ketone $C_5H_{10}O_2$ \approx $CH_3COCH(OH)CH_2CH_3$ (158°), (77° at 35 mm) SG $\frac{17.5}{4}$ 972 Formed by reducing methyl ethyl diketone with zinc and dilute H_2SO_4 (Von Pechmann, *B* 23, 2421) Colourless liquid with sweet smell, sol water, but separated from the solution by NaCl or NaOH Readily oxidised to $CH_3COCO C_2H_5$ Reduced by sodium amalgam to $CH_3CH(OH)CH(OH)C_2H_5$ (187°), which is oxidised by bromine water in sunlight to $CH_3COCO C_2H_5$ Excess of phenyl hydrazine forms $CH_3C(N_2HPh)CO C_2H_5$ [103°]

Methyl α -oxy-propyl ketone $C_5H_{10}O_2$ \approx $CH_3COCH_2CH(OH)CH_3$ (128°) Formed by adding chloro acetic ether to sodium under ether, decomposing the resulting sodium compound by HCl, reducing by zinc dust the ether $C_5H_{11}ClO$, (157° at 45 mm) which is then

formed, and boiling the product $C_5H_{11}O$, (106° at 14 mm), with dilute HCl, when it splits up into CO_2 , alcohol, and the oxy ketone (Fittig, *B* 21, 2188) Colourless mobile liquid, miscible with water Forms compounds with $NaHSO_3$ and with phenyl hydrazine

Methyl α oxy isopropyl ketone **Oxim of the nitrate** $(CH_3)_2C(ONO_2)C(OH)CH_3$ **Pentane nitroso nitrate** Formed by the action of amyl nitrite and nitric acid on amylene $(CH_3)_2CCH_2CH_3$ (38°) dissolved in $HOAc$ (Wallach, *A* 248, 162) Monoclinic crystals, $abc = 977.1 \times 1449$, $\beta = 83^\circ 32'$ (from benzene), or needles (from $HOAc$)

METHYL OXY QUINIZINE v **OXY PHENYL-METHYL-PYRAZOLE**

METHYL OXY QUINOLINE v **OXY METHYL-QUINOLINE**

METHYL OXY SUCCINIC ACID v **OXY-METHYL SUCCINIC ACID**

METHYL PARABANIC ACID v **PARABANIC ACID**

METHYL PARACONIC ACID v **Anhydride of OXY METHYL SUCCINIC ACID**

TETRA METHYL-PARALEUCANILINE v **TETRA METHYL TRI AMIDO TRI PHENYL METHANE**

METHYL PENTADECYL KETONE $C_{15}H_{32}O$ \approx $CH_3CO C_{13}H_{27}$, [48°] (246°) at 110 mm, (320° uncor) at 760 mm SG (liquid) $\frac{25}{4}$ 814 Prepared by distilling a mixture of barium palmitate and barium acetate (Krafft, *B* 12, 1671) On oxidation it gives pentadecic acid

Methyl pentaecyl ketone $CH_3COCH(C_2H_5)_2$ **Di n heptyl acetone** (300°-304°) SG $\frac{47}{5}$ 826 Formed by boiling di heptyl acetoacetic ether with dilute aqueous KOH (Jourdain, *A* 200, 115) Oil, smelling of peppermint Forms a crystalline compound with $NaHSO_3$

METHYL PENTANE v **HEXANE**

DI-METHYL PENTENYLAMINE $C_8H_{17}N$ \approx $CH_3CHCH_2CHCH_2NMe$ This constitution is assigned by Merling to Ladenburg's di methyl piperidine, v **DI-METHYL PYRIDINE HEXAHYDRIDE**

Di methyl pentenylamine **Methylo hydroxide** $C_8H_{17}NMe_2OH$ Formed from amylene bromide and trimethylamine at 55° (Schmiedeberg a Harnack, *J* 1867, 805) — $(C_8H_{17}NMe_2Cl)_2PtCl_6$ aq Irregular laminae

(8) **METHYL PENTHIOPHENE** C_8H_8S \approx $S < \begin{array}{c} CHCMe \\ CHCH \end{array} > CH_3$ (134°) SG $\frac{18}{5}$ 9938

Formed by distilling sodium α methyl glutarate $CO_2NaCHMeCHCH_2CO_2Na$ (5 g) with P_2S_5 (10 g) at 180°-250° (Kreker, *B* 19, 3270) Colourless oil When successively mixed with a solution of isatin in $HOAc$ and conc H_2SO_4 it gives a dark green colour, and on adding water, a green pp Phenanthraquinone and H_2SO_4 (Laubenheimer's reagent) give a dark violet colour Phenyl glyoxylic acid yields a violet dye Alkaline $KMnO_4$ (3 pc) produces aceto and oxalic acid Nitric acid forms a nitro-derivative, and bromine a biomo-derivative

Methyl penthienyl methyl ketone $C_8H_{11}SO$ \approx $SC_2H_4MeCOCH_3$ (284°)

Oxim $SC_2H_4MeC(OH)CH_3$ [68°]

METHYL-PHENACYL-ANILIDE a **PHENYL-AMIDO PHENYL ETHYL KETONE**

METHYL - PHENACYL - BROMIDE

a Bromo phenyl ethyl ketone

METHYL-PHENANTHRIDINE ν **PHENYL-INDOLE****METHYL-PHENANTHROLINE** $C_{11}H_{10}N_2$ ν ϵ
CH N C CH CH C CH CH

CH CH C ——— C N CMe $^{\nu}$ [65°] (above 360°) Formed, together with an isomeride, when *m* amido (Py 3) methyl quinoline (100 g) is boiled with *o* nitro phenol (75 g), glycerin (320 g), and H_2SO_4 (270 g). The product is diluted with water and evaporated to remove *o* nitrophenol. The resinous mass that is ppd on adding NaOH is extracted with benzene, and the dark oily mixture of bases so obtained treated with HCl. The hydrochlorides are washed with alcohol, dissolved in water, decomposed by NH_3 , and the bases again extracted with benzene. From the crystalline mixture of bases left after evaporating the benzene, ether extracts methyl-phenanthroline only (Gerdeissen, B 22, 246).

Properties — White needles (containing 3 aq) Melts at 50° when hydrated, 65° when anhydrous. Sol boiling water, alcohol, and ether, ν sol cold benzene. On oxidation with $KMnO_4$ it yields phenanthroline carboxylic acid $C_{12}H_9N_2COH$ [209°], which decomposes at 210° into CO_2 and phenanthroline.

Salt — $B'HCl$ aq small snow white needles, sl sol cold, ν sol hot, water, *m* sol alcohol, insol ether — $B'H SO_4$ aq slender silky needles (from alcohol) — $B'H_2CrO_4$ red prisms (from hot water) — $B'H_2PtCl_6$ aq flesh coloured crystalline pp, insol hot water, sl sol alcohol — Picrate $B'C_6H_4(NO)_2OH$ [217°] Transparent yellow needles (from boiling alcohol).

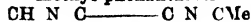
Ethyl iodide $B'EtI$ 2aq [100°–110°] Glistening brown crystals. Yields the platinum chloride $B'EtClH.PtCl_6$ (?) which forms orange red crystals.

Isomeride $C_{11}H_{10}N_2$ ν ϵ 

CH CH C CH C CH CH CH [109°] Occurs in the

preparation of the preceding, from which it differs in being insol ether (G). Crystallises from alcohol in concentric groups of satiny needles (containing 1 aq). Melts at 82° when hydrated and 109° when anhydrous. The anhydrous base is a light white powder, ν sol warm benzene and hot dilute alcohol, insol cold ether, ν sl sol hot ether.

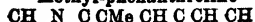
Methyl phenanthroline



CH CH C CH C CH CH CH [76°] Prepared

from *o* amido (Py 3) methyl quinoline, glycerin, *o* nitro phenol, and H_2SO_4 (Gerdeissen). Crystals (containing 2 aq). Melts at 58° when hydrated, and at 76° when anhydrous. ν e sol hot benzene, ν sol chloroform, *m* sol HOAc, sl sol ether and light petroleum. Decomposed on distillation.

Methyl-phenanthroline

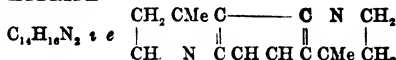


CH CH C ——— C N CH [96°] (above

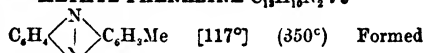
300°) Prepared by boiling *m*-tolylene diamine hydrochloride (40 g) with nitrobenzene (30 g), glycerin (100 g), and H_2SO_4 (100 g), and puri-

fied by means of the chromate (Skraup & Fischer, M 5, 523). Short prisms (containing 5 aq), more soluble in cold than in hot water, ν sol alcohol. Its solutions give a white crystalline pp with $AgNO_3$ and a blue crystalline pp with cupric acetate. Chromic acid mixture oxidises it to the corresponding phenanthroline carboxylic acid.

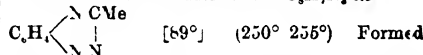
Salts — $B'HCl$ 4aq transparent needles — $B'H_2CrO_4$ red, sparingly soluble needles — $B'H_2PtCl_6$ 2aq yellow crystalline pp — Picrate [253°]

METHYL-PHENANTHROLINE TETRA-HYDRIDE

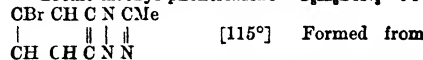
Formed by the action of acetic aldehyde or paraldehyde upon *m* phenylene diamine hydrochloride. The resulting base is ppd by ammonia, dissolved in aqueous HCl, and ppd by platinum chloride as $B'H_2PtCl_6$ (Schiff, A 253, 323).

METHYL PHENAZINE $C_{11}H_{10}N_2$ ν ϵ 

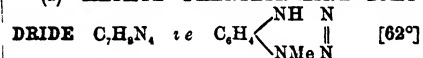
by heating tolylene *o* diamine with pyrocatechin in sealed tubes at 210° (Verz, B 19, 725). Formed also by eliminating the two NH_2 groups by means of the diazo reaction, from the oxidation product ($C_{12}H_{11}N_4$) of phenylene *p* diamine and tolylene *m* diamine (Berntsen & Schweitzer, B 19, 2604, A 236, 345). Needles (by sublimation) ν sl sol water and ligroin, ν sol alcohol and ether. Its solution in conc H_2SO_4 is blood red. It dissolves in conc HCl aq, but the base is ppd again on dilution — *Salts* — $B'H PtCl_6$ aq yellow laminae (from water) — $B'H_2PtCl_6$ 3aq (from dilute alcohol) — Picrate $B'C_6H_4(NO)_2OH$ [168°] Yellow nodules which blacken on fusion.

METHYL PHENTRIAZINE $C_8H_8N_4$ ν ϵ 

by the action of sodium amalgam on acetyl *o* nitro phenyl hydrazine in alcohol, the solution being kept acid with HOAc, and the temperature below 30° (Bischler, B 22, 2806). Crystals, partially decomposed on distillation. ν sol cold water, ν e sol alcohol, sl sol hot ligroin.

Bromo methyl phentriazine $C_8H_8BrN_4$ ν ϵ 

acetyl *p* bromo *o* nitro phenyl hydrazine by like treatment (Bischler & Brodsky, B 22, 2818). Golden plates, sl sol cold, ν sol hot, water.

(a) **METHYL - PHENTETRAZINE DIHY-**

Formed by the action of sodium nitrate on a very dilute hydrochloric acid solution of *o* amido-phenyl methyl hydrazine (Hempel, J pr [2] 41, 176). Colourless plates, ν sol ether and benzene, sl sol alcohol and petroleum ether. Sol hot conc NaOHAq. Conc HNO_3 at 80°–100° yields a product [127°] crystallising from alcohol in golden yellow prisms and needles.

TRI-METHYL PHENENYL TRIKETONE

$C_{15}H_{13}O_3$ *vs* $C_6H_5(COCH_3)_3$ [163°] *Tri acetyl-benzene* Formed by the action of NaOEt in ether on a mixture of acetone (1 mol) and allyl formate (1 mol) Sodium acetoacetic aldehyde slowly separates, and is then dissolved in ice cold water and neutralised by HOAc The free acetoacetic aldehyde changes to tri acetyl benzene and water (Claisen & Stylos, *B* 21, 1145) Small needles, sl sol water, alcohol, and ether, v sol HOAc Oxidised by HNO_3 to trimesic acid

METHYL - ($\alpha\beta$) - PHEN - NAPHTHAZINE

$C_{17}H_{13}N_3$ *vs* C_6H_5 $\begin{array}{c} \text{CH CH C N C CH CMc} \\ \text{CH CH C N C CH CH} \end{array}$ [142°] Formed by the action of (β)-naphthoquinone on tolylene diamine in HOAc (Hinsberg, *A* 237, 342)

Methyl-($\beta\beta$) phen-naphthazine

$C_{17}H_{13}N_3$ $\begin{array}{c} \text{CH C N C CH CMc} \\ \text{CH C N C CH CH} \end{array}$ [180°] Formed by oxidation of a mixture of (β) naphthol and tolylene *o*-diamine with alkaline K_2FeC_4 (Witt, *C N* 49, 404) Pale straw coloured needles, forming an intense red solution in H_2SO_4

METHYL PHENOL *v* CHLOROL**Di-methyl-phenol *v* XYLENOL**

Tri-methyl-phenol $C_9H_7Me_3OH$ [123 5] *Hemmellithenol Hemellithenol* [81°] Formed by fusing hemmellithene sulphonic acid with KOH (O Jacobson, *B* 19, 2518) Needles (from ether) Gives no pp with $FeCl_3$

Tetra-methyl-phenol $C_9H_5Me_4OH$ [123 4 5] [81°] (L), [87°] (T), (249° uncol) (L), (266°) (V) (T) Formed by the action of nitrous acid on the corresponding $C_9H_6Me_4NH$ [66°] (Limpach, *B* 21, 644), or by fusing *c* tetra-methyl-benzene sulphonic acid with potash (Töhl, *B* 21, 907) Long white needles (from very dilute alcohol), v e sol alcohol and ether, m sol petroleum ether Volatile with steam Bromine forms $C_{10}H_7BrOH$ [151°]

Acetyl derivative $C_{10}H_{13}OAc$ [57°]

Prisms

Tetra-methyl-phenol $C_9H(CH_3)_3OH$ [81°] Formed by the action of nitrous acid on tetra-methyl-phenyl amine [14°] (Hofmann, *B* 17, 1916) White crystals Yields a quinone on distillation with MnO_2 and H_2SO_4

Ethylether $C_9H(CH_3)_3OEt$ (236°), liquid Formed by the action of alcohol upon the sulphate of tetra-methyl-diazo benzene (Hofmann, *B* 17, 197)

Penta-methyl-phenol $C_9(CH_3)_5OH$ [125°] (267°) Formed by the action of nitrous acid upon penta-methyl-phenyl amine (Hofmann, *B* 18, 1826) Fine white needles Volatile with steam Soluble in alcohol Sparingly soluble in alkalis

Methyl ether $C_9(CH_3)_5OCH_3$ [64°], long needles

METHYL-DIPHENYL $C_{13}H_{11}$ *vs* $C_6H_5C_6H_5Me$ [3 1] *Phenyl-tolyl* (275°) SG 2 1031 Formed by the action of $MeCl$ upon melted diphenyl in presence of $AlCl_3$ (Adam, *Bl* [2] 47, 689, 49, 98, *A Ch* [6] 15, 239) Limpid colourless liquid, not solidified at -21° V e sol methyl alcohol and acetone Not attacked by $KMnO_4$, either in neutral or alkaline solution. Chromic acid oxidises it to diphenyl

m carboxylic acid [161°] Bromine at 150° forms $C_{12}H_9O_2Br$ which when treated with KOEt yields $C_{12}H_9CH_2OEt$ whence dry gaseous HI liberates phenyl benzyl alcohol

o Methyl-diphenyl $C_{13}H_{11}$ *vs* $C_6H_5C_6H_5Me$ [2 1] (259°) Appears to be formed by treating bromo benzene mixed with liquid bromo toluene with sodium (Barbier, *B* 7, 1548)

p Methyl-diphenyl $C_{13}H_{11}$ Me [4 1] (263°-267°) SG 2 1015 Formed by the action of sodium on an ethereal solution of bromo benzene and *p* bromo toluene (Carnelley, *C J* 29, 13) Formed also by passing a mixture of benzene and toluene through a red hot tube (Carnelley, *C J* 37, 701) Solidifies at -2° Dilute nitric acid oxidises it to diphenyl carboxylic acid. Chromic acid forms telephthalic acid

Di-methyl-diphenyl $C_{14}H_{12}$ *vs* $C_6H_5C_6H_5Me_2$ (284°-290°) SG 2 1025 Formed by the action of methyl chloride on diphenyl in presence of $AlCl_3$ (Adam) Colourless liquid, not solidified at -21° Oxidised by CrO_3 in HOAc yielding an infusible diphenyl dicarboxylic acid which cannot be sublimed

Other di-methyl-diphenyls are described as DIOLALS

DI - METHYL-PHENYLACETAMIDINE

$C_{16}H_{11}N$ *vs* $C_6H_5CH_2C(NHMe)(NMe)$ Formed by the action of alcoholic methylamine on the hydrochloride of phenylacetic imido ether $C_6H_5CH_2C(OEt)(NH)$ (Luckenbach, *B* 17, 1426) Crystalline solid, v sol alcohol - $BHCl$ six sided prisms - $B'H PtCl_4$ small glistening crystals, sol alcohol, sl sol water

u Di-methyl-phenylacetamide $C_6H_5CH_2C(NMe_2)(NH)$ Formed in the same manner as the preceding, using dimethylamine (L) - $B'H.H.PtCl_4$ small needles, sl sol water, m sol alcohol

DI-METHYL-PHENYL-ACETIC ACID

$[5\ 3\ 1]C_9H_7(CH_3)_2CH_2COH$ *Mesityl-acetic acid* [100°] (W), [97°] (Robmet, *Bl* [2] 40, 316) (273° at 735 mm) Formed by the saponification of the nitrile obtained by heating mesityl bromide with HCN (Wispek, *B* 16, 1578) Long white prisms Sol alcohol, ether, and hot water, v sl sol cold water

Salts - $A'K$ aq silky needles - $A'Ba$ 4aq transparent prisms - $A'Ca$ 3aq easily soluble thick needles - $A'Mg$ 5aq long silky needles - $A'Ag$ long thin needles

Tetra-methyl-phenyl-acetic acid $C_{11}H_9O_2$ *vs* $C_9HMe_4CH_2CO_2H$ [5 4 3 2 1] [125° uncol] Formed by reducing the corresponding tetra-methyl mandelic acid with HI (Claus & Fohlisch, *J pr* [2] 88, 234) Slender needles (from hot water), v sol alcohol, ether, and chloroform - $CaA'3aq$ silky needles

TRI-METHYL-PHENYL-AMIDO-CROTONIC ACID *v* ψ CUMYL AMIDO CROTONIC ACID

Tetra-methyl phenyl-amido-crotonic acid *Ethyl ether* $C_9HMe_4CH_2CO_2Et$ [101°] Obtained by the action of tetra-methyl phenyl amine (prepared from ψ -cumidine) on acetoacetic ether (Conrad & Limpach, *B* 21, 1655) Large white prisms (from alcohol and ether) At 280° - 285° it yields oxy tetra-methyl phenyl-di-methyl-pyridine carboxylic acid $C_9HMe_4N \begin{array}{c} \text{CMe} \\ \text{CMe} \end{array} \begin{array}{c} \text{C}(\text{CO}_2H) \\ \text{CO} \end{array}$ [145°]

METHYL-PHENYL AMINE *v* **TOLUIDINE** and **METHYL-ANILINE****Di-methyl-phenyl-amine** *v* **XYLIDINE** and **DI METHYL ANILINE****Tri-methyl-phenyl-amine** *v* **MESIDINE** and **ψ CUMIDINE**

Tetra methyl phenyl amine $C_6HMe_4(NH_2)$ [1 2 3 4 5] (66°) (260° uncor) Obtained by heating isocumidine hydrochloride with MeOH at 250° to 260° under pressure (Limpach, *B* 21, 641) Nacaceous leaflets (from water) May be sublimed The corresponding $C_6HMe_4(OH)$ melts at 81°

Formyl derivative [144°], silky needles (from water)

Acetyl derivative [170°], silky needles

Tetra methyl phenyl-amine $C_6HMe_4NH_2$ [1 2 3 5 6] *Isoduridine* [24°] (255°) Formed by heating the hydrochlorides of ψ cumidine or mesidine with MeOH at 300° (Hofmann, *B* 17, 1912, Noltung a Baumann, *B* 18, 1149, Limpach, *B* 21, 646) — $B'HCl$ — $B'H.PtCl_4$

Formyl derivative [183°], long silky needles

Acetyl derivative C_6HMe_4NHAc [215°]

Tetra methyl-phenyl amine $C_6HMe_4NH_2$ *Duridine* [14°] (253°) *SG* 21 978 A product of the action of MeOH on xylidine hydrochloride at a high temperature (Hofmann, *B* 17, 1913) — $B'HCl$ — $B'H.PtCl_4$

Penta methyl phenyl amine $C_6(CH_3)_5NH_2$ *Imido penta methyl ben ene* [152°] (278°) Prepared by heating dimethyl ψ cumidine with methyl iodide under pressure at 240°–20° (Hofmann, *B* 18, 1821) Large colourless needles *V* sol alcohol and ether, *insol* water On oxidation by acetic acid in conjunction with aniline it yields a homologue of rosaniline MeI at 100° forms $(MeNHMe)_6$, which is not further acted on by MeI even at 170°

Salts — $B'HCl$ long thin needles, easily soluble in hot water, *sl* sol cold — $B.H.Cl.PtCl_4$ sparingly soluble triumctic tablets — The acetate is very soluble the nitrate forms sparingly soluble needles, the sulphate and oxalate very sparingly soluble small scales

Acetyl derivative $C_6(CH_3)_5NHAc$ [213°], needles

METHYL DI PHENYL AMINE $C_6H_5NHCH_3$ *NPh Me Di phenyl methyl amine* (282°) (*G*) (292°) at 741 mm (Brühl, *A* 235, 21) *SG* 24 1 0476 Formed by methylation of diphenyl amine (Bardy, *Z* 1871, 649, Girard, *Bl* [2] 23, 2) Liquid Gives a violet colour with HNO_3 Fuming $HClAq$ at 150° resolves it into $MeCl$ and diphenylamine (Gnehm *B* 8, 1040) By passing through a red hot tube it is converted into diphenylamine, carbazole, benzonitrile, aniline, benzene, methane, hydrogen, nitrogen, and other products (Graebe, *A* 174, 181) Nitrous acid passed into its alcoholic solution yields $NMePh.C_6H_5.N_2.C_6H_5.NMePh$ Boiling nitric acid forms a compound which crystallises from alcohol in yellow prisms [234°]

Derivatives —Tri bromo and Bromo-DI NITRO, METHYL-DIPHENYLAMINE

METHYL-PHENYL-ANTHRACENE *v* **PHENYL-METHYL-ANTHRACENE****METHYL-PHENYL-CARBAMIC ACID** *v* **TOLYL CARBAMIC ACID****Di methyl phenyl carbamic acid** *v* **XYLYL-CARBAMIC ACID**

Tri methyl phenyl carbamic acid *Ethyl ether* $C_6H_3Me_3NH.CO.Et$ ψ *Cumyl-carbamic ether* [91 5°] Formed by the action of chloroformic ether $ClCO.Et$ on ψ cumidine (Frentzel, *C C* 1888, 1361) P_2O_5 converts it into the cyanate $C_6H_3Me_3NCO$ (221°), polymerised by PEt_3 or $KOAc$ into the cyanurate [234°]

Tri methyl phenyl carbamic acid *Ethyl ether* $C_6H_3Me_3NH.CO.Et$ *Mesityl-urethane* [62°] Formed from mesidine and $ClCO.Et$ (Eisenberg, *B* 15, 1016) Long colourless needles, volatile with steam, *sol* alcohol and ether

TETRA METHYL-PHENYL CARBAMINE $C_6H(CH_3)_4NC$ [51°] White crystals Formed by heating tetra methyl phenyl amine [14°] with chloroform and alcoholic KOH It is changed into the nitrile by distillation (Hofmann, *B* 17, 1914)

Penta-methyl phenyl carbamine $C_6(CH_3)_5NC$ [128°] Formed by heating penta-methyl phenyl amine with chloroform and alcoholic NaOH (Hofmann, *B* 18, 1824) Colourless crystals *V* sol alcohol At a few degrees above its melting point it is transformed into the nitrile with evolution of heat

METHYL TRI PHENYL CARBINOL CARBOXYLIC ACID *v* **DI PHENYL TOLYL CARBINOL-CARBOXYLIC ACID****METHYL PHENYLENE DIAMINE***v* **PHENYLENE METHYL DIAMINE****Di methyl phenylene diamine** *v* **PHENYLENE DI METHYL DIAMINE**

Tri methyl phenylene diamine $C_6H_3N_2$ *tc* $C_6HMe_3(NH_2)_2$ [1 2 4 3 5] (81°) Formed from nitro ψ cumidine by reduction (Mayer, *B* 20, 970) Long needles, *v* sol benzene $FeCl_3$ colours its solution deep red

Isomerides : **DI AMIDO MESITYLENE** and **DI-AMIDO ψ CUMYLENE**

Tetra-methyl phenylene diamine $C_6H_2N_2$ *tc* $C_6Me_4(NH_2)_2$ [1 2 4 3 5 6] *Prehnitylene-diamine* [140°] Formed by reducing $C_6Me_4(NH_2)_2(NO_2)_2$ with tin and HCl (Töhl, *B* 21, 906) Plates (from water), or needles (from alcohol), *v* sol alcohol, *sl* sol ether and ligroin — $B'H_2Cl_2Aq$ plates, *v* sol water, *v* sol conc $HClAq$ Coloured dark red by $FeCl_3$

Tetra-methyl phenylene diamine

$C_6Me_4(NH_2)_2$ [1 2 4 3 5 6] *Prehnitylene-diamine* [140°] Formed by reducing $C_6Me_4(NH_2)_2(NO_2)_2$ with tin and HCl (Töhl, *B* 21, 906) Plates (from water), or needles (from alcohol), *v* sol alcohol, *sl* sol ether and ligroin — $B'H_2Cl_2Aq$ plates, *v* sol water, *v* sol conc $HClAq$ Coloured dark red by $FeCl_3$

METHYL-DIPHENYLENE KETONE OXIDE *v* **PHENYLENE TOLYLENE KETONE OXIDE**

DI-METHYL PHENYLENE-DIVINYL DI-KETONE $C_6H_5CH=O_2$ *tc* $C_6H_5(CH=CH.CO.CH_3)_2$ [156°] Formed by the action of $NaOHAq$ on a mixture of acetone (10 pts) and terephthalaldehyde (1 pt) (Loew, *A* 231, 379) Needles, almost *insol* water, alcohol, and ether, *v* sol chloroform, *v* sol acetone With conc H_2SO_4 it yields a deep-red solution

METHYL-PHENYL-ETHYL-ALKINE *v* **METHYL-OXYETHYL ANILINE**

METHYL PHENYL-ETHYL KETONE v**BENZYL-ACETONE****METHYL PHENYL-GLYOXYLIC ACID v****TOLYL-GLYOXYLIC ACID****Di-methyl-phenyl glyoxylic acid v XYLIL-GLYOXYLIC ACID**

Tetra-methyl phenyl glyoxylic acid [6 5 3 2 1] $C_6H_5Me, CO CO_2H$ [124°] Formed by oxidising the ketone $C_6H_5Me, CO CH_3$ with cold aqueous $KMnO_4$ (Claus a Foecking, B 20, 8102) White scales, sl sol water, v sol alcohol and ether Reduced by sodium amalgam to tetra-methyl-mandelic acid $C_6H_5Me, CH(OH) CO_2H$ — KA' 5aq crystalline, v e sol water — CaA' 9aq nodules — BaA' 3aq nodules — AgA' white pp

Tetra-methyl phenyl glyoxylic acid [6 4 3 2 1] $C_6H_5Me, CO CO_2H$ Formed by oxidising the corresponding duryl methyl ketone with cold aqueous $KMnO_4$ (C a F) Yellow liquid, sl sol water, v sol alcohol and ether Solidifies when strongly cooled Sodium-amalgam reduces it in alcoholic solution to the corresponding tetra methyl mandelic acid — NaA' 5aq white crystalline crusts — BaA' 5aq nodules — CaA' 3aq granules — CuA' 5aq green crystals, v sol water — PbA' white pp, v sl sol water — AgA' white pp, insol water

Tetra-methyl-phenyl-glyoxylic acid $C_6H_5, O, \text{ etc } [5 4 3 2 1] C_6H_5Me, CO CO_2H$ Formed by oxidising the corresponding tetra methyl phenyl methyl ketone (Claus, J pr [2] 38, 232) Oil — BaA' 4aq needles — CaA' 4aq needles — CuA' 3aq

Penta-methyl-phenyl-glyoxylic acid $C_6H_5, O, \text{ etc } C_6Me, CO CO_2H$ [122°] Formed by the action of $AlCl_3$ on a mixture of penta methyl benzene and $ClCO CO_2Et$ (Jacobsen, B 22, 1218) Formed also by oxidising $C_6Me, CO CH_3$ with alkaline $KMnO_4$ Prisms, v sl sol cold water, v sol alcohol — NaA' 3aq plates, m sol cold water — BaA' 5aq nodules, v sl sol cold water — CuA' 5aq needles

HEXA METHYL DI-PHENYL GUANIDINE $C_6H_5, N, \text{ etc } HN C(NH C_6H_5Me)_2$ *Di mesityl guanidine* [218°] Formed by heating $SC(NH C_6H_5Me)_2$ with alcoholic NH_3 and lead oxide (Eisenberg, B 15, 1014) Minute prisms, sol. alcohol and ether, insol water

Hexa-methyl-tri-phenyl-guanidine $C_6H_5, N, \text{ etc } C_6H_5Me, N C(NH C_6H_5Me)_3$ [225°] Formed by heating hexa methyl di phenyl thio urea with alcoholic mesidine and lead oxide (E) Small crystals, sol alcohol, insol water

METHYL PHENYL-HYDRAZINE v PHENYL-METHYL-HYDRAZINE

Tri methyl-phenyl-hydrazine $C_6H_5, N, \text{ etc } [1 2 4 5] C_6H_5Me, NH NH_2$ ψ *Cumyl hydrazine* [120°] Formed by boiling its sodium sulphinate with water (Haller, B 18, 91) Needles (from ether), insol water and alkalis, v sol alcohol and ether With aceto acetic ether it yields oxy tetra methyl phenyl pyrazole

Sodium sulphinate $C_6H_5Me, NH NH SO_3Na$ Formed by warming diazo- ψ -cumene chloride with Na_2SO_3 and reducing the resulting $C_6H_5Me, N_2 SO_3Na$ with zinc-dust and $HOAc$ (Haller) White plates (containing 14aq), sl sol cold water and alcohol, v sol hot water

METHYL-PHENYL-DI-KETONE v. PHENYL-METHYL-DI-KETONE**TRI-METHYL-PHENYL MERCAPTAN**

$C_6H_5, S \text{ etc } C_6H_5Me, SH$ *Mesityl sulphhydrate. Thioresitol.* (229°) SG 10192 Formed by reducing mesitylene sulphochloride with zinc and H_2SO_4 (Holtmeyer, Z 1867, 686) Liquid, volatile with steam, v sol alcohol, ether, and benzene — $Hg(S C_6H_5Me)_2$ white silky needles

HEXA-METHYL-DI-PHENYL METHANE

$C_6H_5, \text{ etc } CH_2(C_6H_5Me)_2$ *Di-mesityl methane* [c 130°] Formed by the action of H_2SO_4 on a mixture of $CH_2(OAc)$, and mesitylene dissolved in $HOAc$ (Baeyer, B 5, 1098) Monoclinic prisms (from ether)

Methyl-tri-phenyl methane v Di PHENYL TOLYL METHANE

TETRA-METHYL-PHENYL-DI-METHYL AMINE $C_6H_5, N \text{ etc } C_6H_5Me, NMe_2$ (237°) Formed from tetra methyl phenyl amine [14°] by treatment with MeI and alcoholic soda (Hofmann, B 17, 1914) Colourless liquid — $B_2H_3PtCl_4$ crystalline

Penta-methyl-phenyl-methyl-amine $C_6H_5, N \text{ etc } C_6(CH_3), NHMe$ *Methyl-amido penta-methyl-benzene* [61°] Colourless scales Formed by heating penta methyl phenyl amine with methyl iodide — $B_2H_3Cl_2PtCl_4$ needles (Hofmann, B 18, 1824)

Penta methyl-phenyl-di methyl-amine $C_6(CH_3), NMe_2$ *Di-methyl-amido-penta-methyl-benzene* [54°] Colourless scales Formed by digesting penta methyl phenyl amine with methyl iodide in presence of an alkali Its methyl-iodide could not be obtained — $B_2H_3Cl_2PtCl_4$ needles (Hofmann, B 18, 1824)

TETRA-METHYL-PHENYL-METHYL-CARBINOL [6 4 3 2 1] $C_6H_5Me, CH(OH) CH_3$ *Duryl-methyl carbinol* (above 300°) Formed by reducing duryl methyl ketone with zinc and HCl (Claus a Foecking, B 20, 3099) Pale yellow liquid

Tetra methyl phenyl-methyl carbinol [6 5 3 2 1] $C_6H_5Me, CH(OH) CH_3$ [72°] Formed by reducing s duryl methyl ketone (C a F) White plates

METHYL PHENYL METHYL KETONE v TOLYL METHYL KETONE**Di methyl phenyl methyl ketone v XYLIL METHYL KETONE**

Tetra-methyl-phenyl methyl ketone $C_6H_5, O \text{ etc } [5 4 3 2 1] C_6H_5Me, CO CH_3$ *c Duryl methyl ketone* (260°) Formed from c durenene, $AcCl$, and $AlCl_3$ (Claus, J pr [2] 38, 231) Brownish, strongly refracting, oil, insol water, sol alcohol and ether Oxidised by $KMnO_4$ to tetra methyl-phenyl glyoxylic and c tetra methyl benzoic acids

Phenyl-hydrazide [129°] Plates **Tetra methyl-phenyl methyl ketone** [6 4 3 2 1] $C_6H_5Me, CO CH_3$ (254°) Formed from u durenene, $AcCl$, and $AlCl_3$ in CS_2 (Claus a Foecking, B 20, 3097) Colourless oil, v sol alcohol and ether Volatile with steam Reduced by zinc dust and alcoholic KOH to the carbinol. Oxidised by $KMnO_4$ to tetra methyl phenyl-glyoxylic acid

Phenyl-hydrazide Needles, decomposing at 215°

Oxim [148°] Plates **Tetra-methyl-phenyl methyl ketone** [6.5.3.2.1] $C_6H_5Me, CO CH_3$ [68°]. (251°).

Formed by the action of AlCl_3 upon *s* durenne mixed with AcCl in CS_2 (C & F) Plates

Phenyl hydrazide Crystals, decomposing at 225°

Penta-methyl-phenyl methyl ketone $\text{C}_{11}\text{H}_{14}\text{O}$ & $\text{C}_6\text{Me}_5\text{COCH}_3$ [85°] (286°) Formed by the action of AlCl_3 on a mixture of penta-methylbenzene and AcCl dissolved in CS_2 (Jacobsen, *B* 22, 1218) Pearly plates, v sol alcohol, ether, and HOAc Oxidised by KMnO_4 to penta-methyl phenyl glyoxylic acid


DI-METHYL-PHENYL-PHOSPHINE v

PHENYL DI METHYL PHOSPHINE

METHYL-TRI PHENYL PHOSPHONIUM

IODIDE v *Methyl iodide* of TRI PHENYL PHOSPHINE

TRI METHYL PHENYL PHTHALIDE

$\text{C}_6\text{H}_5\text{O}_2$ & C_6H_5  *Mesityl-phthalide* *Phenyl mesityl carbinol carboxylic anhydride* [164°] Formed by treating an alcoholic solution of [5 3 1 2] $\text{C}_6\text{H}_5\text{Me}$, CO C_6H_5 , CO H with zinc and HCl (Gresly, *A* 234, 237) Short thick needles

Tri-methyl-phenyl-phthalide $\text{C}_6\text{H}_5\text{O}_2$ v *Cumyl phthalide* [140°] Prepared as above, using the derivative of ψ cumene instead of that of mesitylene (G) Small needles

Methyl-di-phenyl-phthalide v **DI PHENYL**

TOLYL CARBINOL CARBOXYLIC-ANHYDRIDE

METHYL-PHENYL-PSEUDO-PICOL-STYRIL v **OXY-PHENYL DI METHYL PYRIDINE**

METHYL-PHENYL-PROPYL-ALKINE v **METHYL-OXYPROPYL ANILINE**

HEXA-METHYL-DI-PHENYL DISULPHIDE

$\text{C}_{12}\text{H}_{18}\text{S}_2$ & $(\text{C}_6\text{H}_5\text{Me})_2\text{S}$ *Mesityl disulphide* [125°] Formed by atmospheric oxidation of tri methyl phenyl mercaptan in alkaline solution (Holtmeyer, *Z* 1867, 688) Light yellow laminae or tables, insol water, sol alcohol, ether, and benzene

OCTA-METHYL-DI-PHENYL SULPHONE v

DI DURYL SULPHONE

Deca-methyl-di-phenyl sulphone $\text{C}_{22}\text{H}_{30}\text{SO}_2$ & $\text{C}_6\text{Me}_5\text{SO}_2\text{C}_6\text{Me}_5$ [93.5°] Formed by the action of ClSO_3H on penta methyl benzene (Jacobsen, *B* 20, 896) Very long, slender needles (from ligroin), v sol alcohol Yields penta methyl benzene when heated with conc HClAq at 170° , and in smaller quantity when distilled

TRI-METHYL-PHENYL-THIO-CARBAMIC

ETHER $\text{C}_6\text{H}_5\text{NSO}$ & $\text{C}_6\text{H}_5\text{Me}_2\text{NHCSOEt}$ or $\text{C}_6\text{H}_5\text{Me}_2\text{N}(\text{CSH})\text{OEt}$ *Mesityl thio urethane* [88°] Formed by heating the corresponding thiocarbimide with alcohol at 140° (Eisenberg, *B* 15, 1015) Slender needles, sol alcohol, ether, and warm alkalis

TRI-METHYL-PHENYL THIOCARBIMIDE

$\text{C}_6\text{H}_5\text{Me}$, NCS *Mesityl mustard oil* [64°] Formed by heating mesidine with CS_2 and some alcoholic potash (Eisenberg, *B* 15, 1012) Long needles, sol alcohol and ether

Tetra-methyl-phenyl-thiocarbimide

$\text{C}_{11}\text{H}_{12}\text{NS}$ & C_6HMe , NCS [65°] Formed by boiling tetra-methyl-phenyl amine [14°] with CS_2 and a little KOH (Hofmann, *B* 17, 1915) Crystalline solid

Penta-methyl-phenyl thiocarbimide

$\text{C}_{12}\text{H}_{13}\text{NS}$ & $\text{C}_6\text{Me}_5\text{NCS}$ [86°] Formed,

together with the corresponding thio-urea, by boiling penta methyl phenyl amine with CS_2 (Hofmann, *B* 18, 1827) Needles (from alcohol), volatile with steam

TRI METHYL PHENYL THIO UREA

$\text{C}_{10}\text{H}_{11}\text{N}_2\text{S}$ & $\text{NH}_2\text{CSNH C}_6\text{H}_5\text{Me}$, *Mesityl thio urea* [222°] Formed by the action of ammonia on the corresponding thiocarbimide (Eisenberg, *B* 15, 1013) Pearly plates, sol ether and hot alcohol, insol water

Hexa methyl di-phenyl thio urea $\text{C}_{12}\text{H}_{13}\text{N}_2\text{S}$

& $\text{CS}(\text{NH C}_6\text{H}_5\text{Me})_2$ *Di-mesityl thio-urea*

Formed by heating mesidine with CS_2 (Eisenberg, *B* 15, 1013) Formed also by digesting tri methyl phenyl thiocarbimide with mesidine in alcoholic solution White needles Conc H_3PO_4 converts it into the thiocarbimide

Isomeride v **DI ψ CUMYL THIO UREA**

Octo methyl di phenyl thio-urea $\text{C}_{12}\text{H}_{13}\text{N}_2\text{S}$

& $\text{SC}(\text{NH C}_6\text{HMe})_2$ [278°] Formed by the action of CS on tetra methyl phenyl amine [14°] (Hofmann, *B* 17, 1916) Four sided plates (from alcohol), sl sol alcohol

Deca methyl di phenyl-thio urea $\text{C}_{22}\text{H}_{22}\text{N}_2\text{S}$

& $\text{SC}(\text{NH C}_6\text{Me}_5)_2$ [252°] Formed by the action of CS upon penta methyl phenyl amine (Hofmann, *B* 18, 1827) White needles, sol HOAc v sl sol alcohol

TRI METHYL PHENYL-UREA v ψ **CUMYL UREA**

Hexa methyl-di phenyl-urea $\text{C}_{10}\text{H}_{11}\text{N}_2\text{O}$ & $\text{OC}(\text{NH C}_6\text{HMe}_5)_2$ *Di mesityl urea* [above 300°] Formed by the action of mesidine on tri methyl phenyl cyanate (Eisenberg, *B* 15, 1017) Minute prisms, sl sol hot alcohol, insol water

Hexa methyl di phenyl urea $\text{OC}(\text{NH C}_6\text{HMe}_5)_2$ *Di ψ cumyl urea* According to Frentzel (*C* 1888, 1361) this substance melts between 260° and 270° (cf vol 11 p 296)

METHYL-PHLOROGLUCINS v **PHLOROGLUCIN**

METHYL PHOSPHATES

Methyl-phosphoric acid $\text{MeO PO}(\text{OH})_2$ Formed by slowly adding methyl alcohol to cooled POCl_3 (H Schiff, *A* 102, 334) — BaA'' 2aq laminae, much less soluble in boiling water than in warm water — CaA'' (dried at 100°)

Di methyl phosphoric acid $(\text{MeO})_2\text{PO OH}$ Formed by pouring POCl_3 into methyl alcohol (Schiff) Acid syrup, sol alcohol and ether Its salts are more soluble than those of the preceding acid — BaA' (dried at 150°) pearly plates, nearly insol alcohol — SrA' 2aq — PbA' 2aq — CaA' (dried at 100°) very soluble nodules

Tri methyl phosphate Me_3PO_4 (197° cor) *SG* 2 1238 (Weger, *A* 221, 89) *SV* 139 5 (Lossen, *A* 254, 74)

METHYL-PHOSPHINE CH_3P & CH_3PH_2 Mol w 48 (-14°) *VD* 24 4 (calc 24) *S* (ether) 70 at 0° Formed, together with di methyl phosphine, by heating PH_3I with MeI and ZnO at 150° When water is added to the product MePH_2I is decomposed, with evolution of MePH_3 , while $\text{Me}_2\text{PH}_2\text{I}$ remains (Hofmann, *B* 4, 605) Formed also by heating chloroform with PH_3I and ZnO (Hofmann, *B* 6, 302) Gas, with powerful odour May be condensed by 2 atmospheres' pressure at 0° Fumes in the air, and takes fire when gently warmed Feeble base, being absorbed by conc HClAq or conc.

HIAq, producing crystalline salts, which are decomposed by water into MePH_2 , and the acid. When passed into fuming HNO_3 , it yields **methane phosphonic acid** or 'methyl-phosphinic acid' $\text{MePO}(\text{OH})_2$, which forms crystals, v sol water, melting at 105° . This acid is not attacked by aqua regia. It yields the salts MePO_3Ba , MePO_3Pb , and MePO_3Ag , and the chloride MePOCl_2 [32?] (163°).

Salts— MePH_2HCl four sided plates, very volatile in the air, being dissociated— MePH_2HI laminae.

Di-methyl phosphine $\text{C}_2\text{H}_5\text{P} \text{ se } (\text{CH}_3)_2\text{PH}$ Mol w 62 (25°) Obtained by the action of caustic soda on its hydro iodide, which is obtained as above described (Hofmann, B 4, 610). Very volatile liquid which takes fire in the air. May be oxidised to Me_2POOH , a waxy solid, v sol water. This acid melts at 76° , forms crystalline Me_2POOAg and is converted by PCl_5 into the chloride Me_2POCl (66°), (204°). This chloride is reconverted by water into the corresponding acid with less energy than MePOCl_2 (Hofmann, B 5, 109, 6, 307).

Tri methyl phosphine $\text{C}_3\text{H}_7\text{P} \text{ se } \text{P}(\text{CH}_3)_3$, Mol w 76 (41°)

Formation—1 By the action of MeCl upon calcium phosphide (Thénard, C R 21, 144, 25, 892)—2 From sodium or zinc phosphide and MeI (Cahours a Hofmann, A Ch [3] 41, 631, Chem Gaz 1855, 11)—3 From PCl_5 and ZnMe , (Hofmann a Cahours, C R 104, 29)—4 From PH_3 and MeI (Drechsel a Finkelstein, B 4, 354, Hofmann, B 4, 205, 430)—5 By heating CS_2 with PH_3I at 140° (Drechsel, J pr [2] 10, 180). 6 From phosphorus and MeI (Friedel a Silva, Wurtz's Dict 2, 938).

Preparation—By treating PCl_5 with ZnMe , in an atmosphere of CO_2 , decomposing the product with caustic soda, and distilling in a very slow current of hydrogen.

Properties—Volatile oil with powerful nauseous odour, heavier than water. Forms readily soluble, crystallisable salts. Unites readily with halogens, oxygen, and sulphur. With $\text{ClCH}_2\text{CO}_2\text{H}$ at 100° it yields $\text{PMe}_2\text{ClCH}_2\text{CO}_2\text{H}$ which forms the platinumchloride $(\text{C}_2\text{H}_5)_2\text{PO}_2\text{PtCl}_4$ (Meyer, B 4, 734, C J 24, 1066).

Salts— $\text{B}'\text{H}_2\text{PtCl}_4$ orange yellow crystalline pp— $\text{B}'\text{H}_2\text{PtCl}_4$ (Cahours a Gal, Z 1870, 662). Combines with CS_2 forming pale red crystals of PMe_2CS_2 , which slowly changes, in ethereal solution, into PMe_2S (Hofmann, A Suppl 1, 59).

Oxide PMe_2O [138°] (Collie, C J 53, 637) (215°). Formed by exposing PMe_2 to a slow current of dry air. Deliquescent crystals. Formed also, together with CH_3 , by distilling PMe_2OH . Yields a platinumchloride $(\text{PMe}_2\text{O})_2\text{H}_2\text{PtCl}_4\text{aq}$ crystallising in orange plates or needles.

Sulphide PMe_2S [105°]. Obtained by gradually adding flowers of sulphur to PMe_2 , or by distilling PMe_2 with cinnabar. Not formed from the oxide and H_2S or ammonium sulphide. Four sided prisms (from concentrated aqueous solution). When warmed with a solution of a silver salt Ag_2S is deposited as a black mirror (Collie).

Selenide PMe_2Se [84°]. Formed from PMe_2 and selenium. Blackens in the air, depositing selenium.

Methylo-hydroxide PMe_2OH Formed

by treating PMe_2I with moist Ag_2O . Caustic base, split up on distillation into PMe_2O and CH_3 .

Methylo-iodide PMe_2I **Tetra methyl-phosphonium iodide** Formed from PMe_2 and MeI (Hofmann a Cahours, A 104, 81). Formed also, together with PMe_2 , by heating PH_3I (1 mol) with MeOH (3 mols) at 180° (Hofmann, B 4, 208). Silvery crystals.

Methylo chloride PMe_2Cl From PMe_2OH and HCl . Deliquescent crystals, decomposed by heat into PMe_2HCl and ethylene (Collie, C J 53, 637)— $(\text{PMe}_2\text{Cl})_2\text{PtCl}_4$ glistening yellow octahedra, insol water— $\text{PMe}_2\text{AuCl}_4$: brilliant yellow needles (from boiling water).

Methylo sulphate $(\text{PMe}_2)_2\text{SO}_4$ Formed from PMe_2I and Ag_2SO_4 (Collie). Thick deliquescent needles. Does not form an alum with $\text{Al}(\text{SO}_4)_3$. Above 300° it decomposes, giving PMe_2O and PMe_2S .

Methylo carbonate $\text{PMe}_2\text{O.CO}_2\text{H}$ Decomposes above 100° into PMe_2O , CO_2 , and CH_3 (Collie).

Methylo acetate PMe_2OAc Decomposed by heat chiefly into PMe_2O and acetone, with traces of PMe_2 and MeOAc .

Benzoate PMe_2OBr Formed by neutralising a solution of PMe_2OH with HOBz (Collie, C J 53, 639). Deliquescent crystals, v sol water. Decomposes between 250° and 300° almost completely into PMe_2O and acetophenone.

Ethylo chloride PMe_2EtCl Obtained from PMe_2EtI (Collie, C J 53, 718). Decomposes above 300° into PMe_2HCl , PMe_2EtHCl , and ethylene— $(\text{PMe}_2\text{EtCl})_2\text{PtCl}_4$ octahedra.

Ethylo iodide PMe_2EtI **Tri-methyl ethyl phosphonium iodide** From PMe_2 and EtI in ether (Cahours a Hofmann, A 104, 33). Crystallises from boiling water. Deliquescent, sol ether.

Isoamyl iodide $\text{PMe}_2\text{C}_4\text{H}_9\text{I}$ Deposited slowly from an ethereal solution of PMe_2 and $\text{C}_4\text{H}_9\text{I}$ (Hofmann). Needles (from alcohol). Yields $(\text{PMe}_2\text{C}_4\text{H}_9\text{Cl})_2\text{PtCl}_4$, crystallising from boiling water in splendid needles aggregated in spherules.

Bromo ethylo-bromide $\text{C}_2\text{H}_5\text{Br}_2 \text{ se } \text{CH}_2\text{BrCH}_2\text{PMe}_2\text{Br}$ From PMe_2 and a large excess of ethylene bromide in alcoholic solution at 60° . Triclinic prisms, $a b c = 56.8 : 1 : 407$ (Hofmann, Tr 1860, 590). With Ag_2O it yields $\text{CH}_2(\text{OH})\text{CH}_2\text{PMe}_2\text{OH}$ whence the platinum chloride $(\text{CH}_2(\text{OH})\text{CH}_2\text{PMe}_2\text{Cl})_2\text{PtCl}_4$ may be got.

Ethyleno di-bromide $(\text{PMe}_2\text{Br})_2\text{O.H}_2$ Formed from ethylene bromide and excess of PMe_2 at 100° . Very deliquescent monoclinic crystals, $a b c = 1.054 : 1.1126, \beta = 87^\circ 49'$. Yields $\text{C}_2\text{H}_4(\text{PMe}_2\text{Cl})_2\text{PtCl}_4$, which crystallises from boiling HClAq in golden yellow laminae. Successive treatment with moist Ag_2O and HI forms $\text{C}_2\text{H}_4(\text{PMe}_2)_2$, crystallising in sparingly soluble needles.

Tri-methyl-tri-ethyl-di-phosphine ethyleno-di-bromide $(\text{PMe}_2\text{Br})_2\text{C}_2\text{H}_4(\text{PEt}_2\text{Br})_2$ Formed from PMe_2 and $\text{CH}_2\text{BrCH}_2\text{PEt}_2\text{Br}$. Yields a caustic hydroxide and the platinumchloride $(\text{PMe}_2\text{Cl})_2\text{C}_2\text{H}_4(\text{PEt}_2\text{Cl})_2\text{PtCl}_4$.

Tetra-methyl-di-phosphine? P_2Me_4 or $(\text{PMe}_2)_2$ (250°) Found among the products of the ac-

tion of MeCl on calcium phosphide or of MeI on sodium phosphide (Thénard, Hofmann a Cahours, *A* 104, 4) Liquid, with unpleasant odour, taking fire in the air Insol water Decomposed by HCl into PMe_3 , and a yellow powder $\text{P}(\text{Me})_2$ (?)

METHYL PHOSPHITE MeH_2PO_3 , *Methyl phosphorous acid* Formed by gradually adding PCl_3 to methyl alcohol (Schiff, *A* 103, 164) Acid syrup, resolved by heat into MeOH and phosphorous acid Its salts are slightly crystalline hygroscopic masses, v sol water, sl sol alcohol, insol ether In aqueous solution they decompose, slowly in the cold, but rapidly on heating, into MeOH and metallic phosphites — $\text{Ba}(\text{MeHPO}_3)_2 \cdot 2\text{aq}$ — $\text{Pb}(\text{MeHPO}_3)_2$

Tri methyl phosphite Me_3PO , (185°) SG 11785 Formed from PCl_3 and NaOMe (Jachin, *A* 256, 281) The compound $\text{Me}_3\text{PO} \cdot \text{PtCl}_3$ is formed by the action of MeOH on PCl_3 , PtCl_3 (Schutzenberger, *B* 2 18, 101 157) It crystallises in orange needles and yields $(\text{Me}_3\text{PO})_2 \cdot \text{PtCl}_3$ and $(\text{Me}_3\text{PO})_2 \cdot \text{PtCl}_3 \cdot \text{H}_2\text{O}$

METHYL-PHTHALIC ACID v *1,2-DIMETHYL-3-CARBOXYLIC ACID*

Di-methyl phthalic acid v *1,2-DIMETHYL-3-CARBOXYLIC ACID*

Tetra methyl phthalic acid v *1,2,3,4-TETRAMETHYL-3-CARBOXYLIC ACID*

METHYL-PHTHALIDE $\text{C}_8\text{H}_6\text{O}$ 12

$\text{C}_8\text{H}_6 \cdot \begin{smallmatrix} \text{CHMe} \\ \text{CO} \end{smallmatrix} > \text{O}$ (276°) VD 74 (obs and calc) Formed by inducing acetophenone carbonylic acid with sodium amalgam and acidifying the resulting $\text{CO} \cdot \text{Na} \cdot \text{C}_6\text{H}_5 \cdot \text{CH}(\text{OH}) \cdot \text{CH}_3$ (Gabriel a Michael, *B* 10 2205, 20 2500) Thick oil solidifying below 0 Insol water and cold alkalis, v sol alcohol and ether sl sol lichen Boiling alkalis convert it into salts of the acid, which forms a crystalline silver salt $\text{CO} \cdot \text{Ag} \cdot \text{C}_6\text{H}_4 \cdot \text{CH}(\text{OH}) \cdot \text{CH}_3$

Di methyl-phthalide $\text{C}_{10}\text{H}_8\text{O}$ 12

$\text{C}_{10}\text{H}_8 \cdot \begin{smallmatrix} \text{CO} \\ \text{CO} \end{smallmatrix} > \text{O}$ [65° (160°) at 25 mm, (271°) at 760 mm Formed by the action of zinc dust and MeI on phthalic anhydride at 100° under a slightly increased pressure (Wislicenus, *A* 218, 66) Crystallises from ether in large crystals which are doubly refracting

Reactions — 1 Converted by the action of conc KOHAq into the k salt of *o* oxy iso propyl benzoic acid — 2 Sodium amalgam yields the dihydride $\text{C}_8\text{H}_8 \cdot \begin{smallmatrix} \text{CMe} \\ \text{CH}(\text{OH}) \end{smallmatrix} > \text{O}$ [90°], a yellow amorphous powder, insol water, sol alcohol and ether, which reduces Fehling's solution and ammoniacal AgNO_3 — 3 Reduction with HI yields *o* iso propyl benzoic acid — 4 Potassium cyanide at 250° forms *o* propenyl benzoic acid $\text{CH}_2 \cdot \text{CMe} \cdot \text{C}_6\text{H}_4 \cdot \text{CO} \cdot \text{H}$ [61°]

METHYL PHTHALIMIDE v *Methylamide of PHTHALIC ACID*

METHYL-PHTHALIMIDINE $\text{C}_8\text{H}_7\text{NO}$ 12

$\text{C}_8\text{H}_7 \cdot \begin{smallmatrix} \text{CH} \\ \text{CO} \end{smallmatrix} > \text{NMe}$ [120°] (300°)

Formation — 1 By reducing methyl phthalimide $\text{C}_8\text{H}_5 \cdot \begin{smallmatrix} \text{CO} \\ \text{CO} \end{smallmatrix} > \text{NMe}$ with tin and HClAq (Graebe, *A* 247, 303) — 2 By heating a solution of phthalimidine in aqueous KOH with excess

of MeI for 6 hours at 100° (Barbier, *C. R* 107, 918) — 3 By heating phthalide with alcoholic methylamine for 12 hours at 220° (B)

Properties — Slender needles or plates, v sol^r water, alcohol, and ether Oxidised by KMnO_4 to methyl phthalimide and, finally, to phthalic acid Combines with bromine forming $(\text{C}_8\text{H}_7\text{NO})_2\text{Br}_2$, crystallising in needles [150°]

Salts — Hydrochloride [120°] Prisms $\text{B} \cdot \text{H} \cdot \text{H} \cdot \text{Cl}$, yellow prisms, sl sol cold water a **METHYL HOMO O PHTHALONITRILE** v *o* **Cyano phenyl propionitrile**

METHYLPYRAZOLENOL v *Organic SELENIUM COMPOUNDS*

METHYL PIAZTHIOLE $\text{C}_4\text{H}_5\text{N}_2\text{S}$ 12

$\begin{bmatrix} 3 & 6 \\ 5 & \end{bmatrix} \text{C}_4\text{H}_5\text{N}_2\text{S}$ [34°] (234°) Mol w

(by Rioult's method) 143 (calc 150) Formed by heating tolylene *o* diamine with H_2SO_4 at 180° (Hinberg, *B* 22 2900) Yields a per iodide when treated with I in HIAq Br in CHCl_3 forms $\text{C}_4\text{H}_5\text{N}_2\text{S}$ which crystallises in white needles; 98° A mixture of conc H_2SO_4 and HNO_3 forms $\text{C}_4\text{H}_5(\text{NO}_2)\text{N}_2\text{S}$ crystallising in colourless needles [156°]

Salt — $\text{B} \cdot \text{H} \cdot \text{PtCl}_3$, reddish yellow crystals, decomposed by water

METHYL PIPECOLINE v *DI METHYL PYR*

IMINI HEXAHYDRIDE

METHYL PIPERIDINE v *METHYL PIPRIDINE*

HEXAHYDRIDE

METHYL PROPARGYL-AMINE $\text{C}_4\text{H}_7\text{N}$ 12

$\text{C}_4\text{H}_7 \cdot \text{NH} \cdot \text{C} \equiv \text{C} \cdot \text{CH}_3$ Formed by the action of MeI on propargylamine in alcoholic solution (Paal a Hermann, *B* 22, 3083) Very volatile yellowish liquid, with ammoniacal odour — $\text{B} \cdot \text{HI}$ [83°] (groups of long hygroscopic needles — $\text{B} \cdot \text{H}_2\text{C} \cdot \text{O}$, [141] Slender white needles, sl sol alcohol

METHYL-PROPIONIC ACID v *BUTYRIC*

ACID

METHYL PROPYL ACETAL v *ALDOPHYDE*

METHYL PROPYL ACETIC ACID v *HEXOIC*

ACID

METHYL PROPYL-ACETOACETIC ETHER

v *ACETOACETIC ACID*

METHYL ISOPROPYL-ACETONE v *METHYL*

ISOPROPYL KETONE

DI-METHYL-PROPYL-ALKINE v *DI-*

METHYL OXYPROPYL AMINE

DI METHYL ISOPROPYL-ALLYL-CAR-

BINOL v *CANFENYL ALCOHOL*

TRI METHYL PROPYL AMMONIUM

IODIDE NMe_3PrI *Propylo iodide of trimethylamine* [190°] Formed by heating propylamine with alcoholic NMe_3 (Langeli, *G* 16, 385) Long needles Yields crystalline NMe_3PrCl and $(\text{NMe}_3\text{PrCl})_2\text{PtCl}_3$ The hydroxide NMe_3PrOH is decomposed by heat into propylene and NMe_3

METHYL-PROPYL ANILINE $\text{C}_{10}\text{H}_9\text{N}$ 12

$\text{C}_6\text{H}_5 \cdot \text{NMePr}$ (212° uncor) Liquid Prepared by the action of methyl iodide upon propylamine (Claus a Hinz, *B* 19, 2785, cf Nolting, *J* 1883, 702) — $\text{B} \cdot \text{HCl}$ very hygroscopic crystals, [106°]

Ethyl-o-iodide v *Propylo iodide* **METHYL-**

ETHYL-ANILINE

METHYL-PROPYL-BENZENE v *CYMENE*

C_9H_{10} s *Methyl di-propyl benzene* $\text{C}_{11}\text{H}_{14}$ 12 $\text{C}_6\text{H}_5\text{Me}(\text{C}_2\text{H}_5)_2$ [35] (248°-248°) Formed by

the action of H_2SO_4 on a mixture of acetone and methyl *n* propyl ketone (Jacobsen, *B* 8, 1259) Yields uvitic acid on oxidation with dilute HNO_3 .

***s*-Di methyl propyl-benzene** $\text{C}_{11}\text{H}_{16}$ *is* $\text{C}_6\text{H}_5\text{Me}_2\text{C}_3\text{H}_7$ [135] (206°-210°) Formed by the action of H_2SO_4 (3 vols) on a mixture of acetone (4 vols) and methyl propyl ketone (2 vols) (Jacobsen, *B* 8, 1259) Oxidised by boiling dilute HNO_3 to mesitylenic acid

***u*-Di-methyl-propyl-benzene** $\text{C}_{11}\text{H}_{16}$ *is* $\text{C}_6\text{H}_5\text{Me}_2\text{Pr}$ [142] (206°) Formed from bromo-*p*-xylene, propyl bromide, and sodium (Uhlhorn, *B* 23, 2350) Liquid, not solidified at -20° Yields a tri-nitro derivative [85°], and a tri-bromo-derivative [49°]

Sulphonic acid $\text{C}_6\text{H}_5\text{Me}_2\text{Pr SO}_3\text{H}$ Salts — BaA' — NaA' 1½aq —

Amide $\text{C}_6\text{H}_5\text{Me}_2\text{Pr SO}_2\text{NH}_2$ [125°] —
Anilide $\text{C}_6\text{H}_5\text{Me}_2\text{Pr SO}_2\text{NHPh}$ [216°]

***u*-Di-methyl-propyl-benzene** $\text{C}_6\text{H}_5\text{Me}_2\text{Pr}$ [134] (208°) Formed from bromo-*m*-xylene, propyl bromide, and Na (Uhlhorn) Liquid, not solid at -20° Yields a tri-nitro derivative [110°] and a tri-bromo-derivative [89°] May be oxidised to (1,3,4) di-methyl-benzoic acid

Sulphonic acid $\text{C}_6\text{H}_5\text{Me}_2\text{Pr SO}_3\text{H}$ Groups of needles — Salts — NaA' 4½aq —
 BaA' 2aq — MgA' 5aq —
Amide $\text{C}_6\text{H}_5\text{Me}_2\text{Pr SO}_2\text{NH}_2$ [102°] —
Anilide $\text{C}_6\text{H}_5\text{Me}_2\text{Pr SO}_2\text{NHPh}$ [182°]

***u*-Di methyl-propyl-benzene** $\text{C}_6\text{H}_5\text{Me}_2\text{Pr}$ [124] (209°) Formed from bromo-*o*-xylene, propyl bromide and sodium (Uhlhorn, *B* 23, 2349) Liquid, not solid at -20° Yields on oxidation (1,2,4) di methyl benzoic acid [162°]

Sulphonic acid $\text{C}_6\text{H}_5\text{Me}_2\text{Pr SO}_3\text{H}$ Thin needles — Salts — NaA' 3½aq —
Amide $\text{C}_6\text{H}_5\text{Me}_2\text{Pr SO}_2\text{NH}_2$ [124°] —
Anilide $\text{C}_6\text{H}_5\text{Me}_2\text{Pr SO}_2\text{NHPh}$ [214°].

***u* Di methyl-isopropyl benzene** $\text{C}_6\text{H}_5\text{Me}_2\text{Pr}$ [134] (194°) Formed from bromo-*m*-xylene, isopropyl bromide, and sodium (U) Liquid Yields a tri-bromo-derivative [261°] and a tri-nitro-derivative [182°]

Sulphonic acid $\text{C}_6\text{H}_5\text{Me}_2\text{Pr SO}_3\text{H}$.

Needles — Salts — NaA' 4aq — BaA' 2 —
Amide $\text{C}_6\text{H}_5\text{Me}_2\text{Pr SO}_2\text{NH}_2$ [163°] —
Anilide $\text{C}_6\text{H}_5\text{Me}_2\text{Pr SO}_2\text{NHPh}$ [207°]

METHYL-PROPYL-BENZOIC ACID $\text{C}_{11}\text{H}_{14}\text{O}_2$ *is* $\text{C}_6\text{H}_5\text{Me}(\text{C}_3\text{H}_7)\text{CO}_2\text{H}$ *Carbocymelic acid* [63°] Obtained from its nitrile, which is produced by distilling potassium cymene sulphonate with KCy (Paterno a Fileti, *B* 8, 442, Paterno a Spica, *G* 9, 400)

Amide $\text{C}_6\text{H}_5\text{Me}(\text{C}_3\text{H}_7)\text{CONH}_2$ [139°] Obtained by boiling the nitrile with alcoholic potash Needles, sl sol cold water

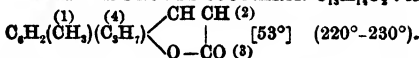
METHYL-PROPYL-CARBINOL *v* *Sec*-AMYL ALCOHOL

Di-methyl-propyl-carbinol *is* *Tert*-HEXYL ALCOHOL

Methyl-di-propyl-carbinol *v* OCTYL ALCOHOL.

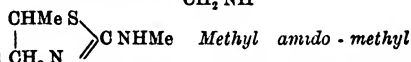
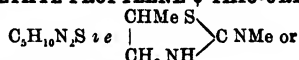
METHYL PROPYL CARBONATE $\text{C}_8\text{H}_{16}\text{O}_3$ *is* $\text{CH}_3\text{O CO OC}_3\text{H}_7$. (131° cor) SG $\frac{d}{4}$ 978 (Röse, *A* 205, 230)

METHYL-PROPYL-COUMARIN $\text{C}_{13}\text{H}_{14}\text{O}_2$ *is* *a*.



Formed by heating a mixture of thymol, malic acid and H_2SO_4 (Pechmann a Welsh, *B* 17, 1647) Fine white needles V sol alcohol, ether, benzene, acetic acid, and chloroform, v sl sol water

METHYL PROPYLENE ψ THIO-UREA



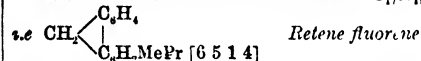
thiazole dihydride [50°] Formed by the action of methyl thiocarbimide on *B* oxy propylamine (Hirsch, *B* 23, 971) Long needles (from ligroin), v sol water Its aqueous solution is strongly alkaline — $\text{B}'\text{H PtCl}_6$ [143°] Large dark-red needles (from hot water) — $\text{B}'\text{C}_6\text{H}_5(\text{NO}_2)_2\text{OH}$ [145°] Needles — Compound with methyl thiocarbimide $\text{B}'\text{MeNCS}$ [64°] White columns (from MeOH)

Methyl propylene ψ thio-urea $\text{C}_3\text{H}_5\text{N}_2\text{S}$ *is* $\text{CHMe S} \text{ } \begin{array}{c} \text{CH}_2 \text{ NMe} \end{array} \text{C NH Imido-di methyl-thiazole}$

tetrahydride Formed from propylene ψ thio-urea, MeI, and KOH (Gabriel, *B* 22, 2989) Converted by bromine water into the acids $\text{NHMe CH}_2\text{CHMe SO}_3\text{H}$ [220°-223°] and $\text{CONH}_2\text{NMe CH}_2\text{CHMe SO}_3\text{H}$ [230°-240°]

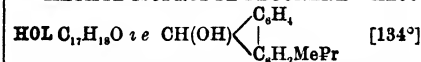
METHYL PROPYL-ETHANE *v* *HEXANE*

METHYL ISOPROPYL FLUORENE $\text{C}_{11}\text{H}_{16}$



[97°] Formed by passing the vapour of methyl isopropyl di phenylene ketone (retene ketone) through a red hot tube (Bamberger a Hooker, *A* 229, 142) Formed also by heating this ketone with HIAq and phosphorus in sealed tubes White plates, v sol cold ether, hot alcohol, and HOAc In the fused state, or in alcoholic solution, it exhibits violet fluorescence It yields a di-nitro derivative [245°]

METHYL ISOPROPYL FLUORENE **ALCO**



Formed by the reduction of methyl isopropyl di phenylene ketone (Bamberger a Hooker, *A* 229, 144) White needles, insol water, v sol alcohol and ether Readily oxidised back to the ketone The acetate melts at 71°

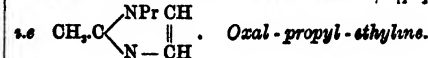
METHYL PROPYL GLUTARIC ACID

$\text{CO}_2\text{H CHPr CH}_2\text{CHMe CO}_2\text{H}$ [102°] Formed from sodium propyl-malonate and bromo isobutyric ether, the product being saponified, and the resulting $(\text{CO}_2\text{H})_2\text{CHPr CH}_2\text{CHMe CO}_2\text{H}$ [168°] being decomposed by heat (Bischoff, *B* 23, 1940) White aggregates of crystals (from petroleum-ether)

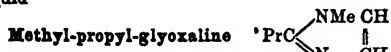
DI METHYL-PROPYL GLYCOLLINE *v* *Di*-

METHYL DI-OXY PROPYL-AMINE

METHYL PROPYL-GLYOXALINE $\text{C}_8\text{H}_{12}\text{N}_2$



(226°). SG 9641 Formed by the action of propyl bromide on methyl glyoxaline (glyoxal-ethylene) (Radziszewski, *B* 16, 489) Colourless liquid



Oxal-methyl-butylene (215° at 722 mm) SG 985 From propyl glyoxaline and MeI (Rieger, *M* 9, 606) Oil, v. sl alcohol, ether, and chloroform—B⁺H₂PtCl₄, orange red prisms

METHYL PROPYL-GLYOXAL v METHYL-

PROPYL-DIKETONE

DI-METHYL PROPYLDIENE DIKETONE C₈H₁₄O₂ i.e. (CH₃CO)₂CHET *Ethyl-acetyl acetone* (179°) Prepared by heating the sodium derivative of methylene di methyl diketone with EtI at 140° (Combes, *A Ch* [6] 12, 248, *C R* 104, 920) Colourless liquid with pleasant odour, sl sol water, miscible with ether, alcohol, and chloroform Decomposed by potash into CH₃CO₂K and CH₃COCH₂Et Combines with NaHSO₄ Sodium yields (CH₃CO)₂CNaEt which reacts with alkyl iodides

METHYL PROPYL KETONE C₈H₁₆O i.e. CH₃COCH₂CH₂CH₂CH₃ *Ethyl acetone* (102°) SG 8124, 88 8044 (Perkin, *C J* 45, 479), 22 805 (F a D) H F p 72,410 H F v 69,400 (Thomson, *Th*) M M 5 499 at 16.1° (P)

Formation—1 By distilling calcic acetate (48 g) mixed with calcic butyrate (65 g), and rectifying the product The yield (10 g) is small (Semljantzin, *J pr* [2] 23, 263, cf Friedel, *A Ch* [4] 16, 366, *A* 108, 124, Grimm, *A* 157, 251)—2 By boiling ethyl acetoacetic ether with potash or baryta (Frankland & Duppa, *A* 133, 216)—3 By the action of ZnMe₂, followed by water, on butyryl chloride (Butlerow, *Bl* [2] 5, 17)—4 By the oxidation of sec n amyl alcohol (Wurtz, *A* 148, 133, Schorlemmer, *C J* 25, 1085, *A* 161, 269, Wagner & Saytzeff, *A* 179, 322)—5 By the action of water and HgBr₂ on valerylène (Kutscheroff, *B* 14, 1542)—6 By the action of KOH on (CH₃CO)₂CHET (Combes, *A Ch* [6] 12, 248)

Properties—Liquid, v. sl sol water Combines with hydrogen sodium sulphite, forming C₈H₁₆(OH)SO₃Na₂aq (Grunm) Reduced by means of sodium amalgam to sec amyl alcohol CH₃CH(OH)CH₂CH₂Et (119° cor) (Belohoubek, *Sitz W* [2] 74, 80) and a pinacone C₁₀H₂₂O (225°–230°) Yields acetic and propionic acids on oxidation (Schorlemmer), Wagner obtained butyric acid (*Bl* [2] 38, 264) PCl₅ forms CH₃CCl₂CH₂CH₂Et, whence alcoholic potash produces CH₃CH₂CH₂CH₂Et, and this is converted by alcoholic potash at 170° into CH₃CCET (56°) (Kavorsky, *J R* 1887 414) CH₃CCl₂CH₂CH₂Et is split up on distillation into HCl and C₈H₁₆Cl (c 95°) Amyl nitrite and HCl or NaOEt forms the nitroso-derivative C₈H₁₅COCHNOH (48°–51°), which is also an oxim of propyl-glyoxal From this oxim may be prepared the compounds C₈H₁₅C(NOH)CH(NOH) [168°] and C₈H₁₅C(N,HPh)CH(N,HPh) [163°] (Claisen & Manasse, *B* 22, 528)

Oxim C₈H₁₅C(NOH)CH₂ Oil Converted by gaseous HCl in HOAc at 100° into propylamine and acetic acid (Beckmann, *B* 20, 2580)

Methyl isopropyl ketone C₈H₁₆O i.e. CH₃COCH(CH₃)₂ *Di methyl acetone* 'Amyl

ene oxide' (94°) SG 2 822, 12 805 (Wino-gradoff), 12 810 (F a D) CE (0°–18°) 00118

Formation—1 By the action of baryta water on di methyl aceto acetic ether (Frankland & Duppa, *Pr* 14, 463, *A* 138, 832)—2 By the dry distillation of a mixture of calcium isobutyrate and calcium acetate (Munch, *B* 7, 1370, *A* 180, 327)—3 By the action of tin and HClAq on di nitro heptonic acid, which is one of the products of the action of nitric acid on camphor (Kachler, *A* 191, 162)—4 From amylene glycol (CH₃)₂C(OH)CH(OH)CH₃ by dehydration with P₂O₅ (Flavitzky, *B* 10, 2240)—5 From (CH₃)₂CHCH(OH)CH₂(OH) by dehydration with ZnCl₂ or P₂O₅ (F)—6 From (CH₃)₂C(OH)CH(OH)CH₃, by heating with dilute HCl at 100°, and treating the product with potash (Bauer, *C R* 51, 55, *A* 115, 91, Eltekoff, *J R* 14, 358)—7 By shaking (CH₃)₂CHCHCH with diluted H₂SO₄ (SG 1 64) (Flavitzky & Kryloff, *J R* 10, 347)—8 By oxidising MeCH(OH)Pr by CrO₃ (Wino-gradoff, *A* 191, 133)—9 By heating (CH₃)₂CB₂CHBrCH₃ with water and PbO at 150° (Eltekoff, *J R* 10, 215), or by digesting it with water alone (Niederist, *A* 196, 360, Nageli, *B* 16, 2983)—10 By heating MeCH(OH)Pr with excess of dilute (1 p c) H₂SO₄ at 100° (Kondakoff, *J R* 17, 300)

Properties—Liquid Gives the iodoform reaction with iodine and potash Forms a crystal line compound with NaHSO₄ On oxidation it yields acetone and acetic acid, and finally CO₂ and acetic acid

Oxim CH₃C(NOH)CHMe. (158°) From the ketone and hydroxylamine (Nageli, *B* 16, 2984) Formed also by heating the oxim of di methyl acetoacetic acid above 97° (Wallach, *A* 248, 178)

Reference—CHLORO METHYL ISOPROPYL KETONE

Methyl propyl diketone C₈H₁₆O₂ i.e. CH₃COCH₂CH₂CH₂CH₂CH₃ *Acetyl butyryl* (128°) SG 9343 Obtained by boiling this mono oxim with dilute H₂SO₄ (Von Pechmann, *B* 21, 2140) Yellow oil, with irritating odour like quinine

Mono oxim CH₃COCH(NOH)C₄H₉ *Isonitroso propyl acetone* [49 5°] Formed by the action of nitrous acid on propyl acetoacetic ether (Treadwell, *B* 14, 2159) Large plates With phenyl cyanate it reacts with formation of CH₃COCP₂NO CO NHPh [93°] (Goldschmidt, *B* 22, 3108), whence hydroxylamine yields CH₃C(NOH)CP₂NO CO NHPh [129°–131°]

Di-oxim CH₃C(NOH)C(NOH)C₄H₉ *Methyl propyl glyoxim* [168°] Formed by the action of hydroxylamine hydrochloride in aqueous alcoholic solution on isonitroso propyl acetone (Schramm, *B* 16, 2185) Small needles With phenyl cyanate it reacts with formation of CMe(NO CO NHPh)CP₂(NO CO NHPh), crystal lising in pearly plates [164°–170°] (Goldschmidt & Strauss, *B* 22, 3108)

Oxim phenyl hydrazide C₁₁H₁₇N₂O i.e. CMe(N,HPh)CP₂(NOH) [130 5°] Formed by the action of phenyl-hydrazine acetate on the mono oxim (Otte & Pechmann, *B* 22, 2121) Needles, sl sol dilute NaOHAq Gives a violet colour with H₂SO₄ and FeCl₃

(a) *Phenyl-hydrazide* $C_{11}H_{18}N_2O$ α Me C(N,HPH) COPr [114°] Formed by the action of phenyl hydrazine on the diketone (O a P) Insol water and ligroin, v sol alcohol and ether With conc H SO₄ and FeCl it gives a red colour

(B) *Phenyl hydrazide* $C_{11}H_{18}N_2O$ α Me CO C(N,HPH) Pr [109°] Formed from propyl acetoacetic acid by the action of diazo benzene chloride in presence of NaOAc (Japp a Klingemann, A 247, 220) Colourless needles (from benzene and ligroin)

Di phenyl-di-hydrazide $C_{15}H_{22}N_4$ α Me C(N,HPH) C(N,HPH) Pr [136.5°] Formed as a yellow pp when excess of phenyl hydrazine acetate is added to the ketone, its phenyl hydrazide, or its oxim phenyl hydrazide (Otte a Pechmann, B 22, 2121) Yellow needles (from benzene), more easily soluble than its homologues with fewer atoms of carbon in the molecule

Methyl isopropyl diketone $C_8H_{14}O_2$ α CH₃ CO CO CH(CH₃)₂ *Acetyl isobutyryl* (116°) Formed by boiling its oxim with dilute H SO₄ (Von Pechmann, B 21, 2140, 22, 2122) Yellow liquid, with irritating odour, sl sol water

Oxim CH₃ CO C(NOH) CHMe *Isontroso methyl isobutyl ketone* [75°] Formed by the action of nitrous acid on isopropyl acetoacetic ether (Westenberger, B 16, 2991) White plates, sol alcohol, ether, alkalis, and hot water

METHYL PROPYL-KETONE CARBOXYLIC ACID ν Ethyl ACETOACETIC ACID

Methyl-propyl-ketone tricarboxylic acid CH₃ CO CH(CH₃ CO H) CH(CO H) α Carboxy β acetyl glutaric acid [121°-124°] The triethyl ether is formed by the action of chloro or bromo levulic (acetyl propionic) ether upon sodio malonic ether Colourless crystalline solid At 160° it splits off CO₂ giving acetoglutaric acid The neutral NH₄ salt gives pps with AgNO₃, BaCl₂, and Pb(OAc) The zinc salt is a very soluble amorphous solid

Triethyl ether A"Et (285°-295°), oil (Conrad a Guthzeit, B 19, 43)

METHYL - ISOPROPYL - KETONE - CARB OXYLIC ALDEHYDE *Di-oxim* $C_{11}H_{18}N_2O$ α CH₃ C(NOH) CH₂ CMe CH NOH [90°] Formed from di methyl pyrrole and hydroxylamine (Ciamician, B 23, 1788) Small needles or prisms, v sol alcohol, and water, sl sol ether Reduces Fehling's solution Sodium reduces it in alcoholic solution to di methyl tetra methylene-diamine

METHYL PROPYL-DI-KETOXIM ν *Oxim of* METHYL PROPYL DIKETONE

METHYL ISOPROPYL MALONIC ACID $C_8H_{12}O_4$ α CO₂H CMePr CO₂H [124°] Formed by saponifying its ether (Van Romburgh, R T C 5, 236) Crystallises from benzene Split up at 150°-200° into CO₂ and a hexoic acid Salts—CaA', V sol water—AgA' white pp

Ethyl ether EtA" (221°) SG 11 990 Formed from sodio malonic ether, MeI, and PrI Colourless liquid with agreeable odour

METHYL PROPYL OXIDE $C_8H_{16}O$ α CH₃ O Pr (38.9°) SG 2 7471 SV 105.1 CE (0° to 10°) 00146 (Dobner, A 243, 2, cf Chancel, A 151, 305)

METHYL-PROPYL PHENOL ν CARBAORCL, CYMENOL, and THYMOL

Tetrahydride ν BORNEOL and CINEOL

DI METHYL PROPYLPHENYL AMINE

$C_{14}H_{18}(C_6H_5)_2$ NMe *Di methyl phenpropyl amine* (230° uncor) Formed by the action of sodium on a mixture of propyl bromide and di methyl p bromo aniline (Claus a Howitz, B 17, 1327). Colourless oil

Methyl iodide B'MeI [168°], plates

METHYL ISOPROPYL PHENYL CARB-AMINE ν Iso CYMYL CARBAMINE

METHYL PROPYL PHENYLENE DIAMINE

$C_6H_5Me(C_6H_5)(NH)$ Formed by reducing the di oxim of thymoquinone with tin and HCl (Liebermann, B 18, 3193) Oxidised by CrO₃ or FeCl₃ to thymoquinone—B'H₂Cl needles

METHYL ISOPROPYL DIPHENYLENE KE-

STONE $C_{11}H_{16}O$ α CO $\begin{matrix} \diagup C_6H_5 \\ \diagdown C_6H_5 \end{matrix}$ Retene-

ketone [90°] Formed by oxidising retene glycolic acid $C_{10}H_{16}O$ C(OH) CO H (Bamberger a Hooker, A 229, 136) Formed also from retene quinone by the action of alkaline KMnO₄ or by passing it over red hot Ba(OH)₂ or PbO (T. stand, B 17, 692) Yellow prismatic needles or rectangular trimetric plates (by spontaneous evaporation), v sol ligroin, benzene, chloroform, alcohol, and HOAc Volatile with steam and with vapour of alcohol Does not react with hydroxylamine or with NaHSO₃ Reacts with phenyl hydrazine Reduced by sodium amalgam to methyl isopropyl fluorene alcohol (q i), and by red hot zinc dust to methyl isopropyl fluorene

METHYL ISOPROPYL-PHENYL ETHYL

GUANIDINE ν Iso CYMYL ETHYL GUANIDINE

METHYL PROPYL PHENYL GLYOXYLIC

ACID $C_{12}H_{16}O_4$ α [2.5.1] C_6H_5MePr CO CO H Formed by oxidising the ketone $C_8H_{16}O$ CO H, with cold aqueous KMnO₄ (Claus, B 19, 233) Thick oil, easily decomposing into CO₂ and C_6H_5MePr CHO Dilute nitric acid yields $C_6H_5Me(CO_2H)_2$

METHYL PROPYL PHENYL METHYL KE-TONE $C_{11}H_{16}O$ α C_6H_5MePr CO CH₃ (248°) Formed from cymene, AcCl, and AlCl₃ (Claus a Cropp, B 19, 242)

METHYL ISOPROPYL - PHENYL - THIO-

ETHYL-UREA ν Iso CYMYL ETHYL THIO UREA

DI METHYL-DI ISOPROPYL DI-PHENYL-

THIO-UREA ν Di Iso CYMYL THIO UREA

METHYL - ISOPROPYL-PHENYL UREA ν

Iso CYMYL UREA

METHYL - ISOPROPYL - PHENYL - URE-

THANE ν Iso CYMYL CARBAMIC ETHER

METHYL ISOPROPYL-PHOSPHINE $C_8H_{11}P$ α CH₃ PH C₆H₅ (79°) Obtained by heating isopropyl phosphine with MeI at 100° (Hofmann, B 6, 299)

DI-METHYL-DI-PROPYL PYRAZINE

$C_{12}H_{20}N_2$ α N $\begin{matrix} \diagup CMe \\ \diagdown CPr \end{matrix}$ N $\begin{matrix} \diagup CMe \\ \diagdown CPr \end{matrix}$ N *Di-propyl-*

ketone (233°-245°) Prepared by the reduction of the oxim of methyl propyl diketone (nitroso methyl-butyl ketone) with tin and HCl (Treadwell, B 14, 1461, 2160, Oeconomides, B 19, 2526) Oil, with narcotic odour, turning brown in the air—B'H₂PtCl₄ red octahedra,

sl sol water — B'AgNO₃aq crystals, insol cold water

METHYL-PROPYL-PYRIDINES Parvoline

The base obtained by the action of P₂O₅ on a mixture of propionic aldehyde and acetamide (Hesekiel, *B* 18, 3097) has been shown by Durkop & Göttisch (*B* 23, 685) to yield pyridine tricarboxylic acid on oxidation, and must therefore be di methyl ethyl pyridine (*q v*). Parvolines, which must be C₆H₅Me₂N, C₆H₄Me(C₂H₅)N, C₆H₃Me₂EtN, or C₆H₂Et₂N, occur in crude paraffin oil (Williams, *C J* 7, 97), in coal tar (Theniuss, *J* 1861, 502), in the product of the distillation of cinchonine with KOH (Oechsner de Coninck, *Bl* [2] 34, 214), and in putrid flesh (Gautier, *Bl* 48, 11)

Di methyl-propyl pyridine C₁₀H₁₃N *z c* C₆H₂NMePr (133) *n* Propyl lutidine *Conradine* (*c* 195°) at 718 mm Formed by distilling potassium di methyl propyl pyridine dicarboxylate with lime (Jaekle, *A* 216, 37) Colourless liquid, sl sol water, v sol alcohol and ether Its hydrochloride is crystalline — B₂H₂PtCl₆ [185°]

Isomeride *v* CORINDINE

v Methyl-isopropyl pyridine hexahydride

C₆H₁₀N *z c* CH <CH CH>NMe (167°) SG 2 8593 Formed by adding the calculated quantity of MeI to a solution of isopropyl pyridine hexahydride in MeOH, evaporating and distilling the residue with potash (Ladenburg *A* 217, 77) Oil, smelling like conine — B.HAuCl₄ [131°] Plates, sl sol water — B.H.LPtCl₆ [100°] Plates, v sol water and alcohol, insol ether — Picrate B'C₆H(NO)₂OH [149°]

Di-methyl-propyl-pyridine hexahydride

C₆H₁₀N *z c* CHPr <CH CHMe>NH Propyl lupetidine (*c* 180°) at 718 mm Obtained by reducing di methyl propyl pyridine in alcoholic solution with sodium (Jaekle, *A* 246, 46) Colourless oil Gives a brownish pp with FeCl₃, and a yellowish pp with MnSO₄ — B'H₂PtCl₆ [197°] Orange stellate needles

DI METHYL PROPYL PYRIDINE DICARBOXYLIC ACID C₈H₁₀NO₄ *z c*

CPr <C(CO₂H) CMe>N [247°] Formed by saponifying its ether, which is obtained by oxidation, with nitrous gas, from the product of the action of ammonia on butyric aldehyde mixed with acetoacetic ether (Jaekle, *A* 246, 36) Colourless prisms (containing aq) When anhydrous it melts at 247°, but at 212° when hydrated V e sol hot, m sol cold, water Yields di methyl propyl pyridine when distilled with lime *Ethyl ether Et.A* (308°) at 715 mm Light yellow oil Saponified by boiling first with aqueous, and then with alcoholic, potash — (C₆H₁₀NO₄)₂H₂PtCl₆ [187°] Orange prisms *Dihydride of the ethyl ether*

CHPr <C(CO₂Et) CMe>NH [118°] Formed by condensation of butyric aldehyde with acetoacetic ether and ammonia (Jaekle, *A* 246, 34) Yellowish white prisms, v sol alcohol and ether, insol water

Di methyl isopropyl-pyridine dicarboxylic ether. *Dihydride*

CHPr <C(CO₂Et) CMe>NH [97°] Formed by the action of alcoholic ammonia on a mixture of isobutyric aldehyde and acetoacetic ether (Engelmann, *A* 231, 47) Long prisms (from 90 p c alcohol) Oxidised by nitrous acid, in presence of alcohol, to di methyl pyridine dicarboxylic ether C₈H₁₀N(CO Et)₂

METHYL DI ISOPROPYL QUINOLINE DIHYDRIDE C₁₆H₁₇N (299°) Formed by heating di iso propyl indole with MeI and MeOH (Dennstedt, *B* 21, 3437) Oil — B'H₂PtCl₆ [177°]

METHYL PROPYL-STYRIL KETONE

C₁₁H₁₅O *z c* C₆H₄(C₂H₅) CH CH CO CH₃ *Cuminal acetone* (181° at 23 mm) Formed by the action of dilute (10 p c) NaOHAq upon a mixture of cumic aldehyde (20 g), acetone (20 g), water (300 g), and alcohol (170 g) (Claisen & Ponder, *A* 223, 147) Yellow oil

METHYL ISOPROPYL SULPHIDE C₆H₁₀S *z c* MeSPi (93°-95°) VD 45.02 Formed by dissolving sodium in an ethereal solution of isopropyl mercaptan, and slowly adding MeI to the product (Obermeyer, *B* 20, 2923)

METHYL PROPYL DITHIOCARBONATE

MePtCS O Methyl propyl xanthate SG 2 1094 (Na-amn a Scala, *G* 17, 66)

METHYL PROPYL THIO-UREA C₄H₇N₂S *z c* CH₃NH CS NHC₂H₅ [79°] Prepared from methyl thiocarbimide and propylamine, or from propyl thiocarbimide and methylamine (Otto Hecht, *B* 23, 294) Transparent glassy plates (from very dilute alcohol), sl sol cold water, m sol hot water, v sol alcohol and wood spirit, extremely sol acetone and chloroform, v sol ether benzene, and CS, v sl sol warm light petroleum

DI METHYL PYRAZINE C₆H₈N₂ *z c*

N <CMe CH>N *Ketone Di methyl aldine* (Meyer, *B* 21 19) (170°-180°) Formed by reduction of nitroso acetone with tin and HCl (Treadwell & Steiger, *B* 15, 1059, 1055) Oecnomides *B* 19, 2526 Wolff, *B* 20, 433 Formed also by heating its dicarboxylic acid Oil, with alkaloidal odour — B H PtCl₆ golden plates, sol hot water

Tetra methyl pyrazine C₈H₁₂N₂ *z c*

N <CMe CMe>N *Methyl ketone Di methyl aldine Tetra methyl aldine* [86°] (190°) Formed by reducing with tin and HCl the oxim of di methyl diketone (methyl nitroso ethyl ketone) CH₃CO C(NO₂H)CH₃ (Gutknecht, *B* 13, 1116, Treadwell, *B* 14, 1469, Braun & V Meyer, 21, 1947) An intermediate base appears to be C₈H₁₂N (Braun, *B* 22, 556) Tetra methyl pyrazine is also formed, together with CO₂ and other products, by heating CH₃CO CHBr CH₃CO₂H or CH₃CO CH(OH)CH₃CO₂H with conc NH₄Aq (Wolff, *B* 20, 427) Glistening prisms (anhydrous) or long white needles (containing 3aq) Very volatile Melts at 74°-77° when hydrated, but at 86° when anhydrous Strong characteristic smell V sol alcohol and ether By alkaline KMnO₄ it is oxidised to pyrazine tetra carboxylic acid C₈N₂(CO₂H)₄

Salts — B'HCl 2aq easily soluble, [91° anhy] — B'H₂Cl₂PtCl₆ red glistening needles, — B'H₂Cl₄PtCl₆ 4aq orange red needles.

Methylo-iodide $B''MeI$ [216°] anhy Yellow needles (containing 2aq), ν sol water and alcohol, insol ether

Methylo chloride $B''MeCl$ [α 105°], needles— $B''MeCl \cdot HCl \cdot PtCl_4$, aq orange red pyra mids (from hot water)

DI METHYL PYRAZINE DICARBOXYLIC

ACID $C_6H_4N_2O_4$, i.e. $N \langle \begin{smallmatrix} CMe & C(CO_2H) \\ C(CO_2H) & CMe \end{smallmatrix} \rangle N$ [201°] Prepared by saponification of its ether, which is formed by reducing nitroso acetoacetic ether with stannous chloride (Wleügel, *B*, 15, 1050) Formed also by oxidising di methyl di-ethyl pyrazine by aqueous $KMnO_4$, and by condensation of imido oximido butyric ether $CH_3 \cdot C(NH) \cdot C(NOH) \cdot CO_2Et$ by warm $ZnCl_2$ (Oeconomides, *B* 19, 2524)

Properties—Colourless crystals (containing 2aq), melting at 201° when anhydrous ν sol alcohol and acetone, m sol hot water and toluene, nearly insol ether, benzene, and ligroin Decomposes when heated above 200° into CO_2 and di methyl pyrazine

Salts— $BaA''3aq$ glistening crystals— $*KA''$ slender felted needles Gives amorphous pps with $FeCl_3$ and $CuSO_4$ — $Ag A''$ yellowish white pp

Ethyl ether Et_2A'' [86°] (316° cor) Long colourless needles, sol alcohol

METHYL-PYRAZOLONE ν OXY METHYL

PIRAZOLE

(a)-METHYL PYRIDINE C_6H_7N i.e.

$N \langle \begin{smallmatrix} CMe & CH \\ CH & CH \end{smallmatrix} \rangle CH$ (a) *Picoline* Mol w 93 (129°) (L), (133 4° cor) (Thorpe, *C J* 37, 223) $SG \frac{1}{2}$ 9656 (L), 9616 (T) CE (0°-10°) 000975, (0°-100°) 0010968 (T) SV 111 5

Occurrence—In bone-oil (Anderson, *A* 60, 86, Weidel, *B* 12, 2008), where it is mixed with a little (β) picoline and di methyl pyridine (Ladenburg, *B* 18, 49) It is also the chief constituent of coal tar picoline (Goldschmidt & Constam, *B* 16, 2976)

Formation—1 By heating pyridine (a) carb oxylic acid with fuming $HIAq$ at 260°, or by treating the acid with zinc and $HOAc$ (Seyfferth, *J pr* [2] 34, 244)—2 By the action of aldehyde on aldehyde ammonia (Dürkopf & Schlangk, *B* 21, 297)—3 By heating pyridine methylo iodide in sealed tubes at 300° (Ladenburg, *A* 247, 7, *C R* 103, 692)

Preparation—Crude pyridine (50 g boiling between 128° and 134°) is dissolved in $HClAq$ (170 g of 11 p.c.) and a hot solution of $HgCl_2$ (312 g) in water (4½ litres) is poured in The double salt crystallises out, and, after recrystallisation, is decomposed by aqueous $NaOH$ (Ladenburg, *A* 247, 6)

Properties—Liquid, inactive to light, miscible with water and alcohol Picoline lessens the heart's action, and acts on the nervous system somewhat like nicotine (Oechsner, *Bl* [2] 38, 547)

Reactions—1 Yields pyridine (a) carboxylic (picoline) acid on oxidation (Weidel)—2 When heated with sodium it yields di methyl dipyrindyl 3 *Methylal* and $ZnCl_2$ yield $CH_2(C_6H_7N)_2$ —4 *Chloral* forms $C_6H_7N \cdot CH_2 \cdot CH(OH) \cdot CCl_3$ [87°], the hydrochloride of which $C_6H_7N \cdot OCl \cdot HCl$ melts at 202° Alcoholic potash converts it into pyridyl acrylic acid (Einhorn & Liebrecht, *B* 20,

1592)—5 *Paraldehyde* at 260° forms allyl pyridine $C_6H_7N \cdot C_3H_5$ (190°), $SG \frac{1}{2}$ 9595 (Ladenburg, *A* 247, 26) Its aurochloride melts at 136°, its platinumchloride at 186° Allyl pyridine is reduced in alcoholic solution by sodium to conine—6 *Furfuraldehyde* and a little $ZnCl_2$ at 170° yields $C_6H_7N \cdot CH \cdot CH \cdot C_4H_3O$ [58°] (Merck, *B* 21, 2709), ν *Furfuryl vinyl pyridine* In alcoholic solution it is reduced by sodium to the hexahydride of furfuryl ethyl pyridine ($q \nu$) 7 *Glycolic chlorhydrin* at 140° forms a compound $C_6H_7N \cdot NO$, which yields the salts $B''H \cdot PtCl_4$ [200°] and $B''H \cdot AuCl_3$ [100°] (Alexander, *B* 23, 2714)

Salts— $B''H \cdot PtCl_4$, aq [178°] Monoclinic tables or prisms, sl sol water— $B''H \cdot PtCl_4 \cdot 2aq$ (Weidel)— $B''H \cdot PtCl_4$ [195°] Monoclinic plates, abc c=6636 1 9078, β =72° 46' (Stohr, *J pr* [2] 42, 420)— $B''H \cdot AuCl_3$ [168°] Needles, sl sol water— $B''HHg \cdot Cl_2$ [164°] Prisms (from dilute $HClAq$), ν sl sol cold, ν sol hot, water— $B'' \cdot ZnCl_2$ (at 120°) Crystals (from alcohol) (Lachovitch & Bandrowsky, *M* 9, 517) Picrate $B''C_6H_4(NO_2)_3OH$ [165°] M sol water (Lange, *B* 18, 3436)

(a) **Methyl pyridine tetrahydride** $C_6H_{11}N$ i.e. $NH \langle \begin{smallmatrix} CMe & CH \\ CH_2 & CH_2 \end{smallmatrix} \rangle CH$ Formed by the action of alcoholic NH_3 upon methyl ω bromo butyl ketone, probably by elimination of water from the intermediate $CH_3 \cdot CO \cdot CH_2 \cdot CH_2 \cdot CH_2 \cdot NH_2$ (Lipp, *B* 19, 284)

(a)-Methyl pyridine hexahydride $C_6H_{13}N$ i.e. $NH \langle \begin{smallmatrix} CMe & CH \\ CH_2 & CH_2 \end{smallmatrix} \rangle CH$ (a)-Methyl piperidine

(a) *Piperidine* (119°) $SG \frac{1}{2}$ 8600 Obtained by reducing (a) methyl pyridine in alcoholic solution with sodium (Ladenburg & Roth, *B* 18, 47, *A* 247, 62, *C R* 103, 747) Colourless liquid, smelling like piperidine, ν sol water, alcohol, and ether Separated by KOH from its aqueous solution Inactive, but if a crystal of hydrogen conine tartrate be added to a conc solution of the acid tartrate, crystals of the acid tartrate of dextrorotatory (a) methyl pyridine hexahydride separate, while the salt of the laevo rotatory isomeride remains as an oil The rotation of the dextro (a) methyl piperidine is $[\alpha]_D^{20}$ =21 44', while that of the laevo base is (probably) equal and opposite

Reactions—1 Treatment with Br and $NaOHAq$ yields (a) piperolein C_6H_7N , an oily base (126°), $SG \frac{1}{2}$ 880 (Ladenburg, *B* 20, 1645) Its acetyl derivative boils at about 230°—2 Combines with sulphide of carbon, forming methyl piperidine methyl piperyl di thio carbamate $C_6H_{13}N \cdot CS \cdot SC_6H_7N$, [119°], ν e sol water and alcohol

Salts— $B''HCl$ [189°] Colourless needles, ν sol water Not deliquescent— $B''HBr$ [182°] Silky matted needles, m sol water—The platinumchloride forms sparingly soluble plates

(β)-Methyl-pyridine C_6H_7N i.e. $N \langle \begin{smallmatrix} CH & CMe \\ CH & CH \end{smallmatrix} \rangle CH$ m *Picoline* (β) *Picoline* (144° cor) $SG \frac{1}{2}$ 9771 (L), 9765 (B)

Occurrence—In bone oil (Weidel, *B* 12, 2008) and in coal tar (Möhler, *B* 21, 1009)

Formation—1 By distillation of acrolein-ammonia (Bayer, *A* 155, 288, cf Claus, *A*,

Suppl 2, 134, 180, 185, 158, 222) —2 By heating CH_3Br CHBr CH_3Br with alcoholic ammonia at 250° (Baeyer) —3 By distilling strychnine with lime (Stoehr, *B* 20, 2728, Löbisch & Malfatti, *M* 9, 632) —4 By heating acetamide (10 g) with glycerin (32 g) and P_2O_5 (26 g) (Zanoni, *J* 1882, 498, Heseckiel, *B* 18, 3091) —5 Together with homologues, by distilling glycerin with $(\text{NH}_4)_2\text{SO}_4$ and a little H_2SO_4 (Storch, *B* 19, 2458) —6 One of the bases got by distilling brucine with lime (Behrend, *J pr* [2] 42, 415) —7 Together with tri methylene imine (66° – 70°) by distilling tri-methylene diamine hydrochloride (Ladenburg & Sieber, *B* 23, 2729)

Purification —By digesting in hydrochloric acid solution with NaNO_2 on a water bath, followed by crystallisation of its mercury double salt (Bachér, *B* 21, 293)

Properties —Liquid When prepared from strychnine it boils at 149° , and the base so obtained (called (β) methyl pyridine) is less soluble in water than the variety boiling at 144° , and forms a platinumchloride melting at 258° instead of 241° (Ladenburg, *B* 23, 2688) Optically inactive (Landolt, *B* 19 157) Less soluble in water than (α) picoline The absorption spectrum has been studied by Hartley (*C J* 41, 45) Oxidised by a 2 p.c. solution of KMnO_4 to pyridine (β) carboxylic (nicotinic acid)

Salts — $\text{B}' \text{H}_2\text{PtCl}_6\text{aq}$ Monoclinic prisms, v sol hot water Melts, when anhydrous, at 191° (L) or 195° (Stoehr) On heating at 120° for some time it gives off HCl , leaving $\text{B}'_2\text{HPtCl}_6$ [214°] A boiling aqueous solution deposits $\text{B}'_2\text{PtCl}_6$, while $\text{B}'_2\text{HPtCl}_6$ crystallises from the nitrate $\text{B}'\text{HAuCl}_4$ [184°] Needles (from hot water), v sol alcohol — $\text{B}'\text{HHg}_2\text{Cl}_4$ [143°] Needles (from water), plates or needles (from HClAq) or prisms (on slow crystallisation), v sol water, more sol HClAq — $\text{B}'_2\text{HgCl}_2$ white pp — $\text{B}'_2\text{H}_2\text{ZnCl}_4$ [158°] Pearly needles, v sol hot water —Picrate $\text{B}'_2\text{C}_6\text{H}_2(\text{NO}_2)_4\text{OH}$ [145°] Needles or plates, m sol water and alcohol

(β) *Methyl-pyridine hexahydrate* $\text{C}_6\text{H}_9\text{N} \cdot 6$
 $\text{NH} \begin{smallmatrix} \text{CH} & \text{CHMe} \\ \text{CH} & \text{CH}_2 \end{smallmatrix} > \text{CH}_2$ (β) *Methyl piperidine*

(β) *Pipecoline* (β) *Picoline hexahydrate* (125°) SG $\frac{2}{3}$ 8684 VD ($\text{H}=1$) 98 14 Obtained by reducing (β) picoline in alcoholic solution with sodium (Ladenburg, *A* 247, 67, Stoehr, *B* 20, 2732, Heseckiel, *B* 18, 910) Colourless liquid, smelling like piperidine, v sol water When heated with MeI it forms $\text{C}_6\text{H}_9\text{NMeI}$ [192°] The hydrochloride of (β) methyl-pyridine hexahydrate is not pptd by HgCl_2 .

Salts — $\text{B}'\text{HCl}$ Colourless needles, v e sol water and alcohol — $\text{B}'\text{HI}$ [131°] Colourless, non deliquescent needles — $\text{B}'_2\text{H}_2\text{PtCl}_6$ [192°] Orange yellow prisms, m sol water — $\text{B}'\text{HAuCl}_4$ [131°] M sol water — $\text{B}'_2\text{H}_2\text{CdI}_4\text{aq}$ White plates, melting at 145° when anhydrous $\text{B}'_2\text{H}_2\text{FeCy}_2\text{aq}$ yellow monoclinic prisms, less soluble than the ferrocyanides of homologous bases Decomposed by water at 75° —Picrate $\text{B}'_2\text{C}_6\text{H}_2(\text{NO}_2)_4\text{OH}$ [138°] Yellow pp, m sol water

(γ) *Methyl-pyridine* $\text{C}_6\text{H}_9\text{N} \cdot 3$
 $\text{N} \begin{smallmatrix} \text{CH} & \text{CH} \\ \text{CH} & \text{CH} \end{smallmatrix} > \text{CMe}$ *p Picoline* (γ) *Picoline*.
 (148 5° cor) SG $\frac{2}{3}$ 9742

Occurrence —In coal tar, from which it may be obtained by preparing its platinumchloride from the so-called lutidine (Schulze, *B* 20, 418, Ladenburg, *B* 21, 285, *A* 247, 11)

Formation —1 By heating di-chloro-pyridine (γ) carboxylic acid with conc HIAq and P at 175° (Behrmann & Hofmann, *B* 17, 2696) —2 In small quantity by heating pyridine methylo-iodide in sealed tubes at 300° , distilling the product with KOH , converting the fraction (142° – 146°) into platinumchloride, and decomposing the Pt salt by H_2S (L) —3 By distilling sparteine with lime (Ahrens, *B* 21, 828)

Properties —Oil, smelling like (α) methyl-pyridine, v sol water, alcohol, and ether Oxidised by dilute KMnO_4 to pyridine (γ) carboxylic (isomonic acid) [307°]

Salts — $\text{B}'_2\text{H}_2\text{PtCl}_6$ [231°] Four sided plates, sl sol cold water — $\text{B}'\text{HAuCl}_4$ [205°] Prisms, v sl sol water — $\text{B}'_2\text{HHg}_2\text{Cl}_4$ [129°] Needles, v sol hot, sl sol cold, water —Picrate $\text{B}'_2\text{C}_6\text{H}_2(\text{NO}_2)_4\text{OH}$ [167°] Tufts of needles, sl sol cold water

(γ) *Methyl-pyridine dihydride* $\text{C}_6\text{H}_7\text{N} \cdot 2$
 $\text{NH} \begin{smallmatrix} \text{CH} & \text{CH} \\ \text{CH} & \text{CH} \end{smallmatrix} > \text{CHMe}$ Formed from ethyl-pyrrole and HClAq at 130° (Dennstedt & Zimmermann, *B* 19, 2197) — $\text{B}'_2\text{H}_2\text{PtCl}_6$, red needles, v sol water

(γ) *Methyl-pyridine hexahydrate* $\text{C}_6\text{H}_9\text{N} \cdot 6$
 $\text{NH} \begin{smallmatrix} \text{CH}_2 & \text{CH}_2 \\ \text{CH}_2 & \text{CH}_2 \end{smallmatrix} > \text{CHMe}$ (γ) *Pipecoline* (126 5° – 129° cor) SG $\frac{2}{3}$ 8674 Obtained by reducing (γ) methyl pyridine in alcoholic solution by sodium (Ladenburg, *B* 21, 288, *A* 247, 69) Colourless hygroscopic liquid which fumes in the air, and smells like piperidine V sol water

Salts — $\text{B}'\text{HCl}$ v e sol water — $\text{B}'_2\text{H}_2\text{PtCl}_6$ [203°] Prisms, m sol water — $\text{B}'\text{HAuCl}_4$ [127°] Yellow needles sl sol water — $\text{B}'_2\text{H}_2\text{CdI}_4$ [135°] —Bismutho iodide characteristic red plates The picrate and mercury double chloride are crystalline

Methyl-pyridine dihydride $\text{C}_6\text{H}_7\text{N} \cdot 2$
 $\text{NMe} \begin{smallmatrix} \text{CH} & \text{CH} \\ \text{CH} & \text{CH} \end{smallmatrix} > \text{CH}_2$ (?) (129°) Obtained by distilling pyridine methylo iodide with KOH (2 pts) and a little water (Hofmann, *B* 14, 1498) Very pungent oil, almost insol water Absorbs oxygen from the air Combines with bromine, with iodine and with sulphur Combines with CS and with mercaptan Conc HClAq at 180° decomposes it, giving off methylamine

Methyl pyridine hexahydrate $\text{C}_6\text{H}_9\text{N} \cdot 6$
 $\text{NMe} \begin{smallmatrix} \text{CH}_2 & \text{CH}_2 \\ \text{CH}_2 & \text{CH}_2 \end{smallmatrix} > \text{CH}_2$ *Methyl piperidine*.
 (105°) (L) Formed by adding potash to the product of spontaneous union of piperidine with MeI (Cahours, *A Ch* [3] 38, 76) Formed also by heating piperidine hydrochloride (10 g) with MeOH (75 g) for 4 hours at 300° , excess of alcohol being distilled off, and the dry residue distilled with aqueous potash (Ladenburg, *A* 247, 56) Liquid — $\text{B}'\text{HCl}$ Needles — $\text{B}'_2\text{H}_2\text{PtCl}_6$ Orange crystals, v sol water

Methylo-iodide $\text{B}'\text{MeI}$ Crystalline, m sol hot alcohol When heated with solid KOH it yields only 'di-methyl-piperidine' (118°), a strongly alkaline base, which is decomposed by gaseous HCl into methyl chloride and methyl-

piperidine 'Di methyl piperidine' forms the salts C_4H_9NHCl and C_4H_9NHAc . It combines with halogens, forming crystalline C_4H_9NI , C_4H_9NCl (which yields $C_4H_9NCl \cdot AlCl_3$), and C_4H_9NBr . The latter is converted by moist Ag_2O into crystalline C_4H_9NBr , whence further treatment with moist Ag_2O produces C_4H_9NOH which is split up on distillation into H_2O and 'di methyl piperidine'. Ladenburg suggests the formula $CH_3CHCHCHCH_2NMe_2$ for 'di methyl piperidine', and Merling (*B* 19, 2628)

suggests $CH_3CH(CH_3)CHCH_2NMe_2$ for the compound $C_4H_9NBr_2$. This dibromide is accompanied by an oily isomeride which may possibly be $CH_3BrCHBrCH_2CH_2NMe_2$, which changes into the other variety on heating its alcoholic solution. The diiodide $C_4H_9NI_2$ is converted, on treatment with Ag_2O into 'di methyl piperidine' C_4H_9N , a liquid (137°–140°) which forms the salts $(C_4H_9N)H_2PtCl_6$ and $(C_4H_9N)HAuCl_4$, and a methylo iodide C_4H_9NMeI . 'Di methyl piperidine' combines with MeI forming crystalline C_4H_9NMeI [200°] whence moist Ag_2O forms strongly alkaline C_4H_9NMeOH which is split up by heat into pyridine C_4H_5N , water, NMe_3 , $MeOH$, and di-methyl piperidine. Di methyl piperidine also combines with methylene iodide forming $C_4H_9NICH_2I_2$ (Ladenburg, *B* 14, 1347). Ladenburg suggests for di methyl piperidine the formula $CH_3CH(CH_3)CHCH_2NMe_2$

Methyl pyridine from Bone oil The following salts described by Ramsay (*P M* Oct 1876 and 1877, July 1878) were prepared from a methyl pyridine obtained from Dippel's oil, which was probably (a) methyl pyridine mixed with a little (b) methyl pyridine — * $B \cdot HCl$ [160°] White deliquescent crystals — $B \cdot HBr$ [187°] Deliquescent Absorbs bromine forming $B \cdot HBr$, golden-yellow needles, sol water — $B \cdot Br$. Needles — * $B \cdot HI$ Decomposed by heat, yielding $B \cdot HI$, which crystallises in brown needles [79°] — $B \cdot ICl$ yellow tables — $B \cdot HgCl$ — $B \cdot H_2PtCl_6$ (at 100°) Converted by water at 170° into yellow flocculent $B \cdot PtCl_6$ and $B \cdot PtCl_4$, a yellowish green insoluble powder — $Platino-cyanide B \cdot H_2PtCy_4$ 4aq yellow crystals The crude bone oil picoline forms the following combinations with alkyl salts — $B \cdot MeCl$ deliquescent needles (from alcohol) — $(B \cdot MeCl)_2PtCl_6$ small cubes — $B \cdot MeI$ [227°] Long white deliquescent needles (from alcohol) — $B \cdot MeI_3$ [129°] Blush black feathery plates, insol water and CS_2 , sol alcohol and ether — $B \cdot MeNO_3$ transparent prisms — $(B \cdot C_2H_5Cl)_2PtCl_6$ — $B \cdot C_2H_5Br$ [276°] Small prisms The same picoline forms with acetyl chloride deliquescent brown crystals of $B \cdot AcCl$ According to Gardner (*B* 23, 1589) crude picoline forms with acetic and formic acids the salts $(C_4H_5N)(HOAc)$, (c 148°) and $C_4H_5N(CO_2H_2)$, (156°–159°) which distil unchanged

Di methyl-di-pyridine ($C_8H_9N_2$), *Dipicoline Parapicoline* (310°–320°) SG 112 H F p 8084 (Ramsay, that for picoline being 3759) Formed by boiling bone oil picoline (6 pts) with sodium (1 pt) for two days (Anderson, *A* 105, 844). Pale-yellow oil miscible with alcohol and ether. Bromine water gives a pp of

$C_{12}H_{11}BrN_2H_2Br_2$ — $C_{12}H_{11}N_2H_2PtCl_6$ pale yellow powder Its aurochloride is decomposed by boiling water (O de Coninck, *B* [2] 45, 131)

Methylo iodide $C_4H_9N(Me)I$ Yellow powder, v sol water, almost insol alcohol and ether Yields $C_4H_9NMe_2PtCl_6$ and $C_4H_9NMe_2I_3$

(aa) **Di methyl pyridine** C_4H_5N : e
 $N \begin{smallmatrix} \text{CMe} \text{CH} \\ \text{CMe} \text{CH} \end{smallmatrix} \text{CH}$ (aa) *Lutidine* oo *Lutidine*
 Mol w 107 (142° cor) (L), 145° (E) SG
 § 942

Occurrence — In coal tar, being obtained by extracting with H_2SO_4 ppq by alkali, and separating from the isomeride (157°) by fractionating (Lunge a Rosenberg, *B* 20, 127, Ladenburg a Roth, *A* 247, 28) It occurs also in bone oil, and may be obtained from the fraction (135°–145°) (Roth, *B* 19, 786)

Formation — 1 By the action of cinnamic aldehyde and alcoholic NH_3 on acetoacetic ether, the resulting dihydride of styryl di methyl pyridine dicarboxylic ether being saponified, oxidised by $KMnO_4$, and the di methyl pyridine tetracarboxylic acid so produced distilled with lime in a current of hydrogen (Epstein, *A* 231 18) — 2 By distilling its dicarboxylic acid with lime (Engelmann, *A* 231, 54) — 3 By distilling oxy di methyl pyridine (lutidine) with zinc dust (Courad a Epstein, *B* 20, 162)

Preparation — Claude picoline (139°–142°) from bone oil is dissolved in excess of $HClAq$ and a hot solution of $HgCl_2$ is added The double salt which is ppd is recrystallised and decomposed by aqueous $NaOH$ The base is finally separated by solid KOH (Ladenburg *A* 247, 30)

Properties — Liquid smelling and tasting like pyridine, sol cold water, the base separating again on warming Its aqueous solution ppts solutions of $ZnSO_4$, $CdSO_4$, $FeSO_4$, and $FeCl_3$ With $CuSO_4$ it gives a pale blue pp not turned black by heating With $AgNO_3$ it gives minute needles of $B \cdot AgNO_3$ Dilute $KMnO_4$ oxidises it to pyridine dicarboxylic acid [227°]

Salts — * $B \cdot HCl$ Deliquescent needles — $B \cdot H_2PtCl_6$ [208°] Orange red monoclinic plates, $a b c = 892.1 \ 660. \ 81^\circ 55'$ v sol hot, m sol cold, water, insol alcohol — $B \cdot HAuCl_4$ [124°] Yellow needles (from very dilute HCl) — $B \cdot HHgCl_2$ [186°] Thin plates (from acidulated water) — $B \cdot HHgCl_2$ (Mohler, *B* 21, 1008) — $B \cdot H_2CrO_4$ [92°] Orange prisms — $B \cdot (NH_4)_2H_2O_4$ [c 160°] — *Picrate* [159°] Yellow needles or thin plates

(aa) **Di methyl pyridine hexahydride**
 $C_8H_9N_2$: e $NH \begin{smallmatrix} \text{CHMe} \text{CH}_2 \\ \text{CHMe} \text{CH}_2 \end{smallmatrix} CH_2$ (127°–130°) SG § 8492 Formed by reducing the corresponding di methyl pyridine in alcoholic solution with sodium (Ladenburg, *A* 247, 87 *B* 18, 54) Colourless liquid, miscible with water, alcohol, and ether — $B \cdot HCl$ non deliquescent needles, m sol water — $B \cdot HBr$ Needles, v sol water — $B \cdot H_2PtCl_6$ [212°], orange red crystals

(ay) **Dimethyl pyridine** C_6H_7N : e
 $N \begin{smallmatrix} \text{CMe} \text{CH} \\ \text{CH} \text{CH} \end{smallmatrix} \text{CMe}$ *Lutidine* (157°) SG
 § 9493 S 20

Occurrence—In coal tar oil, being extracted with other bases by H_2SO_4 , (Ladenburg a Roth, *B* 18, 918, Lunge a Rosenberg, *B* 20, 131), Ladenburg, 21, 286)

Formation—1 By distilling oxy di methyl pyridine $CMe \begin{smallmatrix} \text{CH CO} \\ \text{CH CMe} \end{smallmatrix} > NH$ with zinc dust (Hantzsch, *B* 17, 2908)—2 By distilling its tri-carboxylic acid with lime (Hantzsch, *A* 215, 56)—3 By distilling with lime the acid $N \begin{smallmatrix} \text{CMe C(CO}_2\text{H)} \\ \text{CH CH} \end{smallmatrix} > CMe$ the ether of which is obtained by condensation of an equal number of molecules of acetoacetic ether, acetic aldehyde, and acetic aldehyde ammonia (Michael, *B* 18, 2020)

Preparation—The fraction of coal tar bases boiling between 155° and 160° is dissolved in dilute $HClAq$, conc $HClAq$ is then added, followed by a hot concentrated solution of $HgCl_2$. The double salt which then crystallises out is decomposed by distilling with $NaOHAq$ (Ladenburg, *A* 247, 35)

Properties—Liquid, smelling like pyridine, dissolving in 5 pts of cold water, less soluble in hot water. Miscible with alcohol and ether. Easily volatile with steam. Oxidised by $KMnO_4$ to pyridine dicarboxylic (lutidine) acid [235°]. Reacts with benzoaldehyde and $ZnCl_2$ forming styryl methyl pyridine (Bachér, *B* 21, 3071)

Salts— $B'HgCl_2$, aq Needles. Melts at 132° when anhydrous. Mohler (*B* 21, 1008) obtained from coal tar lutidine in acid solution a salt $B'HHgCl_2 \cdot B'H.PtCl_4$ [220°]. Plates or prisms— $B'HAuCl_4$, amorphous pp, changing to prisms— $*B'HCl$ slender needles— $*B'HBr$ Needles—Picrate [179°] Needles, sl sol cold water

($\alpha\gamma$) Di-methyl-pyridine hexahydrate

$NH \begin{smallmatrix} \text{CHMe CH} \\ \text{CH}_2, \text{CH}_2 \end{smallmatrix} > CHMe$ ($\alpha\gamma$) Di methyl pyridine (141°) SG 2 8615 Obtained by reducing the corresponding di methyl pyridine in alcoholic solution with sodium (Ladenburg, *A* 217, 88). Colourless strongly alkaline liquid, smelling like piperidine. Fumes with HCl . M sol water, v e sol alcohol and ether

Salts— $B'HCl$ [235°] Long colourless needles, v sol water— $B'HBr$ [142°] Short needles, v e sol water— $B'_2H.PtCl_4$ groups of yellow needles

($\beta\delta$) Di-methyl-pyridine C_4H_8N &c

$N \begin{smallmatrix} \text{CH CMe} \\ \text{CH CMe} \end{smallmatrix} > CH$ (170°) SG 2 9614 Obtained by heating with lime the di methyl pyridine carboxylic acid formed by oxidation of the ($\beta\delta$) di methyl-ethyl pyridine produced by heating propionic aldehyde ammonia with propionic aldehyde (Dunkopf a Götsch, *B* 23, 1113). Transparent, strongly refracting liquid, with a pleasant odour characteristic of (β) alkyl pyridines, m sol cold water, sl sol hot water. Yields on oxidation a pyridine dicarboxylic acid [315°]

Salts— $B'_2H_2.PtCl_4$ [256°] Dark red needles and plates, sl sol water— $B'HAuCl_4$ [149°] Yellow needles, sl sol water—Mercury double chloride [176°] Long needles, sl sol cold water

($\alpha\delta$)-Di-methyl-pyridine $N \begin{smallmatrix} \text{CMe CH} \\ \text{CH CMe} \end{smallmatrix} > OH$

(162° – 166°) Occurs in coal tar (Lunge a Rosenberg, *B* 20, 134) Yields isomochomeronic (pyridine dicarboxylic) acid on oxidation

($\alpha\gamma$)-Di-methyl-pyridine tetrahydride?

C_4H_8N &c $NMe \begin{smallmatrix} \text{CHMe CH}_2 \\ \text{CH=CH} \end{smallmatrix} > CH_2$ Formed, together with $MeCl$, hydrocarbons, methylamine, NH_3 , and other bases, by heating anhydro ecgonine with conc $HClAq$ (Einhorn, *B* 22, 1362). Oil. When heated with conc $HClAq$ at 280° it yields a mixture of bases which appear to yield methyl pyridine when distilled over zinc dust— $B'HAuCl_4$ [212°] Small needles, m sol water— $B'_2C_2H_5(NO_3)_2.OH$ Long needles, v sl sol hot water— $*B'HCl$ very hygroscopic

Di-methyl-pyridine of bone oil (156° – 159°) is a mixture of ($\alpha\gamma$) di methyl pyridine, ($\alpha\delta$)-di methyl-pyridine, (β)-ethyl pyridine, and (γ)-ethyl pyridine. The existence of these bases is shown by the production of the corresponding pyridine di and mono carboxylic acids on oxidation by $KMnO_4$ (Weidel a Herzig, *M* 1, 1, Weidel a Pick, *M* 5, 658, cf Anderson, *A* 80, 5). Bone oil also contains ($\alpha\alpha$) di methyl pyridine (*v supra*) Greville Williams (*C J* 7, 97, *P* 13, 311) obtained a lutidine from coal tar, this has since been shown to contain ($\alpha\alpha$), ($\alpha\gamma$), and ($\alpha\delta$) di methyl pyridines (*v supra*) Oechsner de Coninck (*Bl* [2] 41, 249) found (γ) ethyl pyridine (154°) in coal tar. Lutidines have also been obtained by distilling the bituminous shale of Dorsetshire (Williams) and peat (Church a Owen, *P M* [4] 20, 110). Among the products obtained by distilling cinchonine with KOH Oechsner de Coninck (*C R* 91, 296) obtained a lutidine (165°), VD 38 (calc 3.7), SG 2 959, which formed a deliquescent crystalline hydrochloride and a platinochloride $B'_2H_2.PtCl_4$ crystallising in orange red needles, converted by boiling water into $B'_2H.PtCl_4$ crystallising in yellow needles. When brucine is distilled with KOH it yields a lutidine (166°) whence $B'_2H.PtCl_4$ [180°] and $B'.PtCl_4$ [205°] (Oechsner de Coninck, *C R* 95, 298, 96, 437). Lutidine aureochloride $B'HAuCl_4$ is decomposed on boiling with water, yielding first thin red plates of $B'HAuCl_4 \cdot B'_2AuCl_4$, and then a red crystalline pp B'_2AuCl_4 (O de Coninck, *Bl* [2] 31, 634)

Di-methyl pyridine dihydride C_4H_8N (199°) VD 33 Occurs in cod liver oil (Gautier a Mourgues, *C R* 107, 111, *Bl* [3] 2, 213). Colourless strongly alkaline and caustic oil, absorbs CO_2 from the air. Poisonous. Its salts taste bitter. Oxidised by boiling aqueous $KMnO_4$ to methyl pyridine carboxylic acid and a little pyridine carboxylic acid

Salts— $*B'HCl$ confused needles, v sol water— $*B'HN_3$ Reduces $AgNO_3$ — $B'_2H_2SO_4$ Groups of deliquescent needles— $B'_2H.PtCl_4$ Lozenge shaped plates, loses HCl on boiling with water

Methyl iodide $B'MeI$ Colourless needles, sol water and alcohol. Yields a tri-methyl-pyridine dihydride on treatment with potash

Di-methyl-pyridine hexahydrate **Methyl iodide** C_4H_8NMeI [192°] Formed by heating (β) pyridine hexahydrate with MeI and $MeOH$ at 100° (Hessekel, *B* 18, 8099, *A* 247, 69). Needles (from acetone). Not decomposed by aqueous KOH . Yields $(C_4H_8NMeCl).PtCl_4$, an orange crystalline pp turned black at 234°

(e) Tri-methyl-pyridine $C_6H_3N \cdot 3e$

$N \begin{smallmatrix} \text{CMe CH} \\ \text{CMe CH} \end{smallmatrix} \text{CMe}$ (γ)-Collidine (172° cor) (H, M); [168°] (D) SG λ 917 (H), 922 (M), τ 9312 (D) Occurs in coal tar, from which it may be obtained by fractional distillation followed by ppn of the bases by K_2FeC_4 (Mohler, B 21, 1011) Formed by heating with quicklime the potassium salt of its dicarboxylic acid, which is obtained by the action of nitrous acid on its dihydride produced from acetoacetic ether and aldehyde ammonia (Hantzsch, A 215, 32) Formed also by heating acetone with NH_4Cl for 3 days at 265°, CH_4 being evolved (Riehm, A 238, 16), and by heating acetone with aldehyde ammonia for 10 hours at 200° (Dürkopf, B 21, 2713)

Properties—Liquid which turns brown in the air More than 3 times as soluble in water as aldehyde collidine Less soluble in hot than in cold water It differs also from aldehyde collidine in giving a pp with $AgNO_3$, an orange crystalline pp with CrO_3 , and an aurochloride that melts under water It is oxidised by $KMnO_4$ to $N \begin{smallmatrix} \text{C(CH}_3)_2 \text{ CH} \\ \text{C(CO}_2H)_2 \text{ CH} \end{smallmatrix} \text{CMe}$ and uvitonic acid Bromine added to its solution in CS forms unstable orange crystals of $C_6H_3NBr_2$ (Pfeiffer, B 20, 1844)

Salts— B^+HCl Slender, non deliquescent needles— $B^+H_2PtCl_6$ yellow crystalline pp— B^+HAuCl_4 [113°] (H), [115°] (M), [106°] (D) Needles (from hot water)— B^+HHgCl_2 [155°]— B^+HI — B^+TNO , [above 300°]— $B^+H_2CrO_4$ Yellow prisms, decomposing at 190°— $B^+H_2SO_4$ [203°]—Picrate [156°] (M) Silky yellow needles, sl sol water, v sol alcohol

Reference—DI BROMO TRI METHYL PYRIDINE

Tri methyl pyridine dihydride $C_8H_{12}N$ Di-hydro collidine (175°–180°) Formed by heating its dicarboxylic ether (obtained from aldehyde ammonia and acetoacetic ether) with dilute HCl aq at 130° (Hantzsch, A 215, 44) Pungent oil with alkaloidal smell Alkaline to litmus, v sol cold water Precipitates the hydrides of Mg, Zn, and Fe from solutions of their salts— $B^+H_2PtCl_6$ Minute needles which blacken at 200°— B^+HI

Polymeride $C_6H_2N_2$ Tetrahydriodicollidine (255°–260°) Formed at the same time as the preceding— $C_6H_2N_2H PtCl_6$ — $C_6H_2N_2HI$

s -Tri-methyl-pyridine hexahydride $C_8H_{11}N$ νe $NH \begin{smallmatrix} \text{CHMe CH}_2 \\ \text{CHMe CH} \end{smallmatrix} CHMe$ Copellidine (146°) SG τ 8475 Formed by reducing s tri methyl pyridine in alcoholic solution with sodium (Jaacke, A 246, 43) Formed also, together with s tri methyl pyridine, by heating acetone with aldehyde ammonia at 200° (Dürkopf, B 21, 2715) Liquid, smelling like piperidine, sl sol water, miscible with alcohol and ether Unlike s tri methyl-pyridine it gives a brownish-black pp with $Hg_2(NO_3)_2$. Gives no pps with $HgCl_2$ or picric acid

Salts— B^+HCl Needles or prisms, v e sol. water and alcohol— B^+HBr — $B^+H_2PtCl_6$ [205°] (J); [244°] (D)

Isomerides of tri-methyl-pyridine ν METHYL-ETHYL-PYRIDINE, where (α)- and (β) collidines,

aldehyde-collidine and other isomerides are described

A collidine dihydride $C_8H_{11}N$, (210°), SG τ 1029, is contained in putrid horseflesh and putrid beef, and forms a crystalline hydrochloride and platinocchloride (Gautier, Bl [2] 48, 12)

Tetra methyl pyridine dihydride $C_8H_{11}N \cdot 2e$

$NH \begin{smallmatrix} \text{CMe CMe} \\ \text{CH}_2 \text{ CMe} \end{smallmatrix} \text{CMe}$ Dihydroparvoline

(159°) Formed by heating potassium pyrrole carboxylate with MeI and MeOH at 120° (Ciamician & Anderlini, B 21, 2462) Basic liquid— B^+HAuCl_4 [110°] Yellow needles

Tetra methyl pyridine hexahydride $C_8H_{11}N$

νe $NH \begin{smallmatrix} \text{CHMe CHMe} \\ \text{CH}_2 \text{ CHMe} \end{smallmatrix} CHMe$ Parpervoline

(151°) Obtained by reducing the preceding in alcoholic solution with sodium (C & A) Liquid, smelling like pyridine Does not turn brown in air— B^+HAuCl_4 [119°] Yellow needles

Methylo iodide B^+MeI [262°] Prisms, v sol water, insol ether

A parvoline $C_8H_{11}N$ (c 200°) is present among the products of the putrefaction of horseflesh (Gautier, Bl [2] 48, 11) It is an oil which resinifies in the air, and forms a flesh coloured platinocchloride

A parvoline $C_8H_{11}N$ (188°) occurs among the bases obtained by distilling cinchonine with KOH (O de Coninck, C R 91, 296)

Penta methyl pyridine dihydride $C_9H_{11}N$

νe $NMe \begin{smallmatrix} \text{CMe CMe} \\ \text{CH CH} \end{smallmatrix} \text{CMe}$ (189°), (46° at 7 mm) Prepared by heating ν methyl pyrrole with MeI, K_2CO_3 , and methyl alcohol at 140° (Ciamician & Anderlini, Rend Accad Linc [4] 203, B 21, 2863, 22, 658) Formed also by heating tetra methyl pyridine dihydride with MeI (Anderlini, B 22, 2507) Oil, with strong alkaline reaction— B^+HAuCl_4 [100°] Yellow needles Reacts energetically with MeI, forming an oily product, whence successive treatment with $AgCl$ and $AuCl_3$ yields $C_9H_{11}NHAuCl_4$, which crystallises in thin golden yellow needles [100°]

METHYL PYRIDINE CARBOXYLIC ACID

$C_7H_7NO_2 \cdot 2e$ $N \begin{smallmatrix} \text{CMe CH} \\ \text{CH CH} \end{smallmatrix} \text{C CO}_2H$ Picoline

carboxylic acid Formed by heating methyl pyridine dicarboxylic (uvitonic) acid at 275° (Bottinger, B 14, 67, 17, 92) Fimetric crystals (containing aq), sol hot, sl sol cold, water and alcohol, nearly insol ether Sublimes without melting Forms salts both with acids and bases Yields on oxidation with $KMnO_4$ pyridine dicarboxylic (lutidmic) acid

Salts— HA^+HCl prisms— BaA^+ 11aq very soluble needles— CaA^+ aq very soluble four sided colourless prisms— CuA^+ aq blue pp— AgA^+ white pp

Methyl-pyridine carboxylic acid $C_7H_7NO_2 \cdot 2e$

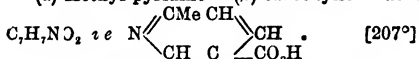
$N \begin{smallmatrix} \text{CH C(CO}_2H) \\ \text{CH CH} \end{smallmatrix} \text{CMe}$ Homomocotinic acid

Methyl-carbopyridine acid [212°] Obtained by heating methyl pyridine dicarboxylic (methyl-quinolinic) acid at 170°, or by warming it with HOAc (Hoogewerf & Van Dorp, R T C 2, 21) Formed also by the slow oxidation of (β) collidine by $KMnO_4$ (O de Coninck, A Ch [5] 27, 493, Bl [2] 43, 107) Needles, v sol hot

water Oxidised by KMnO_4 to pyridine dicarboxylic (cinchononic) acid

Salts — KA' small plates — CuA' small blue crystals — AgA' needles — $\text{HA}'\text{HCl}$ small prisms — $(\text{HA}')_2\text{H}_2\text{PtCl}_6$ orange prisms — $\text{HA}'\text{HAuCl}_4$ yellow needles

(a) Methyl pyridine (β) carboxylic acid



Formed by oxidising (a) methyl (β) ethyl pyridine (aldehyde collidine) with a 2 p.c. solution of KMnO_4 , allowing the mixture to stand for 48 hours, and then heating to 60° . The filtrate from MnO_2 is neutralised by H_2SO_4 and evaporated, the residue is extracted with alcohol and the acid purified by means of its silver salt (Dukopf, *B* 18, 3132, Ladenburg, *A* 217, 43). Prisms, v.e. sol water and alcohol. On distillation with lime it yields (a) methyl pyridine (128°). KMnO_4 oxidises it to pyridine dicarboxylic (isocinchononic) acid.

Salts — $(\text{HA}')_2\text{H}_2\text{PtCl}_6$ [240°] Aggregates of needles, v. sol water, insol ether alcohol — $(\text{CuA}')_2\text{Cu}(\text{OAc})_2$ Crystalline powder, formed by boiling a solution of the acid with cupric acetate. The silver salt is amorphous. The aurichloride forms yellow needles [c 202°].

Methyl pyridine carboxylic acid $\text{C}_6\text{H}_7\text{NO}_2$ is

$\text{N} \begin{array}{c} \text{C}(\text{CO}_2\text{H}) \text{CH} \\ \diagup \quad \diagdown \\ \text{CH} \quad \text{CH} \end{array} \text{CMe (?)}$ [c 260°] Formed in small quantity, together with pyridine (α)-dicarboxylic acid, by oxidising (α) di-methylpyridine with KMnO_4 (Bichr, *B* 21, 30-0). Plates (from alcohol), v.e. sol water. It is possibly identical with the isomeride described by Bottinger (*supra*).

(β) Methyl-pyridine (β') carboxylic acid

$\text{N} \begin{array}{c} \text{CH CMe} \\ \diagup \quad \diagdown \\ \text{CH C}(\text{CO}_2\text{H}) \end{array} \text{CH}$ [216°] Formed by heating (β) methyl pyridine (αβ') dicarboxylic acid with HOAc and Ac_2O at 225° (Dukopf a. Gottsch, *B* 23, 1113). White mass, m. sol water. Its aqueous solution is not coloured by FeSO_4 .

Methyl pyridine dicarboxylic acid

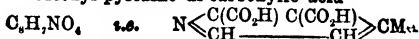
$\text{C}_6\text{H}_5\text{NO}_4$ is $\text{N} \begin{array}{c} \text{CMe} \text{---} \text{CH} \\ \diagup \quad \diagdown \\ \text{C}(\text{CO}_2\text{H}) \text{CH} \end{array} \text{CO}_2\text{H}$ *Uti* tonic acid [274°] Formed by the action of ammonia on pyruvic acid (Bottinger, *A* 188, 330, 208, 138, *B* 13, 2092, 16, 35, 17, 144). Formed also by oxidising s tri methyl pyridine,

or di methyl ethyl pyridine, $\text{N} \begin{array}{c} \text{CMe CH} \\ \diagup \quad \diagdown \\ \text{CMe CH} \end{array} \text{CLt}$, with KMnO_4 (Altar, *A* 237, 191, Dukopf, *B* 21, 2717). Minute six sided trimetric plates (Friedlander, *J* 1882 367), v. sl sol cold, sl sol hot, water, m. sol NH_4Ac and HClAc , v. sol aniline, phenol, HOAc , and glycerin, sl sol isoamyl alcohol and chloroform, insol benzene and CS_2 . Gives a violet red colour with FeSO_4 . It is a powerful antiseptic. Yields (a) methyl pyridine on distillation with lime. Split up by heat into CO_2 and $\text{N} \begin{array}{c} \text{CMe CH} \\ \diagup \quad \diagdown \\ \text{CH CH} \end{array} \text{CO}_2\text{H}$. Alkaline KMnO_4 oxidises it to pyridine tricarboxylic acid. Bromine water forms bromoform, CO_2 , and formic acid.

Salts — $(\text{NH}_4)\text{HA}'$ powder — CaA' 6aq: amorphous — CaA' 4aq: prisms — BaA' 2aq:

slender needles — CuA' 4aq — $\text{Cu}(\text{OH})_2\text{A}'_2$ 9aq * PbA' dense pp — AgA' 7aq gelatinous pp

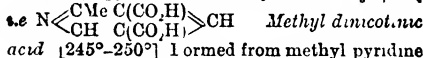
Methyl pyridine di carboxylic acid



Methyl quinolinic acid [c 183°] *S* 84 at 10° . Formed by the oxidation of (*Py* 1) methyl-quinoline (lepidine) (1 pt) by KMnO_4 ($7\frac{1}{2}$ pts) (Königs, *B* 12, 983, 14, 103, Hoogewerff a. Van Dorp, *R T C* 2, 15, *B* 13, 1639, 14, 645). Tables or prisms, sol hot, sl sol cold, water, sl sol alcohol, ether, and benzene. Decomposes on fusion into CO and methyl pyridine carboxylic acid [210°], which on further oxidation yields cinchononic acid. KMnO_4 oxidises it to pyridine (a) tricarboxylic acid. The aqueous solution of the acid gives pps with the acetates of Pb , Ba , and Cu , and a yellow colour with 1cSO_4 .

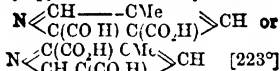
Salts — KHA' 2aq needles — KHA' 3aq — AgA' 7aq crystalline powder

Methyl pyridine dicarboxylic acid $\text{C}_6\text{H}_7\text{NO}_4$



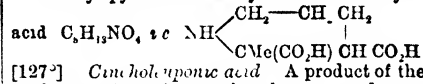
Formed from methyl pyridine tricarboxylic acid $\text{N} \begin{array}{c} \text{CMe} \text{---} \text{C}(\text{CO}_2\text{H}) \\ \diagup \quad \diagdown \\ \text{C}(\text{CO}_2\text{H}) \text{C}(\text{CO}_2\text{H}) \end{array} \text{CH}$ by heating at 150° (Weber, *A* 241, 9). Spherical groups of needles (containing aq), sl sol cold water. KMnO_4 oxidises it to a pyridine tricarboxylic acid — $\text{HA}'\text{HCl}$ aq transparent efflorescent crystals — PbA' 2aq crystalline pp.

Methyl pyridine dicarboxylic acid



Obtained by oxidising the di methyl ethyl pyridine, which is formed by the action of NH_3 on propionic aldehyde (Dukopf a. Gottsch, *B* 23, 6-8, 1110), and by the action of paraldehyde on propionic aldehyde ammonia (Dukopf a. Schlaun, *B* 21, 834). White powder (from hot water). The K , Ag , and Cu salts are sl sol water.

Methyl pyridine hexahydrate dicarboxylic



[127°] Cincholeuponic acid. A product of the oxidation of cinchonine by chromic acid mixture (Skiaup, *M* 9, 780). Prisms (containing aq), v.e. sol water, insol alcohol and ether. Ac_2O at 125° forms an amorphous acetyl derivative $\text{C}_6\text{H}_5\text{AcNO}_3$. Distillation of the lead salt over zinc dust yields a small quantity of pyridine.

Salts — PbA' powder, v.e. sol water — $\text{HA}'\text{HCl}$ [194°] Trimetric crystals [$\alpha_D = 34.4^\circ$ at 18°].

Nitrosamine $\text{C}_6\text{H}_5(\text{NO})\text{NO}_4$ [163°] Trimetric crystals, sl sol cold water, m. sol alcohol. Warm conc HClAc reproduces $\text{C}_6\text{H}_5\text{NO}_4$ and nitrous acid — BaA' (at 115°). Deliquescent amorphous powder, in sol alcohol.

In the preparation of cincholeuponic acid a base called cincholeupone $\text{C}_6\text{H}_7\text{NO}_3$ is also formed. It yields cincholeuponic acid on oxidation with chromic acid mixture, and ethylpyridine on distillation over red hot zinc dust. It forms the salts $(\text{C}_6\text{H}_7\text{NO}_3)\text{HCl}$ [200°], $\text{B}'_2\text{H}_2\text{PtCl}_6$ 8½aq and $\text{B}'\text{HAuCl}_4$ [203°], an acetyl

derivative $C_6H_4AcNO_2$ [121°], and a nitrosamine $C_6H_4(NO)NO_2$ [84°]

Methyl-pyridine tricarboxylic acid $C_6H_4NO_3$, *s.e.* $N \left\langle \begin{array}{c} CMe \\ C(CO_2H) \end{array} \right\rangle CH$ [226°] Formed

by oxidising $N \left\langle \begin{array}{c} CMe \\ C(CO_2H) \end{array} \right\rangle CH$ with $KMnO_4$ (Weber, *A* 241, 6) Spherical aggregates (containing aq) At 150° it slowly gives off CO_2 , yielding methyl pyridine dicarboxylic acid [245°–250°] — KH_2A'' , 6aq spherical aggregates — AgH_2A'' 2aq needles (from hot water)

Methyl-pyridine tricarboxylic acid

$N \left\langle \begin{array}{c} C(CO_2H) \\ C(CO_2H) \end{array} \right\rangle CMe$ *Methyl carbo-nicotinic acid* Formed by the oxidation of (α)-di-methyl pyridine dicarboxylic acid by $KMnO_4$ (Weber, *A* 241, 25) Needles (containing aq) or prisms (containing 2aq) Less soluble in water than the preceding acid Turns yellow at 205°, and completely decomposes at 260° $FeSO_4$ colours its solutions deep red The calcium salt yields (γ) methyl pyridine on distillation

Methyl-pyridine tricarboxylic acid

$N \left\langle \begin{array}{c} C(CO_2H) \\ C(CO_2H) \end{array} \right\rangle CH$ *Picoline tricarboxylic acid* [238°] Formed by the oxidation of flavenol (1 mol) with $KMnO_4$ (9 mols) (Fischer & Tauber, *B* 17, 2926) Formed also by oxidising potassium tri methyl pyridine carboxylate with aqueous $KMnO_4$ (Michael, *A* 225, 140) Slender needles (from water) Does not combine with acids On further oxidation with $KMnO_4$ it yields pyridine tetra carboxylic acid [227°] $FeSO_4$ colours its solution brownish-red

Salts — BA'' amorphous pp — Ag_2A''

Methyl-pyridine tetracarboxylic acid

$C_6H_4NO_4$, *s.e.* $N \left\langle \begin{array}{c} C(CO_2H) \\ C(CO_2H) \end{array} \right\rangle CMe$ [199°] Obtained by boiling potassium tri-methyl-pyridine dicarboxylate with a solution of $KMnO_4$ (Hantzsch, *A* 215, 57) Small prisms (from water), v e sol water, m sol alcohol, sl sol ether Gives (γ) picoline on distillation with lime Its neutral salts crystallise with difficulty Neutral solutions give pps with salts of Pb and Ag, mercurous salts, and $Ba(OAc)_2$, but no pps with dilute $BaCl_2$ or with salts of Mg, Zn, Mn, Ni, Co, and Cu, and mercuric salts — K_2H_4A'' 4aq large trimetric tables, v sol hot water, with acid reaction — KH_2A'' 2aq — Ca_2A'' 4aq, ppd by adding NH_3 and $CaCl_2$ — Mg_2A'' 6aq

Di-methyl-pyridine carboxylic acid

$C_6H_4NO_3$, *s.e.* $N \left\langle \begin{array}{c} CMe \\ CH \end{array} \right\rangle CMe$ *Lutidine carboxylic acid* Obtained by saponifying with alcoholic potash its ether, which is formed by adding acetic aldehyde (50 pts) to a mixture of aceto acetic ether (130 pts) and aldehyde-ammonia (61 pts), the reaction, which sets in at once, being completed by heating to 100° (Michael, *B* 18, 2020) Transparent prisms (containing 2 aq), v sol water and alcohol Gives (α)-lutidine on distillation with lime On oxidation with $KMnO_4$ it yields pyridine dicarboxylic (carboanichomeronic) acid

Salts. — $HA''HCl$ [166°] Large prisms or thick tables — $(HA'')_2H_2PtCl_4$ 2aq [216°]. Reddish-yellow prisms

Ethyl ether EtA' (246° uncor) Yellowish

oil, not volatile with steam — $(EtA')_2H_2PtCl_4$ [191°] Thin yellow pointed prisms, v . sl sol water and strong alcohol

Di-methyl-pyridine carboxylic acid

$N \left\langle \begin{array}{c} CMe \\ C(CO_2H) \end{array} \right\rangle CH$ *Di-methyl-picolinic acid* [153°] Formed by oxidising *s*-tri-methyl-pyridine with $KMnO_4$ (Altar, *A* 237, 188) Small crystals, v e sol water and alcohol, m sol ether Yields (α) di methyl pyridine on distillation with lime Its metallic salts are v sol water — $B'HCl$ aq minute white needles, v e sol water — $B'_2H_2PtCl_4$ 4EtOH [221°] Prisms

Di-methyl pyridine carboxylic acid

$N \left\langle \begin{array}{c} CMe \\ CMe \end{array} \right\rangle C(CO_2H)CH$ *Di-methyl-nicotinic acid* [160°] Prepared by distilling the mono ethylic ether of di methyl pyridine dicarboxylic acid $N \left\langle \begin{array}{c} CMe \\ CMe \end{array} \right\rangle C(CO_2H)CH$ (Weiss, *B* 19, 1308)

Needles (containing $\frac{1}{2}$ aq), v e sol water Oxidised by $KMnO_4$ to the corresponding pyridine tricarboxylic acid — AgA' — $HAHCl$ small prisms — $(HA')_2H_2PtCl_4$ 2aq orange needles

Di-methyl-pyridine carboxylic acid

$N \left\langle \begin{array}{c} CH \\ C(CO_2H) \end{array} \right\rangle CMe$ [151°] Formed by oxidising the parvoline obtained from propionic aldehyde and NH_3 (Dürkopff a Götsch, *B* 23, 687, 1110) — $(HA')_2H_2PtCl_4$ EtOH v sol water, sl sol alcohol Decomposes at 260°

Di-methyl-pyridine carboxylic acid

$C_6H_4Me_2NCO_2H$ Formed by saponifying, by alcoholic potash, its ethyl ether, which is produced from acetoacetic ether (30 g) by heating with formic aldehyde (5 g) and $ZnCl_2$ (30 g) for 12 hours at 100° (Canzonieri a Spica, *G* 14, 449) Silky needles, v e sol water — $HA'HCl$ aq [220°] Transparent rhombohedra (from water) — $(HA')_2H_2PtCl_4$ red crystals

Ethyl ether EtA' (260°) Oil

Di-methyl-pyridine dicarboxylic acid

$C_6H_4NO_4$, *s.e.* $N \left\langle \begin{array}{c} CMe \\ CH \end{array} \right\rangle CMe$ (α) *Di-methyl-di-methyl-pyridine dicarboxylic acid* [258°] Obtained by heating (α) di methyl pyridine tricarboxylic acid at 175° (Weber, *A* 241, 20) Formed also by oxidation of the corresponding tetra methyl pyridine (Dürkopff a Götsch, *B* 23, 1112) Needles (containing 2 aq (W) or anhydrous (D a G) Its solution is not coloured by $FeSO_4$ — PbA'' gelatinous pp, becoming crystalline on boiling — $HA''HCl$ aq slender needles, decomposed by water — $(H_2A'')_2H_2PtCl_4$ orange tables, melting above 300°

Di methyl pyridine dicarboxylic acid

$N \left\langle \begin{array}{c} CMe \\ C(CO_2H) \end{array} \right\rangle CH$ *or*
 $N \left\langle \begin{array}{c} CMe \\ C(CO_2H) \end{array} \right\rangle CMe$ [245°].

Formed by oxidising *s* tri methyl pyridine carboxylic acid in neutral solution with $KMnO_4$ (Michael, *A* 225, 187). Prisms (from water). — CaA'' — MgA'' 8aq — $(HA'')_2H_2PtCl_4$ 6aq

Di-methyl-pyridine dicarboxylic acid

$N \left\langle \begin{array}{c} CMe \\ CMe \end{array} \right\rangle C(CO_2H)CH$ *Lutidine dicarboxylic acid* Formed by saponifying with alcoholic potash its ether, which is produced by passing nitrous acid gas into di methyl isopropyl dicarboxylic acid

oxylic ether in alcohol (Engelmann, *A* 231, 50) Its ether is also one of the products of the action of acetoacetic ether on hexamethylene tetramine at 170° (Griess, *B* 21, 2740) The same ether appears to be formed by oxidising with nitrous acid gas the product obtained by the action of phenyl acetic aldehyde on aceto acetic ether and ammonia (Jeauraud, *B* 21, 1784) Slender needles (containing $\frac{1}{2}$ aq) Melts at a very high temperature ν sol cold water, alcohol, and ether KMnO_4 oxidises it to pyridine tetracarboxylic acid On distillation it yields CO_2 and di-methyl pyridine carboxylic acid Distillation over CaO yields (aa) di methyl pyridine

Salts — $\text{BaA}''2\text{aq}$ — $\text{PbA}''2\text{aq}$ amorphous pp changing to stout prisms — $\text{H}_2\text{A}''\text{HCl}2\text{aq}$ prisms

Mono ethyl ether EtHA'' [131°] Obtained by decomposing the di ethyl ether (1 mol) with alcoholic KOH (1 mol) (Weiss, *B* 19, 1308) Needles (from water) Its neutral solution is ppd by salts of Ag , Hg , and Cu — $\text{EtHA}''\text{HCl}2\text{aq}$ [90°] Needles (from water)

Diethyl ether Et.A'' [72°] (302°) Long white needles — $\text{Et.A}''\text{HAcCl}$, ν sol alcohol

Di methyl pyridine dicarboxylic ether di hydride $\text{C}_6\text{H}_7\text{NMe}_2(\text{CO}_2\text{Et})_2$ [170°] Formed by heating acetoacetic ether (20 g) with ZnCl_2 (20 g) and hexamethylene tetramine (4 g) at 100° (Griess a Harrow, *B* 21, 2740) Four sided plates or needles, almost insol water, sl sol cold alcohol and ether, ν sol chloroform By treatment with nitrous acid, or even by solution in hot dilute HCl , it is oxidised to $\text{C}_6\text{H}_7\text{NMe}_2(\text{CO}_2\text{Et})_2$.

Di methyl pyridine tricarboxylic acid

$\text{C}_{10}\text{H}_9\text{NO}_5 \cdot \frac{1}{2} \text{N} \left\langle \begin{smallmatrix} \text{CMe} \\ \text{C}(\text{CO}_2\text{H}) \end{smallmatrix} \right\rangle \text{CMe}$ *Lu thidine tricarboxylic acid Dimethylcarbodinicotinic acid* [212°] Formed by oxidation of potassium tri methyl pyridine dicarboxylate with KMnO_4 in a solution kept neutral by CO_2 (Hantzsch, *A* 215, 52, Weber, *A* 241, 20) Crystallises from water in hard crusts composed of rhombohedra (containing 2aq) Split up by heat into CO_2 and ($\alpha\gamma$)-di-methyl pyridine di carboxylic acid Yields ($\alpha\gamma$) di methyl pyridine on distillation with lime Its neutral solutions give no pps with salts of Cu , Ag , or Pb , nor with FeCl_3 , but with $\text{Hg}(\text{NO}_3)_2$ a pp soluble in hot water

Salts — $\text{KH}_2\text{A}'''2\text{aq}$ — $\text{Ba.A}'''(8?)\text{aq}$ minute needles — $\text{Mg.A}'''10\text{aq}$ — $\text{Ca.A}'''8\text{aq}$ — $\text{Ag.A}'''$

Di methyl pyridine tricarboxylic acid

$\text{N} \left\langle \begin{smallmatrix} \text{CMe} \\ \text{C}(\text{CO}_2\text{H}) \end{smallmatrix} \right\rangle \text{CMe}$ S 178 at 8° Formed by oxidising potassium styryl di methyl pyridine dicarboxylate with cold aqueous KMnO_4 (Epstein, *A* 231, 11) Crystallises from water in prisms (containing aq), almost insol water, alcohol, ether, benzene, and chloroform Turns brown at about 220° but decomposes without melting Gives (aa) di methyl pyridine on distillation with lime A solution of its ammonium salt gives pps with CaCl_2 , CaSO_4 , $\text{Hg}_2(\text{NO}_3)_2$, AgNO_3 , SnCl_4 , $\text{Pb}(\text{OAc})_2$ and $\text{Bi}(\text{NO}_3)_3$, but not with BaCl_2 , MgSO_4 , MnSO_4 , and alum, nor, in the cold, with ZnSO_4 or CuSO_4 FeCl_3 gives a reddish colour

Salts.— $\text{H}_2\text{A}'''\text{HCl}$ needles, decomposed by

water and by alcohol — $\text{Ph.A}'''\text{aq}$. — $\text{Cu}(\text{NH}_4)\text{A}'''4\text{aq}$ — $\text{Ag}_2\text{A}'''8\text{aq}$

Tri-methyl pyridine carboxylic acid

$\text{C}_6\text{H}_{11}\text{NO}_5 \cdot \frac{1}{2} \text{N} \left\langle \begin{smallmatrix} \text{CMe} \\ \text{C}(\text{CO}_2\text{H}) \end{smallmatrix} \right\rangle \text{CMe}$ *Collidine carboxylic acid* [155°] Obtained by saponifying with alcoholic potash its ethyl ether, which is obtained by passing nitrous acid into its dihydride (Hantzsch, *A* 215, 42) and by heating the mono-ethyl ether of tri methyl pyridine dicarboxylic acid (Michael, *A* 225, 131) Crystallises from water in short needles or cubes (containing 2aq) Melts at 110° when hydrated, or 155° when anhydrous On oxidation with KMnO_4 it yields di methyl pyridine dicarboxylic acid, methyl pyridine tricarboxylic acid, and pyridine tetracarboxylic acid

Salts — KA' needles (from alcohol) — $\text{CaA}'\text{aq}$ — $\text{HA}'\text{HCl}$ needles or prisms. — $(\text{HA}')_2\text{H}_2\text{PtCl}_6\text{aq}$

Ethylether EtA' (256°) SG 15 10815 Liquid, ν sol ether, alcohol, benzene, and chloroform, insol water, sol dilute acids — $(\text{EtA}')_2\text{H}_2\text{PtCl}_6$ [193°] Prisms (from alcohol) — $\text{EtA}'\text{MeI}$ [128°] Needles, ν sol water and alcohol, insol ether With AgO it yields crystalline $\text{C}_{10}\text{H}_{11}\text{NO}_53\text{aq}$, whence $\text{C}_{10}\text{H}_{11}\text{NO}_5\text{HCl aq}$ (Hantzsch, *B* 19, 35)

Tri methyl pyridine dihydride carboxylic ether $\text{C}_6\text{H}_7\text{NMe}_2\text{CO}_2\text{Et}$ Formed by warming $\text{C}_6\text{H}_7\text{NMe}_2(\text{CO}_2\text{Et})_2$ with HClAq (of 25 p.c.) at 100° (Hantzsch, *A* 215, 40) Feebly basic oil. — $(\text{HA})_2\text{H}_2\text{PtCl}_6$ minute needles

Tri methyl pyridine dicarboxylic acid

$\text{C}_{10}\text{H}_{11}\text{NO}_5 \cdot \frac{1}{2} \text{N} \left\langle \begin{smallmatrix} \text{CMe} \\ \text{C}(\text{CO}_2\text{H}) \end{smallmatrix} \right\rangle \text{CMe}$

Formed by saponifying with alcoholic potash its ether, which is obtained by the action of nitrous acid gas and alcohol upon its dihydride, which is itself produced by the action of aldehyde-ammonia on aceto acetic ether (Hantzsch, *A* 215, 26) Fluffy mass of needles (from hot water), melting above 300°, ν sl sol cold, ν sol hot, water, ν sl sol alcohol and ether Oxidised by KMnO_4 to di methyl pyridine tricarboxylic acid, methyl pyridine tetra carboxylic acid, and pyridine pentacarboxylic acid Decomposed by heat into CO_2 and tri methyl pyridine carboxylic acid Bromine acting on an aqueous solution of the K salt yields di bromo- α tri methyl pyridine The salts which it forms with alkalis and alkaline earths are very soluble, crystallise badly, have an alkaline reaction, and split up on heating into α tri-methyl-pyridine and carbonates FeCl_3 gives an intense red colour to a solution of the K salt. — KA'' — $(\text{CaA}'')_2\text{CuO}11\text{aq}$ — $\text{Ag}_2\text{A}''8\text{aq}$ — $\text{H}_2\text{A}''\text{HCl}2\text{aq}$ — $(\text{H}_2\text{A}'')_2\text{H}_2\text{PtCl}_6$

Di-methyl ether Me.A'' [32°] (386°). Formed by oxidation of its dihydride by nitrous acid (Hantzsch, *B* 16, 1947) White needles, ν sol water — $\text{Me.A}''\text{HCl}2\text{aq}$ long glistening prisms melting, when anhydrous, at 142° — $(\text{Me.A}'')_2\text{H}_2\text{PtCl}_6$ [200°] Orange spangles. — $\text{Me}_2\text{A}''\text{HAcCl}$ [105°] Slender light yellow needles. — $\text{Me}_2\text{A}''\text{HNO}_3$ [140°] Prisms

Mono ethyl ether EtHA'' 2aq Formed from the neutral ether by boiling with the calculated quantity of potash dissolved in alcohol. Needles in radial groups (from alcohol), ν sol

water, v. al sol ether Melts, when anhydrous, at 157° — $\text{EtEtA}''\text{HCl}$ [178°] Cubes — (EtEtA''), H_2PtCl_6 2aq prisms (from cold water) Melts, when anhydrous, at 219° — BaEtA'' , 3aq — $\text{BaEt}_2\text{A}''$, aq — $\text{CaEt}_2\text{A}''$, 8aq needles — $\text{CuEt}_2\text{A}''$ — $\text{CdEt}_2\text{A}''$, 4aq — $\text{ZnEt}_2\text{A}''$, 6aq — $\text{AgHEt}_2\text{A}''$, aq monoclinic prisms

Di-ethyl ether EtA'' (308°–310°) SG 15 1087 Formed from its dihydride (*v. infra*) by treatment with HNO_3 , or, better, by alcohol and nitrous acid gas (Hantzsch, *A* 215, 21) Bright yellow viscid oil with faint odour and burning taste Not volatile with steam It has no action on litmus It does not ppt metallic salts With HgCl_2 it gives, only after a time, silky needles of a double compound It dissolves in dilute acids — $\text{EtA}''\text{HCl}$ very hygroscopic — (EtA''), H_2PtCl_6 rose-coloured tables [184°] — $\text{EtA}''\text{HNO}_3$ needles [92°] $\text{Et}_2\text{A}''\text{HI}$ [170° – 173°] Dark plates, which dissolve with rotation in cold water — $\text{Et}_2\text{A}''\text{HI}$, violet pyramids

Methylo iodide of the diethyl ether EtA''MeI [140°] Formed by heating the ether with MeI and MeOH at 120° Crystalline mass, more soluble in water or alcohol than $\text{Et}_2\text{A}''\text{HI}$ It is ppd unaltered from its aqueous solution by NaOH Its aqueous solution is acid to litmus It is not affected by fuming HCl at 150° or by alcoholic NH_3 at 150° Concentrated aqueous KOH forms methyl di-carbo-collidylum dehydride $\text{C}_{11}\text{H}_{11}\text{NO}_4$ This substance crystallises in needles, [92°], v. e sol alcohol, v. al sol cold water, and distils without decomposition at a temperature far above 360° It has neither acid nor basic properties When heated with H_2SO_4 at 150° – 180° it yields CO_2 , HOAc, and methyl pseudo lutidostyryl $\text{C}_9\text{H}_{11}\text{NO}$ When 'methyl dicarbo collidylum dehydride' is heated in a current of HCl it loses CO_2 and forms methyl-carbo collidylum dehydride $\text{C}_9\text{H}_{11}\text{NO}_3$ This is also an indifferent body, it crystallises in needles [103°], v. sol water, and boils above 340° On heating with H_2SO_4 it evolves acetic acid and gives methyl pseudo lutidostyryl (Hantzsch, *B* 17, 1023)

Methylo-chloride of the diethyl ether EtA''MeCl Formed by treating $\text{Et}_2\text{A}''\text{MeI}$ with AgCl (Hantzsch, *B* 17, 1019) Crystals Yields (EtA''), Me_2PtCl_6 , which crystallises in thick yellow prisms

Tri methyl-pyridine dihydride dicarboxylic acid $\text{C}_{10}\text{H}_8\text{NO}_4$, $\text{e NH} < \text{CMe C(CO}_2\text{H)} > \text{CHMe}$

Methyl ether MeA'' [156°] Formed by the action of methyl acetoacetate on aldehyde ammonia (Hantzsch, *B* 16, 1946) When heated with HCl it exchanges CO_2Me for H and yields the methyl ether of the monocarboxylic acid On oxidation with nitrous acid it gives $\text{C}_9\text{Me}_2(\text{CO}_2\text{Me})\text{N}$

Di-ethyl ether EtA'' [131°] Formed by heating acetoacetic ether (52 g) with aldehyde ammonia (13.5 g) (Hantzsch, *A* 215, 8)

Properties — Compact colourless tables with many facets (from alcohol), with bright-blue fluorescence V. al sol boiling water, al sol cold alcohol, ether and CS_2 , m sol benzene, v. sol boiling alcohol, v. e sol chloroform. Boils above 315° with decomposition

Reactions — 1 Boiling conc KOHAq has no action — 2 Alcoholic NH_3 at 150° has no action — 3 Alcoholic potash slowly decomposes it, forming NH_3 and K_2CO_3 . — 4 Boiling dilute acids do not dissolve it, it forms no platinumchloride — 5 Conc H_2SO_4 dissolves it, but on pouring into water it is reppd unchanged — 6 Warm conc HCl completely destroys it, forming CO_2 , EtCl , NH_3 , aldehyde, and acetone — 7 HCl passed into an ethereal solution removes the H_2 (Hantzsch, *A* 215, 37), and forms other products — 8 Aqueous HCl (25 p.c.) at 100° forms hydro tri methyl pyridine mono carboxylic ether — 9 Dilute HCl at 130° forms ethyl chloride, CO_2 , and tri methyl pyridine dihydride Another product is an oil composed of a ketone $\text{C}_9\text{H}_{11}\text{O}$ (208° – 209°), V D 4.20, which combines with NaHSO₃, and with bromine, and with hydroxylamine yields crystals [76°] — 10 Nitrous acid removes H_2 , leaving tri methyl pyridine dicarboxylic ether — 11 Bromine dissolved in CS_2 forms a dibromide of di bromo tri methyl pyridine dihydride carboxylic ether $\text{NC}_9\text{H}_8\text{Br}_2(\text{CO}_2\text{Et})_2\text{H}_2\text{Br}_2$ [88°] This is extremely soluble in boiling alcohol, and crystallises as yellow twin crystals Fuming HNO_3 converts it into the di bromide of di bromo tri methyl pyridine di carboxylic ether [102°] — 12 Chloride forms the di chloride of penta chloro tri methyl pyridine dicarboxylic ether, crystallising as woolly needles [150°]

Tetra methyl pyridine dihydride carboxylic ether $\text{C}_{10}\text{H}_{12}\text{NO}_4$, e e
 $\text{NMe} < \text{CMe C(CO}_2\text{Et)} > \text{CHMe}$ [86°] Formed by the action of paraldehyde and H_2SO_4 on methyl amido oxy butyric ether (Kuckert, *B* 18, 620), and, in small quantity, by the action of methylamine on a mixture of acetoacetic ether and aldehyde (Hantzsch, *B* 18, 2580) Crystals, with blue fluorescence

METHYL-PYRIDINE-HYDRIDES *v.* METHYL PYRIDINE

DI - METHYL - DIPYRIDYL $\text{C}_{10}\text{H}_{12}\text{N}_2$, e e
 $\text{NO}_2\text{H}_2\text{Me C}_9\text{H}_8\text{MeN}$ *Dipicolyl* [84°] (230° – 298°) Obtained by treating (a) picoline with sodium at 80° – 90° , exposing the product to air, and fractionally distilling (Ahrens, *B* 21, 2930, Heuser, *J pr* [2] 42, 430) Very deliquescent yellowish needles With water it forms a compound (containing 4aq) melting at 38° When oxidised by potassium permanganate it yields $\text{NC}_9\text{H}_8\text{Me C}_9\text{H}_8\text{N CO}_2\text{H}$ [193°], which on heating by itself or with HOAc yields methyl dipyridyl [94°] — $\text{B}''\text{H}_2\text{Cl}_2$ Hygroscopic plates (from alcohol) Picrate $\text{B}''2\text{C}_9\text{H}_8(\text{NO}_2)_3\text{OH}$ [240°] — $\text{B}''\text{H}_2\text{Cl}_6\text{HgCl}_2$ [220°] — $\text{B}''\text{H}_2\text{PtCl}_6$ small plates, v. al sol water, v. sol HClAq Discolours at 235° , but only partially decomposed at 275° — $\text{B}''2\text{HAuCl}_4$ nodules (from conc HClAq) decomposes at 200° (A), [210°] (H) — The tin double salt melts at 180°

Di methyl-dipyridyl dodecahydride $\text{C}_{12}\text{H}_{18}\text{N}_2$, e e $\text{MeNC}_9\text{H}_8\text{C}_9\text{H}_8\text{NMe}$ *Di-methyl dipyridyl*, (230° – 235°) Formed, together with tri methyl dipyridyl dodecahydride, by treating the product of the action of MeI upon dipyridyl dodecahydride (dipyridyl) with Ag_2O (Liebrecht, *B* 19, 2595) Liquid, miscible with water, but separating on addition of NaOH — $\text{B}''\text{H}_2\text{Cl}_2(\text{HgCl}_2)_2$

flocculent pp, sol hot water — $B'H_2PtCl_4$, small dark red crystals

Tri-methyl dipyriddy dodecahydride $C_{11}H_{22}N_2$, α, β NMe C_4H_7 , C_4H_7 Me NMe (205° — 212°) Formed as above Yellowish oil, insol water — $B'H_2PtCl_4$, yellowish red crystals, v e sol water

TRI METHYL-PYRIDYLENE DIHYDRIDE DI-METHYL DIKETONE $C_{12}H_{11}NO_2$ α, β N $\langle CMe CH(CO CH_3) \rangle CMe$ [153°] (250° *in vacuo*) Formed by heating methylene di-methyl diketone (2 mols) with aldehyde-ammonia (1 mol) on the water bath (Combes, *B* [2] 51, 15) Hexagonal prisms (from alcohol), insol water

METHYL PYRIDYL ACETYLENE TETRA-HYDRIDE $C_8H_9NMe C H$ Formed by boiling the hydrobromide of anhydro eegonine dibromide with aqueous K_2CO_3 (Eichengrün a Einhorn, *B* 23, 2879) Oil — $B'H_2AuCl_4$ [179°]

DI METHYL DI PYRIDYL PROPANE DO DECAHYDRIDE $C_{11}H_{12}N_2$ α, β ($C_4H_7NMe CH$), CH_2 *Methyl iodide* $B'Me I$ Obtained by heating ($C_4H_7N CH_2$), CH_2 , with MeI (Ladenburg, *B* 21, 3102) White non hygroscopic crystals, v sol hot water The aurochloride $B'(H_2AuCl_4)$ [171°] obtained from the methyl iodide, crystallises from water in needles

(*Py* 15) **DI METHYL PYRIDYL-(Py 3)-QUINOLINE** $C_{10}H_{11}N$ α, β

$C_2NH_2Me, C_4H_7 \begin{matrix} \diagup CH CH \\ \diagdown N CH \end{matrix}$ *Lutidyl quinoline*

[109°] Colourless glistening crystals Formed by heating *m*-amido phenyl di methyl pyridine $C_2NH(CH_3) C_4H_7NH$ with glycerine, nitrobenzene, and H_2SO_4 The chloride and nitrate form white needles, the platino chloride small orange needles, the auro chloride long golden yellow needles (215°) (Lepetit, *G* 20, 2399, (*G* 17, 473))

METHYL - PYROCATECHIN $C_8H_9O_2$ α, β $C_4H_7Me(OH)$ [134] Formed by heating creosol $C_8H_9Me(OH)(OH)$ [134] with HCl (H Müller, *C N* 10, 269) Formed also by the dry distillation of di oxy toluene ((α) homo protocatechuic acid (Tiemann a Nagai, *B* 10, 210) and from $C_4H_7Me(NO)(OH)$ [134] by displacement of NO by OH (Neville a Winther, *B* 15, 2983) Liquid, which may be distilled, v e sol water, alcohol, and ether Reduces Fehling's solution and ammoniacal $AgNO_3$ in the cold $FeCl_3$ gives a green colouration, turned reddish-violet by ammonia

Mono methyl ether $C_8H_9Me(OH)(OMe)$ [134] *Isocresol* (α 185°) Prepared from $C_4H_7Me(NH_2)(OMe)$ [134] by the diazo reaction (Limpach, *B* 22, 350) Yellow liquid, v sol alcohol and ether, sl sol water Volatile with steam

Mono methyl ether $C_8H_9Me(OMe)(OH)$ [134] v CREOSOL

Isomerides v Methyl ethers of PYROCATECHIN

TETRA-METHYL PYROCROLL $C_8H_9N_2O_2$ α, β $NO, HMe, \begin{matrix} CO \\ CO \end{matrix} C_4H_7Me, N$ [272 5°] Formed, together with ($\alpha\beta$)-di methyl pyrrole, by heating ($\alpha\beta$)-di-methyl pyrrole ($\alpha\beta$) dicarboxylic anhydride at 850° (Magnanini, *B* 21, 2877, 22, 2502) Trimetric yellow needles (by sublimation), insol

water, v sol chloroform Its molecular weight was determined by Raoult's method Boiling alcoholic KOH gives

$NO, C_4H_7Me, CO NO, HMe, CO, K$

METHYL - PYROGALLOL $C_8H_9O_2$ α, β $C_4H_7(CH_3)(OH)$, [129°] Prepared by heating the dimethyl ether with HCl to 150° Sublimes in small needles Sl sol benzene

Dimethyl ether $C_8H_9(CH_3)(OH)(OMe)$, [36°] (265°) Crystalline Occurs together with the dimethyl ether of pyrogallol in beech wood tar, oil, from which it can be isolated by means of its benzoyl compound [118°] (Hofmann, *B* 12, 1371) With bromine it yields $C_6Br_2(CH_3)(OH)(OMe)_2$ [126°] If a mixture of the sodium salts of the dimethyl ethers of pyrogallol and methyl pyrogallol is oxidised by heat in the air or with C_2Cl_6 eupitonic acid (pitakal) is formed

METHYL-PYROMUCIC ACID $C_8H_9O_4$ α, β $C_4H_7MeO CO H$ [108°] Formed by the oxidation of methyl furfuraldehyde (Hill, *B* 22, 608) and by the action of conc KOH aq upon that aldehyde (Bicler a Tollens, *A* 28, 123) Needles, more soluble than pyromucic acid in water, benzene, and chloroform With $NaOH$ and H_2SO_4 it gives a deep green colour on warming $FeCl_3$ gives a brown pp

Reaction — Bromine (2 mols) added to its aqueous solution gives off CO and forms an acid $C_8H_9O_4$ (? acetylacrylic), which crystallises in broad flat needles [123°], s 67 (in the cold). This acid is v sol alcohol, ether, and hot water, it forms an addition product [108°] with bromine, a silver salt $AgC_4H_7O_3$, crystallising in slender needles, and a phenyl hydrazide [157°] apparently identical with that of acetyl acrylic acid (Hill a Hendrixson, *B* 23, 422, cf Bender, *B* 21, 2494, Decker, *B* 21, 2937)

Salt — $AgC_4H_7O_3$, slender needles (from hot water), sl sol hot water

DI-METHYL-PYRONE v Anhydride of DI-METHYLENE DI-METHYL DIKETOYL

Di-methyl-pyrone carboxylic acid v Di-HYDRACTIC ACID

Di-methyl-pyrone dicarboxylic acid v *Anhydride of DI-METHYLENE DI-METHYL TRIKETOYL*

METHYL PYRROLE C_4H_5N α, β $NMe \begin{matrix} CH CH \\ CH CH \end{matrix}$ *Methyl pyrrolone* (113°)

(*B*), (114° i v) (*C* a *D*) SG 12 9203 (*B*) Formed by the dry distillation of methylamine mucate (*C* a Bell, *B* 10, 1866) Formed also by treating potassium pyrrole with MeI (Ciamician a Dennstedt, *B* 17, 2951) Liquid Resolved by heating with KOH into pyrrole and $MeOH$ When v methyl pyrrole (3 g) is heated with $MeOH$ (5 g), MeI (7 g), and K_2CO_3 (3 g) for 10 hours at 140° there is formed a tri methyl pyrrol (160°-165°) and penta methyl pyridine dihydride C_8H_9Me, NMe (Ciamician a Anderlin, *B* 22, 656) Methyl pyrrole acts on alloxan in warm aqueous solution forming $C_8H_9N_2O$, α, β $NH_2, CO NH CO CO CO C_4H_7NMe$, which crystallises in white plates, sol hot water, sl sol alcohol and cold water (Ciamician a Silber, *B* 19, 1710)

v-Methyl-pyrrole dihydride C_8H_9NMe , *Methyl-pyrrolone* (80°) Formed by reduction of *v*-methyl pyrrole with zinc dust and $HOAc$ (Ciamician a Magnaghi, *B* 18, 725) Colour-

less, strongly alkaline liquid, miscible with water. Tertiary base. Its hydrochloride forms colourless crystals. — $B^*H_2PtCl_4$, trimetric crystals, v. s. sol water (Le Valle, *G* 15, 490)

Methylpyrrole iodide C_4H_5NMeI [286°] Obtained by the action of MeI on the above or on pyrrole dihydride (Ciamician, *B* 16, 1541, *G* 15, 492) Pearly plates. Yields the platinum-chloride ($C_4H_5NMe_2Cl$), $PtCl_2 \cdot xaq$

Methylpyrrole tetrahydride C_4H_9N v. s. $NMe \begin{smallmatrix} CH_2CH_2 \\ CH_2CH_2 \end{smallmatrix}$. **Methyl pyrrolidine** (82°)

Formed by heating the dihydride with fuming $HIAq$ and amorphous phosphorus at 250° (Ciamician & Magnaghi, *G* 15, 493) Liquid, miscible with water

Methylpyrrole iodide C_4H_5NMeI Formed by the action of MeI on methyl pyrrole tetrahydride or on pyrrole tetrahydride. Deliquescent needles, v. s. sol alcohol, insol ether and $NaOHaq$. Ag_2O yields $C_4H_5NMe_2OH$, which is decomposed by distillation into MeOH and C_4H_5NMe . Distillation of the iodide with solid KOH yields C_4H_5NMe [89°–92°]

(a) **Methyl pyrrole** C_4H_5N v. s. C_4H_5NMe NH (a) **Homopyrrole** (148°) Occurs, together with the (b) isomeride, in bone oil. The fraction 140°–150° is boiled with KOH, and the potassium derivatives that separate are washed with ether, decomposed by water, and fractionally distilled (Ciamician, Dennstedt, a Zimmermann, *B* 19, 178, 2200, 22, 1918, Weidel a Ciamician, *B* 13, 77) The isomerides may also be separated by conversion into their carboxylic acids, separation of these by means of their lead salts, and dry distillation of their calcium salts (Ciamician, *B* 14, 1053)

Properties—Liquid, smelling like chloroform. Turns brown in air. Resinified by HCl, but more slowly than pyrrole, gives a white pp. with $HgCl_2$. Forms a potassium compound C_4H_5NMeK , which combines with CO_2 at 200°, forming (a) methyl pyrrole carboxylic acid. The potassium compound reacts with chloroform, yielding chloro methyl pyridine. Potash fusion yields pyrrole (a) carboxylic acid. Oxidising agents form HOAc, ammonia, and CO_2 . HCl passed into its ethereal solution ppts di-methyl di-pyrrole hydrochloride ($C_4H_5N_2$), HCl, whence cold dilute H_2SO_4 forms di-methyl indole (c 275°), of which the picrate melts at 156° (Dennstedt, *B* 21, 3439). On heating with phthalic anhydride and HOAc at 240° there is formed a yellow compound $C_8H_5NO_2$ [157°], which appears to be the anhydride of an acid, which may be obtained from it by boiling with dilute KOH aq, acidifying, and extracting with ether. This acid forms colourless crystals, and melts at 170°–172° (Ciamician a Zimmermann, *B* 19, 2203)

(b) **Methylpyrrole** C_4H_5NMe (143°) Occurs in bone oil (v. supra) Liquid. Resembles the (a) isomeride in its reactions. Potash fusion yields pyrrole (b) carboxylic acid. CO_2 acting on its potassium compound forms (b) methyl pyrrole carboxylic acid. HCl passed into its ethereal solution forms the hydrochloride of (b)-dimethylpyrrole ($C_4H_5N_2$), HCl, and when dilute H_2SO_4 is allowed to act for 24 hours on this hydrochloride there is formed a di-methyl indole of which the picrate melts at 149° (Dennstedt, *B* 21, 3439). On heating (b) methyl pyrrole (5 g) dissolved in

HOAc (3 vols) with phthalic anhydride (10 g) for 5 hours at 200° there is formed a compound $C_{12}H_9NO_2$ [215°], which may be purified by crystallisation from alcohol and sublimation, and obtained as lemon yellow needles, insol water (Ciamician a Dennstedt, *B* 17, 2957, *I* 9, 2201) Dilute KOH converts the compound $C_4H_5NO_2$ into an acid $C_4H_5NO_3$, which forms colourless crystals (from ether), and melts at 159°

Acetyl derivative of methyl pyrrole C_4H_5NMeAc (197°) Formed, together with methyl pyrrol methyl ketone, by boiling methylpyrrole with NaOAc and Ac_2O (Ciamician a Silber, *B* 19, 1409) Liquid. Volatile with steam, almost insol water. Easily saponified by alkalis

Methylpyrrole-tetrahydride C_4H_9N v. s. $CH_2CHMe \begin{smallmatrix} CH_2CH_2 \\ CH_2CH_2 \end{smallmatrix} NH$ **Methyl pyrrolidine** (97° at

737 mm iV) Formed by reduction of oxy methyl pyrrole dihydride (from γ amino valeric acid) in boiling amyl alcohol with metallic sodium (Tafel, *B* 20, 250) Colourless mobile liquid. Very volatile at the ordinary temperature. Boiling MeI and MeOH form crystalline $C_4H_5NMeNHMeI$

Salts (Tafel a Neugebauer, *B* 22, 1865) — B^*HCl [210°–220°] Colourless prisms — $B^*H_2PtCl_4$ aq golden needles (from hot water) — $B^*H_2AuCl_4$ [140°–144°] Golden crystals, v. sol water and alcohol — $B^*H_2C_2O_4$ [165°–168°] Small needles

Nitrosamine $C_4H_5(NO)N$ Yellow oil. **Methyl pyrrole tetrahydride** $NH \begin{smallmatrix} CH_2CHMe \\ CH_2CH_2 \end{smallmatrix}$ (104°) SG $\frac{d}{4}$ 8654 This base is formed by the dry distillation of the hydrochloride of methyl tetramethylene-diamine $NH_2CH_2CH_2CH_2CH_2NH_2$ (Oldach, *B* 20, 1657) Liquid, fuming in the air, and smelling like piperidine

Salts—Hydrochloride very deliquescent — $B^*H_2PtCl_4$ [194°] Long prisms — $B^*H_2AuCl_4$ [170°] Minute tables, v. sol water — $B^*HI(BI)_2$ — $B^*C_6H_5(NO_2)_2OH$ [105°]

Nitrosamine $C_4H_5(NO)N$ (224°)

Di-methylpyrrole $C_4H_5N_2$ v. s. $NH \begin{smallmatrix} CMeCH \\ CMeCH \end{smallmatrix}$ (165°) Occurs in bone oil (Weidel a Ciamician, *B* 13, 78)

Formation—1 By the action of alcoholic potash at 150°–160° on its dicarboxylic ether, which is formed by reduction of a mixture of acetoacetic and isonitroso acetoacetic ethers with zinc dust and acetic acid (Knorr, *B* 17, 1638) — 2 By heating its mono (or di) carboxylic acid, obtained from di acetyl succinic ether (Knorr, *B* 18, 1565) — 3 By heating acetyl acetone $CH_3COCH_2CH_2COCH_3$ with a slight excess of alcoholic NH_3 for an hour at 150° (Paal, *B* 18, 2254)

Properties—Colourless liquid with unpleasant odour, almost insol water, v. s. sol alcohol and ether. Very volatile with steam. Gives a white pp. with $HgCl_2$. Slowly resinified by HCl. Its vapour colours pine-wood moistened with HCl red. Bromine-water gives a white pp. $FeCl_3$ gives a brownish red colouration. H_2SO_4 added to its acetic acid solution mixed with phenanthraquinone gives a deep brownish red colour. H_2SO_4 and isatin give a green colour, changing on warming to brownish red. Phenyl glyoxylic

acid and sulphuric acid also give a brownish-red colour. Hydroxylamine acting on its alcoholic solution forms the di-oxim of acetylacetone $\text{CH}_3\text{C}(\text{NOH})\text{CH}_2\text{CH}_2\text{C}(\text{NOH})\text{CH}_3$ [186 5°] (Ciamician & Zanetti, *B* 22, 3177).

Acetyl derivative $\text{C}_6\text{H}_5\text{MeNAc}$ Liquid, not solidified at -20° , v sl sol water. Saponified by potash. Does not reduce AgNO_3 in very dilute ammoniacal alcoholic solution.

Di-methyl-pyrrole tetrahydride $\text{C}_6\text{H}_{11}\text{N}$ $\text{NH} \begin{smallmatrix} \text{CHMeCH}_2 \\ \text{CHMeCH}_2 \end{smallmatrix}$ (107° 1 V) Formed by distilling the hydrochloride of hexylene diamine $\text{NH}_2\text{CHMeCH}_2\text{CH}_2\text{CHMeNH}_2$ obtained by reducing the phenyl hydrazide of acetylacetone (Tafel, *B* 22, 1854; Tafel & Neugebauer, *B* 23, 1547). Colourless oil, smelling like piperidine, miscible with water, alcohol, and ether. Its hydrochloride crystallises in needles [188° – 190°]— $\text{B}'_2\text{H}_3\text{C}_2\text{O}_4$, small needles (from alcohol)— $\text{B}'_2\text{H}_3\text{PtCl}_6$, golden prisms.

Nitrosamine $\text{C}_6\text{H}_8\text{MeNNO}$ (135° at 60 mm) Yellow oil, lighter than water, sl sol cold water, v sol ether and alcohol. Conc H_2SO_4 dissolves it without becoming coloured, but on warming the liquid becomes yellow and gives off gas. The nitrosamine may be reduced to an only hydrazine, which on oxidation with HgO yields the tetrazone $\text{C}_4\text{H}_4\text{N}_4$ [43°].

Di-methyl-pyrrole $\text{C}_6\text{H}_8\text{N}$ $\text{NH} \begin{smallmatrix} \text{CMeCH} \\ \text{CHCMe} \end{smallmatrix}$ (171° cor) Formed by distilling its carboxylic acid (Knorr, *A* 236, 326). Liquid, with blue fluorescence and characteristic sweet odour, sl sol water, v sol alcohol, ether, and benzene. Its aqueous solution becomes red on warming with FeCl_3 .

Di-methyl-pyrrole tetrahydride $\text{C}_6\text{H}_{10}\text{MeNMe}$ (89°–92°) Formed by distilling the methyl iodide of methyl pyrrole tetrahydride with KOH (Ciamician & Magnaghi, *G* 15, 485). Liquid, miscible with water. Its hydrochloride is deliquescent.

Methyl iodide $\text{C}_6\text{H}_8\text{MeNMe}_2\text{I}$ Small needles (from alcohol), v e sol water. Ppd from its aqueous solution by addition of KOH . On distillation with KOH it yields trimethylamine and butene C_4H_8 .

Tri-methyl-pyrrole $\text{C}_6\text{H}_{11}\text{N}$ (180°–195°) Occurs in bone oil (fraction 180°–205°), from which it is isolated by conversion into the potassium compound (Ciamician & Dennstedt, *B* 14, 1340). Colourless oil. Sol acids, sl sol water. Turns brown in air. Its vapour turns pine wood moistened by HCl red. Reduces PtCl_4 . Gives a white pp with HgCl_2 . Potassium acts on it very slowly with production of the solid compound $\text{C}_6\text{H}_{10}\text{NK}$. Conc HClAq at 120° forms ammonia and a di-methyl pyridine dihydride $\text{C}_6\text{H}_{10}\text{N}$.

Tri-methyl-pyrrole $\text{C}_6\text{H}_{11}\text{N}$ (150°–165°) Prepared by heating methyl pyrrole (3 g) dissolved in MeOH (5 g) with MeI (7 g) and K_2CO_3 (3 g) for 10 hours at 140° . The product is acidified and distilled with steam (Ciamician & Anderlini, *B* 22, 656).

Tri-methyl-pyrrole $\text{C}_6\text{H}_{11}\text{N}$ $\text{NMe} \begin{smallmatrix} \text{CMeCH} \\ \text{CMeCH} \end{smallmatrix}$ (173° cor) Formed by heating its dicarboxylic acid at 260° (Knorr, *A* 236,

304). Liquid, volatile with steam; v sol alcohol and ether. Boiling aqueous FeCl_3 gives an intense red colour.

Tri-methyl-pyrrole tetra-hydride $\text{C}_6\text{H}_{11}\text{N}$ $\text{NH} \begin{smallmatrix} \text{CHMeCH}_2 \\ \text{CHMeCH}_2 \end{smallmatrix}$ (116°) at 750 mm Formed

by the action of MeI on $\text{NH} \begin{smallmatrix} \text{CHMeCH}_2 \\ \text{CHMeCH}_2 \end{smallmatrix}$ (Tafel & Neugebauer, *B* 23, 1549). Liquid, sl sol water— $\text{B}'\text{HCl}$ glittering white prisms— $\text{B}'_2\text{H}_3\text{PtCl}_6$, yellow oil, sl sol alcohol.

Methyl iodide $\text{B}'\text{MeI}$ [256°] Colourless prisms, v sol water, sl sol warm alcohol.

A tri-methyl pyrrole tetrahydride

$\text{NH} \begin{smallmatrix} \text{CH}_2\text{CMe} \\ \text{CMeCH}_2 \end{smallmatrix}$ appears to be formed by distilling oxy tri-methyl pyrrole dihydride with zinc dust (Weil, *A* 232, 213).

METHYL-PYRROLE-DISAZO COMPOUNDS v Dis azo compounds

(a) **METHYL-PYRROLE CARBOXYLIC ACID** $\text{C}_6\text{H}_8\text{NO}_2$, $\text{e C}_6\text{H}_8\text{MeN CO}_2\text{H}$ (a) *Homo pyrroline carboxylic acid* *Carbohomo-pyrroline acid* [169 5°] When crude potassium methyl pyrrole is heated in a current of CO_2 at 190° there is formed a mixture of the potassium salts of (a) and (b)-methyl pyrrole carboxylic acids. These acids may be separated by lead acetate, which ppt (b) acid only (Ciamician, *G* 11, 230, *B* 14, 1056). Colourless scales, sol water. Yields (a) methyl pyrrole on distillation with lime (b) **Methyl-pyrrole carboxylic acid** $\text{C}_6\text{H}_8\text{NO}_2$, $\text{e C}_6\text{H}_8\text{MeN CO}_2\text{H}$ [142 4°] Obtained as above, forms a minutely crystalline mass. Its Ca salt yields (b) methyl pyrrole on distillation with lime.

Methyl pyrrole carboxylic acid $\text{C}_6\text{H}_8\text{NO}_2$, $\text{e C}_6\text{H}_8\text{MeN CO}_2\text{H}$ [135°] Formed by heating its methylamide with alcoholic potash at 120° (Chichester & Bell, *B* 10, 1861, 11, 1810).

Methylamide $\text{C}_6\text{H}_8\text{MeN CO NHMe}$ [90°] Formed, together with methyl pyrrole, by heating methylamine mucate in a paraffin bath (Bell). Scales or prisms, sol water, volatile with steam.

Di-methyl-pyrrole carboxylic acid $\text{C}_6\text{H}_8\text{NO}_2$, $\text{e NH} \begin{smallmatrix} \text{CMe}=\text{CH} \\ \text{C(CO}_2\text{H)} \end{smallmatrix}$ [137°] Formed by the prolonged action of boiling aqueous KOH on tetra-methyl pyrrolyl pyrrole carboxylic acid or on tetramethylpyrrocol, which is itself obtained by heating di-methyl pyrrole dicarboxylic acid (Magnanini, *B* 22, 38, *Rend Accad Linc* [4] 4, 475). Crystals, v sl sol cold water, dissolves in warm water with partial decomposition into CO_2 and di-methyl pyrrole. Decomposes on fusion. Boiling Ac_2O converts it into tetra-methyl pyrrocol and di-methyl pyrrolyl methyl ketone. A solution of its ammonium salt gives with lead acetate a white pp, sol excess, with cupric acetate a green crystalline pp, and with FeCl_3 a dark red pulverulent pp.

Di-methyl pyrrole carboxyl derivative $\text{C}_6\text{H}_8\text{N}_2\text{O}_2$, $\text{e CHCMe} \begin{smallmatrix} \text{C CO N} \\ \text{C(CO}_2\text{H)} \end{smallmatrix} \begin{smallmatrix} \text{CMe}=\text{CMe} \\ \text{CH} \end{smallmatrix}$. **Tetra-methyl pyrrolyl-pyrrole carboxylic acid.** Formed by boiling tetra-methyl-pyrrocol with alcoholic

potash, diluting with water, and ppg with acetic acid (Magnanini, *B* 22, 35, *Rend Accad Linc* [4] 4, 468). At 145° it gives off CO₂ and leaves a residue which may be crystallised from alcohol. Boiling aqueous potash forms di methyl pyrrole carboxylic acid. On warming solutions of the salts a pp of tetra methyl pyrrocol is formed. The lead and silver salts are white pps, the ferric salt a red pp, and the cupric salt a green pp —BaA'. Tables

Methyl ether MeA' [163°] Formed from the silver salt and MeI. Monoclinic tables, $a b c = 701 \text{ } 1 \text{ } 443, \beta = 80^\circ 59'$. Insol water, sl sol benzene, v sol CHCl₃. Yields MeOH and tetramethylpyrrocol on fusion.

Di methyl pyrrole carboxylic acid C₅H₅NO₂ $\text{NH} \begin{array}{c} \text{CMe} \text{ C}(\text{CO}_2\text{H}) \\ \text{CH} = \text{CMe} \end{array}$ [183°] Obtained from its ether, which is produced by heating the mono ethyl ether of di methyl pyrrole dicarboxylic acid $\text{NH} \begin{array}{c} \text{CMe} \text{ C}(\text{CO}_2\text{Et}) \\ \text{C}(\text{CO}_2\text{H}) \text{ CMe} \end{array}$ (Knorr, *A* 236, 325). Crystalline flocculi. Split up on melting into CO₂ and di-methyl pyrrole. Boiling Ac₂O forms di methyl pyrrol methyl ketone CH₃CO C₅H₄NMe₂ [122°].

Ethyl ether EtA' [76°] (291° cor) Crystalline mass, v sol alcohol and ether. *Anilide* C₅H₄Me₂N CONHPh [80°] Formed by heating the mono anilide of di methyl pyrrole dicarboxylic acid.

Di methyl-pyrrole carboxylic acid C₅H₅NO₂ $\text{NH} \begin{array}{c} \text{CMe} \text{ CH} \\ \text{CMe} \text{ C} \text{ CO}_2\text{H} \end{array}$ Obtained by saponifying with aqueous NaOH its ether, which is formed by heating the mono ethyl ether of di methyl-pyrrole dicarboxylic acid $\text{NH} \begin{array}{c} \text{CMe} \text{ C} \text{ CO}_2\text{H} \\ \text{CMe} \text{ C} \text{ CO}_2\text{Et} \end{array}$ (Knorr, *B* 18, 1564). Slender needles. Split up at 210–213° into CO₂ and the corresponding di-methyl pyrrole.

Salts —PbA' microcrystalline pp —AgA' white pp.

Ethyl ether EtA' [118°] (290° at 731 mm) Formed as above, and also by slowly adding conc NH₄Aq to a mixture of molecular proportions of chloro acetone and acetoacetic ether. Flat prisms, volatile with steam. Sol alcohol and ether, insol water.

Di-methyl-pyrrole dicarboxylic acid C₅H₄NO₂ $\text{NH} \begin{array}{c} \text{CMe} \text{ C}(\text{CO}_2\text{H}) \\ \text{C}(\text{CO}_2\text{H}) \text{ CMe} \end{array}$

Preparation —NaNO₂ (2pts) in conc aqueous solution is added to acetoacetic ether (7pts) dissolved in HOAc. Zinc dust (25 pts) is added to the well cooled product. On adding water the di ethyl ether is deposited in needles, and this is saponified by boiling NaOHaq (Knorr, *A* 236, 317, *B* 17, 1638).

Properties —Crystalline flakes, v sol water and alcohol, less sol ether. Rapidly turns red in air. Decomposes completely at 260° into CO₂ and di-methyl pyrrole, without melting. It reduces boiling ammoniacal AgNO₃. It forms normal and acid salts which are mostly amorphous.

Mono-ethyl ether HEtA' $\text{NH} \begin{array}{c} \text{CMe} \text{ C}(\text{CO}_2\text{Et}) \\ \text{C}(\text{CO}_2\text{H}) \text{ CMe} \end{array}$ [202°]. Formed by

boiling the di-ethyl ether with alcoholic potash. Needles (from alcohol), sl sol cold alcohol, insol water. Split up on fusion into CO₂ and di methyl pyrrole carboxylic ether. Its lead and silver salts are white amorphous pps. When heated with Ac₂O in a sealed tube at 200° it yields $\text{NH} \begin{array}{c} \text{CMe} \text{ C}(\text{CO}_2\text{Et}) \\ \text{C}(\text{CO}_2\text{H}) \text{ CMe} \end{array}$ [148°], and this ether gives on saponification a crystalline acid [150°–158°], which on dry distillation is split up into CO₂ and di methyl pyrrol methyl ketone [123°] (Magnanini, *B* 21, 2865).

Di-ethyl ether C₅H₄NO₂ or Et₂A' [135°] Formed as above. White matted needles, insol water, acids, and alkalis, sol alcohol and ether. Split up by heat into CO and di methyl pyrrole. Alcoholic potash forms a pp of C₅H₄N₂KNO₄.

Mono anilide C₅H₄N₂O₂ $\text{NH} \begin{array}{c} \text{CMe} \text{ C}(\text{CONHPh}) \\ \text{C}(\text{CO}_2\text{H}) \text{ CMe} \end{array}$ Formed by boiling its ethyl ether (*v infra*) with alcoholic potash. Slender needles which soften at 180° and decompose at 198° into CO₂ and the anilide of di-methyl pyrrole carboxylic acid. Boiling dilute H₂SO₄ decomposes it in the same way.

Ethyl ether of the mono-anilide C₅H₄N₂O₂Et [216°] Formed by the action of zinc dust on a mixture of equivalent quantities of acetoacetic anilide and nitroso acetoacetic ether dissolved in HOAc (Knorr, *A* 236, 327). Crystals, sol hot alcohol and HOAc. Yields di methyl pyrrole on warming with conc H₂SO₄.

Ethyl ether of the isomeric mono anilide C₅H₄N₂O₂Et $\text{NH} \begin{array}{c} \text{CMe} \text{ C}(\text{CO}_2\text{Et}) \\ \text{C}(\text{CONHPh}) \text{ CMe} \end{array}$ [180°] Formed by the action of zinc dust on a mixture of equivalent quantities of acetoacetic ether and nitroso acetoacetic ether dissolved in HOAc (Knorr). Needles (from alcohol). Yields di methyl pyrrole on warming with H₂SO₄.

Di-anilide $\text{NH} \begin{array}{c} \text{CMe} \text{ C}(\text{CONHPh}) \\ \text{C}(\text{CONHPh}) \text{ CMe} \end{array}$ [255°] Formed by the action of zinc dust on a mixture of equivalent quantities of acetoacetic anilide and nitroso acetoacetic anilide (Knorr). Needles (from alcohol). Yields di methyl pyrrole on warming with H₂SO₄.

Anhydride C₅H₄NO₂ $\text{N} \begin{array}{c} \text{CMe} \text{ C}(\text{CO}_2\text{H}) \\ \text{G} = \text{CMe} \end{array}$ Formed by boiling di CO

methyl pyrrole dicarboxylic acid with Ac₂O (Magnanini, *B* 21, 2876). Powder, almost completely insol water, alcohol, and ether. Turns brown at 300° and at a higher temperature it splits up into di methyl pyrrole and tetra methyl pyrrocol —MgA₂ needles, m sol water —AgA' yellow amorphous sediment.

Ethyl ether of the anhydride C₅H₄EtNO₂ [270°] Formed by boiling di methyl pyrrole dicarboxylic ether with Ac₂O. White, sparingly soluble needles.

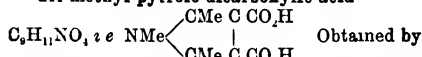
Di-methyl-pyrrole dicarboxylic acid $\text{NH} \begin{array}{c} \text{CMe} \text{ C} \text{ CO}_2\text{H} \\ \text{CMe} \text{ C} \text{ CO}_2\text{H} \end{array}$ [250°] Formed by dissolving di acetyl succinic ether in aqueous NH₃, and saponifying the resulting ether with alcoholic potash (Knorr, *B* 18, 302, 1558). Needles

(from alcohol) Decomposes at its melting point into CO₂ and (aa') di-methyl pyrrole — BaA'' small needles — CuA'' 3ag slender green needles — xAg₂A'' microcrystalline powder

Mono ethyl ether HEtA'' [227°] Slender crystals Evolves CO₂ at its melting point, giving di methyl pyrrole carboxylic acid It has acid properties, and its Cu, Co, and Ni salts form white felted needles

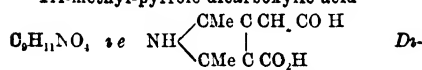
Di ethyl ether EtA'' [99°] Colourless crystals, sol alcohol, CHCl₃, and HOAc, sl sol ether, nearly insol water Has weak basic and acid properties — (C₄H₇NO, HCl), PtCl₂, Orange red crystals — C₄H₇NO, KNO₃, slender felted needles

Tri-methyl-pyrrole dicarboxylic acid



saponifying with alcoholic potash its di ethyl ether which is produced by boiling di aceto succinic ether with methylamine in glacial acetic acid solution (Knorr, B 18, 303, A 236, 304) Slender needles, insol water, sl sol ether, sol alcohol Split up at 260° into CO₂ and tri methyl pyrrole — KHA'' — NH₄HA'' — BaA' — CoA''

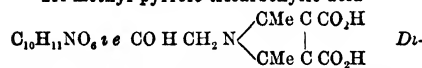
Ethyl ether EtA'' [72°] Tri-methyl-pyrrole dicarboxylic acid



methyl pyrrole carboxy acetic acid [196°] Formed by saponification by aqueous NaOH liq from its ether which is obtained by boiling aβ di acetyl glutaric ether with a solution of NH₃ in HOAc (Knorr, B 19, 48) Slender prisms Imparts a red colouration to acidified pine wood

Di ethyl ether EtA'' [110°] Glistening plates, v sol alcohol and ether, insol water

Tri-methyl-pyrrole tricarboxylic acid



methyl pyrrole dicarboxy acetic acid Obtained by saponifying its ether which is produced by boiling di acetyl succinic ether with glyccoll and HOAc (Knorr, A 236, 315) Crystalline pp, decomposed at 214° giving off CO₂ — K₂A''' — Ag HA''' crystalline pp

Di ethyl ether EtA''' [169°] Formed as above Plates, insol water, v sol alcohol, ether, and alkalis — Pb(EtA'''), prisms

METHYL-PYRROLIDINE v METHYL PYRROLE TETRAHYDRIDE

METHYL-PYRROLINE v METHYL PYRROLE and its Dihydrate

TETRA- METHYL- PYRROYL- PYRROLE CARBOXYLIC ACID v DI METHYL PYRROLE CARBOXYLIC ACID

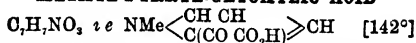
DI-METHYL PYRRYL-BENZOIC ACID v PHENYL DI-METHYL-PYRROLE CARBOXYLIC ACID

METHYL-PYRRENE-DIBENZOIC ACID v DI PHENYL-METHYL PYRROLE DICARBOXYLIC ACID

v METHYL-PYRRENE DI METHYL DI-KETONE C₈H₁₁NMe(CO Me)₂ v *Methyl di acetyl pyrrole* [184°] Formed by heating v-methyl pyrrole with acetic anhydride for 8 hrs at 250° Colourless needles V sol alcohol, ether chloro

form, benzene, and hot water (Ciamician a. Silber, B 20, 1368, G 17, 134)

METHYL PYRRYL-GLYOXYLIC ACID

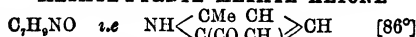


Formed by boiling methyl pyrrol methyl ketone with dilute alkaline KMnO₄ (De Varda, B 21, 2871, *Rend Accad Linc* [4] 4, 755, 758) Light yellow needles (from benzene) Bromine added to its solution in glacial acetic acid forms C₆HBr MeN CO CO₂H crystallising in yellow prisms [160°], oxidised by fuming HNO₃ to the methylamide of di bromo maleic acid — AgA' white pp

METHYL PYRRYL KETONE v PYRRYL

METHYL KFTONE

METHYL-PYRRYL METHYL KETONE



(210°) Formed by the action of AcO and NaOAc upon methyl pyrrole from bone oil (Ciamician a. Silber, B 19, 1408, 20, 2604, G 16, 352, 17, 269) Small white needles (from water), v sol alcohol, ether, and benzene Not decomposed by boiling alkalis Yields a methyl pyrrol glyoxylic acid on oxidation Bromine in CS₂ gives C₆HBr NO crystallising in needles [162°], which is oxidised by HNO₃ to di bromo maleic imide at — 18° Ammoniacal AgNO₃ gives with the ketone a white pp of C₆H₇MeAcNag

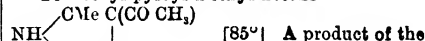
v METHYL- PYRRYL METHYL KETONE

CH₃CO C₆H₇NMe *Pseudo acetyl methyl pyrrole* (201°) Prepared by boiling v methyl pyrrole (10 g) with AcO (70 g) and NaOAc (12 g) for 12 hours (Weidel a. Crammian, B 13, 76, Ciamician a. Dennstedt, B 17, 2932, De Varda, B 21, 2972) Colourless liquid, sl sol water Not hydrolised by boiling KOHAq Reduces warm ammoniacal AgNO₃, forming a silver mirror

Di-methyl-pyrryl methyl ketone C₆H₇NO₂ i.e.

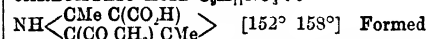
NH $\begin{array}{c} \text{CMe CH} \\ | \\ \text{C(CO CH}_3) \end{array}$ CMe [128°] Obtained by distillation of its carboxylic acid (Magnanini, *Rend Accad Linc* [4] 4, 832, B 21, 2867) Monoclinic crystals, a b c = 402.1 857, β = 78° 16' At 100 it sublimes in needles V sol hot water, v sol alcohol, benzene, and HOAc, sl sol petroleum Not affected by boiling conc KOHAq Yields a phenyl hydrazide

Di-methyl-pyrryl methyl ketone



action of AcO on the corresponding di methyl pyrrole (Dennstedt a. Zimmermann, B 19, 2195) Crystals Reduces AgNO₃ in very dilute alcoholic solution

DI-METHYL-PYRRYL METHYL KETONE CARBOXYLIC ACID C₆H₇NO₃ i.e.



by saponification of its ethyl ether, which is prepared by heating the mono ethyl ether of di methyl pyrrole dicarboxylic acid (1 pt) with Ac₂O (5 pts) at 205° in sealed tubes (Magnanini, *Rend Accad Linc* [4] 4, 830, B 21, 2865) Needles (from HOAc), almost insol water and cold alcohol, v sl sol ether, CHCl₃, and benzene Completely decomposed on fusion into CO₂ and di methyl pyrrol methyl ketone When heated with isatin and conc H₂SO₄ it gives a green

colour A solution of its ammonium salt gives pps with lead, copper, ferric, cobalt, and mercuric salts

Ethyl ether EtA' [143°] Slender needles, v. sol hot water, alcohol, and ether

DI-METHYL-o-PYRRYL-PHENOL $C_{12}H_{13}NO$

1.6 [2 1] $C_6H_4(OH)N < \begin{smallmatrix} CMe \\ CMe \end{smallmatrix} CH >$ *Oxy phenyl-di methyl pyrrole* [95°] Formed by warming acetyl acetone with *o* amido phenol in alcoholic solution (Paal, B 19, 558) White plates (from alcohol), turning red in air, sl sol water, v sol alcohol and ether Dissolves in alkalis, and is reprecipitated as needles by CO_2 —NaA' crystalline—The picrate crystallises in reddish brown plates

(a) **METHYL-PYRRYL STYRYL KETONE** $C_9H_9(CH_3)COCHCHC_6H_5$, [193°] Formed by boiling (a) methyl pyrrol methyl ketone and benzoic aldehyde with dilute caustic potash (Dennstedt a Lehne, B 22, 1918) Sulphur yellow needles (from alcohol)

(b) **Methyl-pyrrol styryl ketone** $C_{11}H_{13}NO$ $\approx C_9H_9(CH_3)COCHCHC_6H_5$, [156°–157°] Formed from (b) methyl pyrrol methyl ketone, benzoic aldehyde, and dilute caustic potash (D a L) Small light yellow plates, v sol alcohol

(aB') **Dimethyl-pyrrol styryl ketone**

$< \begin{smallmatrix} CH \\ CMe \end{smallmatrix} CMe > CCOCHCHC_6H_5$, [188°] Formed from (aB')-di-methyl-pyrrol methyl ketone, benzoic aldehyde, and KOHAq Yellow shining plates (from alcohol) (Dennstedt, B 22, 1921)

(aB') (?) **Di-methyl-pyrrol styryl ketone** $C_9H_9(CH_3)_2COCHCHC_6H_5$, [166°] Formed like the preceding Small yellow plates or needles (from boiling alcohol) (Dennstedt, B 22, 1926)

METHYL-PYRUVIC ACID v **ETHYL GLYOXYLIC ACID** Its nitrile is described as **PROPRIONYL CYANIDE**

Tri-methyl-pyruvic acid $C_6H_9O_3$ $\approx CMe_3COCO_2H$ [88°] (186°) Formed, together with a little CMe_3CO_2H , by oxidising pinacol with warm alkaline $KMnO_4$ (Glucksmann, M 10, 770) Small colourless prisms (from ether), sl sol cold water, v sol hot water, m sol ether Volatile with steam Gives a silver mirror with ammoniacal $AgNO_3$ May be reduced to $CMe_3CH(OH)CO_2H$ and oxidised to CMe_3CO_2H

Salts—NaA'—CaA' 3aq—AgA'

Phenyl-hydrazide $C_{12}H_{15}N_3O_2$ [158°] Long needles

METHYL-QUINALDINE v **DI METHYL QUINOLINE**

METHYL-QUINALDINIUM HYDRATE v **Methyl hydrate of (Py 3) METHYL-QUINOLINE**

METHYL-QUINAZOLINE DIHYDRIDE C_8H_9N , $\approx C_6H_5 < \begin{smallmatrix} CH_2NH \\ N \\ CMe \end{smallmatrix} >$ Formed by distilling the acetyl derivative of *o* amido benzyl amine (Gabriela Jansen, B 23, 2812) Colourless tenacious liquid, sol water Its solution has a bitter taste and alkaline reaction—B'HCl prisms, v sol water—B'H₂PtCl₄—Chromate reddish-yellow needles—Picrate [200°] Long needles

DI METHYL-QUINOGEN v **DI METHYL DIKETONE**, *Reaction 6*

METHYL-QUININE v **QUININE**

METHYL-(pseudo)-QUINISATIN $C_{16}H_{19}NO$

$\approx C_6H_5 < \begin{smallmatrix} CO \\ NMe \end{smallmatrix} CO > (?)$ [c 120°–122°]. Formed

by oxidation of (Py 1 2) di oxy (Py 4) methyl-(Py 3) pseudo oxy-quinoline

$C_6H_5 < \begin{smallmatrix} C(OH) \\ NMe \end{smallmatrix} C(OH) >$ with Fe_2Cl_6 V. sol ordinary solvents

Dissolves in alkalis with a yellow colour (Friedlander a Muller, B 20, 2015)

METHYL-QUINIZABIN v **DI OXY METHYL-ANTHRAQUINONE**

METHYL QUINOLINIC ACID v **METHYL-PYRIDINE DICARBOXYLIC ACID**

(Py 1) **METHYL-QUINOLINE** $C_{10}H_9N$ \approx

$C_6H_5 < \begin{smallmatrix} CMe \\ N \\ CMe \end{smallmatrix} CH >$ *Lepidine Cincholepidine*

Mol w 143 (265° V) (K), (263° V) (D) S G \approx 1.0995, d_4^{20} 1.0862 (K)

Formation—1 Obtained, together with quinoline, by distilling cinchonine with aqueous KOH (Greville Williams, *Pr E* [3] 21, 377)—2 By distilling the hydrochloride of quinoline tetrahydride (Py 1) carboxylic acid (tetrahydrocinchonic acid) with zinc dust (Weidel, M 3, 75)—3 Formic aldehyde or methylal is mixed with acetone, the mixture is saturated with HCl, and heated with a solution of aniline in conc HCl The bases are separated from the product by potash and fractionally distilled, the equations being $CH_3O + CH_3COCH_3 = H_2O + CH_3CHCOCH_3$, and the following $CH_3CHCOCH_3 + C_6H_5NH_2 = C_{10}H_9N + H_2O + H_2$ (Beyer, *J pr* [2] 82, 127)—4 By heating oxy-(Py 1) methyl-quinoline with zinc dust (Knorr, A 236, 94)—4 By heating cinchene with HOAc at 200° (Koenigs, B 23, 2677)

Preparation—1 The fraction 250°–260° of the distillate from cinchonine and KOHAq is converted into acid sulphate, the acid sulphate of (Py 1) methyl quinoline being insoluble in alcohol, whilst the acid sulphate of quinoline is soluble (Hoogewerf a Van Dorp, R T C 2, 1) 2 Obtained in pure state by the action of KOH upon cinchonine in presence of superheated steam (Krakau, Bl [2] 45, 248)

Properties—Oil, solidifying when cooled below 0°, sl sol water, miscible with alcohol, ether, benzene, and ligroin Smells like quinoline Turns brown in air and light According to Oechsner de Coninck (Bl [2] 38, 546), two lepidines (268°) and (268°) are obtained by distilling cinchonine or brucine, and the base of higher boiling point alone solidifies in a freezing mixture

Reactions—1 *Oxidation* with $KMnO_4$ in alkaline solution gives methyl pyridine dicarboxylic acid and, finally, pyridine tricarboxylic acid (Hoogewerf a Van Dorp, B 13, 1639) Oxidation by $K_2Cr_2O_7$ and H_2SO_4 yields quinoline (Py 1) carboxylic (cinchonine) acid—2 By adding sodium amalgam to lepidine O Greville Williams (C N 37, 85) obtained a base which formed a red crystalline nitrate $C_{10}H_9N_2HNO_3$ 3 When heated with an equivalent quantity of benzoic aldehyde and $ZnCl_2$ it yields styryl-quinoline (Heymann a Königs, B 21, 1424)—4 When heated with isoamyl iodide it yields the amylo iodide (which may be extracted by

repeatedly boiling the product with water) and also an insoluble residue whence boiling alkalis liberate 'lepamine' $C_{20}H_{21}N$, (Greville Williams, *C. J.* 16, 375) Lepamine is a fragrant oil (d 275°), V D (obs and calc 10.4) Lepamine yields the salts $C_{20}H_{21}N \cdot H_2Cl_2$ (melting below 100°) and $C_{20}H_{21}N \cdot H \cdot PtCl_5$.—5 By the action of a boiling aqueous solution of caustic potash upon mixtures of the alkyl iodides of lepidine and quinoline, blue or green substances, called cyanines are obtained (*v* QUINOLINE)

Salts — $B \cdot HCl$ needles — $B \cdot H \cdot PtCl_5 \cdot 2aq$ orange-red triclinic needles [226° – 230°] (Knorr), [α 253°] (Königs) — $B \cdot H \cdot AuCl_4$ [188°] Prismatic needles Decomposed when heated for some time at 110° — $B \cdot H \cdot CdCl_2$ — $B \cdot HNO_3$ slender prisms — $B \cdot H_2SO_4$ [229°] S (alcohol) 4 at 18° , 1.1 at 78° (Krakau, *J. R.* 17, 362) — $B \cdot H_2Cr_2O_7$, yellow needles (from hot water) which decompose at 110° — $B \cdot AgNO_3$, white needles — Tartrate $B \cdot C_4H_4O_6 \cdot aq$ Crystallises from alcohol — Picrate $B \cdot C_6H_3(NO_2)_3OH$ [208°] (H & D), [213°] (Krakau) Small yellow needles (from alcoholic solutions)

Methyl iodide $B \cdot MeI$ [174°] Yellow prisms (from alcohol) (Hoogewerf & Van Dorp, *R. T. C.* 2, 41, 318)

Ethyl iodide $B \cdot EtI$ [143°] Prisms

Isoamyl iodide $B \cdot C_5H_{11}I$ [160°] Yellow prisms (Hoogewerf & Van Dorp, *R. T. C.* 3, 352)

Chloro-lepidine *v* CHLORO METHYL QUINOLINE

(Py 1)-Methyl quinoline tetrahydride

$C_{10}H_{13}N$ *vs* $C_6H_5 \cdot \begin{smallmatrix} CHMe \\ NH \end{smallmatrix} \begin{smallmatrix} CH_2 \\ CH_2 \end{smallmatrix}$ (250° – 253° i V) at 740 mm Formed by adding sodium to a boiling alcoholic solution of oxy (Py 1) methyl-quinoline (Knorr & Klotz, *B.* 19, 3300) Formed also by the action of tin and HCl on lepidine Colourless oil, with pungent odour

(Py 2) Methyl quinoline $C_6H_5 \cdot \begin{smallmatrix} CH \\ N \end{smallmatrix} \begin{smallmatrix} CH \\ CH \end{smallmatrix}$

[10° – 14°] (250° at 710 mm) Obtained by heating (Py 2) methyl quinoline (Py 3) carboxylic acid with soda lime or alone at 160° By CrO_3 and H_2SO_4 it is oxidised to quinoline (Py 2) carboxylic acid [273°] (Doebner & Miller, *B.* 17, 1716, 18, 1642) Formed also by saturating a mixture of propionic aldehyde and methylal with HCl and heating the product with aniline and conc HClAq (Miller & Kinkel, *B.* 20, 1916) Prisms or colourless liquid

Salts — $B \cdot H_2PtCl_6 \cdot 2aq$ orange-yellow needles — $B \cdot H \cdot AuCl_4$ [145°] Prismatic yellow needles, sol hot water, sl sol cold — Picrate $B \cdot C_6H_3(NO_2)_3OH$ [187°], fine yellow needles

Methyl iodide $B \cdot MeI$ [221°], long yellow needles, sparingly sol alcohol

Amyl iodide $B \cdot C_5H_{11}I$ [215°], yellow needles

(Py 2) Methyl quinoline dihydride

$C_{10}H_{11}N$ *vs* $C_6H_5 \cdot \begin{smallmatrix} CH \\ NH \end{smallmatrix} \begin{smallmatrix} CHMe \\ CH_2 \end{smallmatrix}$ Is one of the

products formed by the action of methyl iodide (*In* 2) methyl indole (methyl ketole) (Fischer & Steche, *A.* 242, 858) Oil Forms a nitrosamine.

(Py 3) Methyl quinoline $C_{10}H_9N$ *vs*

$C_6H_5 \cdot \begin{smallmatrix} CH \\ N \end{smallmatrix} \begin{smallmatrix} CH \\ CMe \end{smallmatrix}$ (2) Methyl-quinoline Quinaldine (240° at 720 mm) (D & M), (246° 5° i V) at 724 mm (P & B), (240° uncor) (Drewson) Occurs in coal tar (Jacobsen & Reimer, *B.* 16, 1084)

Formation — 1 By heating a mixture of aniline, nitrobenzene, and H_2SO_4 with ethylene glycol or paraldehyde (Doebner & Miller, *B.* 14, 2812) — 2 By adding a little NaOH to an aqueous solution of equivalent quantities of *o* amido benzoic aldehyde and acetone (Friedlander & Gohring, *B.* 16, 1835) — 3 By the reduction of *o* nitro styryl methyl ketone with $SnCl_2$ (Drewson, *B.* 16, 1953) — 4 By heating a mixture of aniline, aldol, and HCl (Doebner & Miller, *B.* 16, 2464, 17, 1699) — 5 By heating a mixture of aniline, acetal, and H_2SO_4 (D & M) — 6 By heating a mixture of aniline, nitrobenzene, lactic acid, and H_2SO_4 (Wallach & Wusten, *B.* 16, 2007) — 7 By heating crotonic aldehyde with aniline, nitrobenzene, and H_2SO_4 (Skraup, *B.* 15, 897) — 8 By distilling its carboxylic acid with lime (Beyer, *J. pr.* [2] 33, 413, Rohde, *B.* 22, 267) — 9 By heating chloro methyl quinoline (vol ii p 82) with HI in HOAc in sealed tubes at 260° (Conrad & Lumpach, *B.* 20, 955) — 10 By heating methyl indole (methyl ketole) with bromoform and alcoholic NaOH and reducing the resulting bromo methyl quinoline [78°] with P and HI at 180° (Magnanini, *B.* 20, 2610, 21, 1940, *G.* 17, 246) — 11 By heating methyl indole (methyl ketole) with conc HCl at 225° (Magnanini, *B.* 20, 2609) — 12 By heating ethyl acetanilide with $ZnCl_2$ at 225° (Pictet & Bunzl, *B.* 22, 1847)

Preparation — A mixture of aniline (2 pts), paraldehyde (3 pts), and conc HClAq (4 pts), is heated on a water bath for a few hours (Doebner & Miller, *B.* 16, 2464)

Properties — Oil, smelling like quinoline, al sol water It gives with $AgNO_3$ a compound crystallising in white needles Not affected by nitrous acid A solution of chloride of iodine in HCl forms deep yellow needles of a chloro iodide [151°] (Dittmar, *B.* 18, 1612)

Reactions — 1 Scarcely attacked even after long boiling with chromic acid mixture, but completely oxidised by CrO_3 in HOAc CrO_3 and H_2SO_4 yield quinoline (Py 3) carboxylic acid — 2 $KMnO_4$ oxidises it to acetyl *o* amido benzoic acid (Doebner & Miller, *B.* 15, 3075) — 3 Cold conc HNO_3 forms (B 3) and (B 4) nitro (Py 3) methyl quinolines Boiling conc HNO_3 forms nitro quinoline carboxylic acid [220°] — 4 Tin and HCl reduce it to a tetrahydride — 5 On heating with sulphur H_2S is evolved and the product contains a base $C_{10}H_{11}N \cdot aq$ [162°] which crystallises from alcohol in white needles (Von Miller, *B.* 21, 1828) — 6 Combines at once with formamide forming a white mass, which crystallises from alcohol or ether in needles of $C_6H_5 \cdot MeN \cdot C(OH) \cdot NH_2$ [76°], and is decomposed by water (Cleva, *B.* 20, 76) — 7 Combines with aldehydes, forming aldol like products which readily split off water (Von Miller, *B.* 20, 2041) Thus paraldehyde when heated with quinaldine for 5 hours at 210° yields (Py 3) allyl quinoline (249° – 253°) (Eisele, *B.* 20, 2043), isobutyric aldehyde with $ZnCl_2$,

forms $C_{11}H_{11}NO$ [98°] (Brünner, *B* 20, 2041); *p* nitro benzaldehyde at 120° unites forming $C_8H_7NCH_2CH(OH)C_6H_4NO_2$ [160°] which when heated with Ac_2O loses water and becomes *p* nitro styryl quinoline (Bulach, *B* 20, 2046, *B* 22, 285) In like manner, when quinaldine (1 mol) is heated with furfuraldehyde (1 mol) and a little $ZnCl_2$ for 2 hours at 100° the product is $C_8H_7NCHCH_2C_4H_3O$ which crystallises from ligroin in tufts of needles (Srpek, *B* 20, 2044) — 8 When heated with *phthalic anhydride* and $ZnCl_2$ at 200° it yields 'quinoline yellow' $C_8H_7NCHC_2O_2C_6H_4$ which crystallises from alcohol in golden needles [235°], may be sublimed, and is insol water, v sl sol ether, m sol boiling alcohol, v sol $HOAc$ It dyes silk and wool yellow (Jacobsen & Reimer, *B* 16, 1082)

Salts — The nitrate and hydrochloride are easily soluble — $B'H_2PtCl_6$ [226°] (F & G), [229°] (P & B), [230°] (Beyer) Sparingly soluble orange prisms — *Aurochloride* yellow crystalline pp — $B'HHgCl_2$ [165.5°] White needles — $B'HI$ [186°] V sol water — $B'HSO_4$ [213°] Deliquescent prisms (Hoogewerf & Van Dorp, *R T C* 3, 344) — $B'H_2CrO_4$ long yellowish red needles, sol hot, sl sol cold, water — *Picrate* $B'CH(NO_2)_2OH$ [191°] Needles, sl sol water and cold alcohol

Methylo-iodide $B'MeI$ [195°] Lemon-yellow needles, v sol water, sol alcohol, insol ether Yields $(B'MeCl)_2PtCl_6$ crystallising in orange tables, $(B'MeCl)_2AuCl_3$ crystallising from hot water in lemon yellow needles, and $(B'Me)_2Cr_2O_7$ crystallising in orange plates, sl sol water, exploding at 90°

Methylo oxide $(B'Me)_2O$ Formed by adding aqueous KOH to the methylo iodide (Bernthsen & Hess, *B* 18, 32, Moller, *A* 242, 302) Yellow amorphous flakes, sl sol water, v sol alcohol and ether Easily decomposes, becoming red *HIAq* converts it into $B'MeI$, while *HCl* yields $B'MeCl$

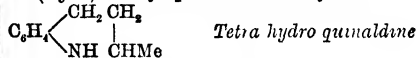
Ethylo iodide $B'EtI$ *Ethyl quinaldinum iodide* [234°] Prepared by heating quinaldine with *EtI* at 100° Yellow prisms (from alcohol), decomposed on fusion (Hoogewerf & Van Dorp, *R T C* 3, 345, 350, Spaltholz, *B* 16, 1851) Aqueous KOH converts it into the flocculent oxide $(B'Et)_2O$ which readily changes on exposure to air to a crimson resin The oxide yields the following salts $B'EtAuCl_4$ golden needles — $(B'EtCl)_2PtCl_6$ ruby red prisms, sol hot water — $(B'Et)_2Cr_2O_7$ Reddish yellow needles, exploding at 100° On heating quinoline ethylo iodide (2 pts) with quinaldine ethylo iodide (1 pt) and aqueous KOH diethyl isocyanine iodide is formed, which crystallises from its crimson alcoholic solution in lustrous green prisms (v QUINOLINE)

Propylo iodide $B'PrI$ [167°] Obtained by heating its components on the water bath (Moller, *A* 242, 306) Small greenish yellow prisms (from alcohol), sl sol cold, v sol hot, alcohol and water Gives with aqueous KOH a yellow amorphous oxide $(B'Pr)_2O$, sol alcohol and ether It yields the salts — $(B'PrCl)_2PtCl_6$ crystallising in orange plates, $B'PrAuCl_4$ crystallising in canary yellow needles, v sl sol cold water, and $(B'Pr)_2Cr_2O_7$ crystallising in brownish red prisms

Isobutylo-iodide $B'CH_3I$ [172°] Obtained by cohobating its components at 115° (M). Straw yellow plates, sol hot alcohol Yields an unstable oxide which on heating changes to a violet colouring matter

Isoamyllo-iodide $B'C_4H_9I$ [175°] Formed by cohobating its components for 12 hours at 145° Small yellow prisms, v sol water and hot alcohol Converted by aqueous KOH into an oxide which condenses to a crimson dye on heating

(Py 3)-Methyl-quinoline tetrahydride

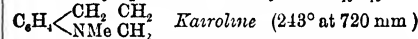


[248° at 710 mm] Formed by reducing $CH_3COCH_2CH_2C_6H_4NO_2$ [1.2] (Jackson, *B* 14, 890) It is also a product of the action of *HCl* on a mixture of aldehyde and aniline (Doebner & Miller, *B* 17, 1698) Prepared by reducing quinaldine with tin and *HCl* (Doebner & Miller, *B* 16, 2467) Colourless liquid with pleasant smell, sl sol water, v sol alcohol, ether, and benzene $FeCl_3$ gives a blood red colour to aqueous solutions of its salts, CrO_3 acts in like manner Nitrous acid produces an oily nitrosamine and a crystalline nitro nitrosamine [152°] (Moller, *A* 242, 314) Heated with benzoic chloride it gives a green dye stuff — $B'H PtCl_6$ yellow needles

v Methyl quinoline dihydride $C_{10}H_{11}N$ & c

$C_6H_7 \begin{array}{l} \swarrow CHCH_3 \\ | \\ \searrow NMeCH_3 \end{array}$ This base is probably formed by heating v methyl indole with *MeI* (Fischer & Steche, *B* 20, 2201)

v Methyl quinoline tetrahydride $C_{10}H_{11}N$ & c



Prepared by heating quinoline tetrahydride with *MeI* (Hoffmann & Konigs, *B* 16, 731), or by reduction of quinoline methylo iodide with tin and *HCl*, the yield by the latter process being 50 p c of the theoretical (Feer & Konigs, *B* 18, 2388) Oil Resembles quinine in physiological action, and has been used as a febrifuge Nitrous acid gives a yellowish red colouration in dilute solutions Gives a green dye stuff on heating with benzo trichloride

Salts — $B'HClaq$ [244°] Prepared by heating quinoline tetrahydride with *MeOH* and conc *HClAq* at 160° (Ostermayer, *B* 18, 595) Crystallises from alcohol or chloroform in tetrahedra, v sol alcohol, sl sol ether — $B'HCuCl$ [86°] Large yellow needles — $B'HPtCl_6$ [177°] Red crystalline pp — *Picrate* [125°] Long yellow needles

Nitroso derivative $C_{10}H_{12}(NO)N$ Green plates, v sol alcohol, ether, and benzene, sl sol ligroin Does not give Liebermann's reaction Not affected by boiling *NaOHAc*

Methylo hydroxide $*B'MeOH$ Strong alkaline base, sol water

Ethylo-iodide $C_6H_9NMeEtI$ [179°] Formed from ethyl quinoline tetrahydride and *MeI* (Claus & Stegehlitz, *B* 17, 1331) Crystals Not affected by *KOHAc* With *Ag_2O* it gives a crystalline base V sol water, insol ether — $(C_6H_7MeEtCl)_2PtCl_6$

References — AMIDO METHYL HYDRO QUINOLINE, NITRO METHYL QUINOLINE TETRAHYDRIDE, and OXY METHYL QUINOLINE TETRAHYDRIDE

(B 1) Methyl quinoline $C_{10}H_9N$ s.s.
 $\begin{array}{c} CH\ CMe\ C\ CH\ CH \\ CH\ CH\ CN\ CH \end{array}$ (251°) This is probably the constitution of the base which is formed, together with (B 3) methyl quinoline, by heating *m* toluidine, *o* nitro phenol, glycerin, and H_2SO_4 , together (Skraup, a Brunner, *M* 7, 139) Its acid sulphate is more soluble in alcohol than that of the (B 3) isomeride The platinum chloride melts at 224°, the picrate at 198° The picrate is much less soluble than that of the accompanying isomeride The chromate is v sol water

Derivative — v CHLORO (B 1) METHYL QUINOLINE, which may, however, perhaps be CHLORO (B 3) METHYL QUINOLINE

(B 2) — Methyl - quinoline $C_{10}H_9N$ s.s.
 $\begin{array}{c} (Me\ CH\ C\ CH\ CH \\ CH\ CH\ CN\ CH \end{array}$ *p* Toluquinoline (258°) at 745 mm SG $\frac{1}{4}$ 10815, $\frac{2}{3}$ 10681 Formed by heating a mixture of *p* toluidine, *p* nitro toluene, glycerin, and H_2SO_4 (Skraup, *M* 2, 158) Liquid, smelling like quinoline Yields pyridine di carboxylic (quinolinic) acid on oxidation with $KMnO_4$ HCl forms chloro oxy methyl quinoline $C_9H_8Me \begin{array}{c} CH\ CH \\ \diagup\ NCl\ CO \end{array}$

Salts — $B'HCl \frac{1}{2} aq$ slender needles, v sol water and alcohol — $B'H\ PtCl_2 \cdot 2aq$ slender yellow prisms — $B'HI$ [186°] (Moller, *A* 242, 307) — $B'H_2SO_4$ aq Prisms, v e sol water — Picrate $B'CO_2H(NO_2)_2OH$ [229°] Yellow powder

Chloro iodide $C_{10}H_8NClI$ [c 170°], easily soluble in alcohol The hydrochloride $C_{10}H_8NClI \cdot HCl$ is formed by adding a solution of chloride of iodine in HCl to a solution of (B 2) methyl quinoline It forms easily soluble reddish-yellow needles [c 112°], blackened by NH_3 (Dittmar, *B* 18, 1612)

Methylo iodide $B'MeI$ Slender yellowish prisms When heated with (Py 1) methyl-quinoline methylo iodide, and aqueous KOH it forms di methyl methylo cyanine $C_{22}H_{18}N_2I_2aq$, which crystallises from alcohol in violet blue needles, sl sol water, melting (when anhydrous) at 270°–277° (Hoogewerff a Van Dorp, *R T C* 3, 342)

Derivatives — V TRI CHLORO (B 2) METHYL-QUINOLINE, CHLORO OXY (B 2) METHYL-QUINOLINE, DI CHLORO DI NITRO OXY (B 2) METHYL-QUINOLINE, and OXY (B 2) METHYL-QUINOLINE

(B 3) Methyl - quinoline $C_{10}H_9N$ s.s.
 $\begin{array}{c} CH\ CH\ C\ CH\ CH \\ CMe\ CH\ CN\ CH \end{array}$ *m* Toluquinoline (250° uncor), 259 7° cor at 747 mm) SG $\frac{1}{4}$ 10839, $\frac{2}{3}$ 10722 Formed, together with smaller quantities of an isomeride, probably (B 1) methyl-quinoline, by heating a mixture of *m* toluidine, *m* nitro toluene, glycerin, and H_2SO_4 (Skraup, *M* 3, 381, *B* 15, 893), or of *m* toluidine, *o* nitrophenol, glycerin, and H_2SO_4 (Skraup a Brunner, *M* 7, 139) The two bases may be separated by crystallising their acid sulphates from alcohol, in which the sulphate of the base here described is the less soluble Yellowish, highly refractive liquid, not solid at –20° Yields on oxidation quinoline (B 3) carboxylic acid [247°]

Salts — Hydrochloride large transparent prisms — $B'H \cdot PtCl_2 \cdot 2aq$ [224°] Orange prisms — $B'H_2SO_4$ — $B'_2(H_2SO_4)_2 \cdot 2aq$ — Chromate [89°]: v. sl. sol water — Picrate. [287°].

Methylo iodide $B'MeI \frac{1}{2} aq$ golden needles

(B 4) — Methyl - quinoline $C_{10}H_9N$ s.s.
 $\begin{array}{c} CH\ CH\ C\ CH\ CH \\ CH\ CMe\ CN\ CH \end{array}$ *o*-Toluquinoline (248° cor) at 751 mm SG $\frac{1}{4}$ 1085, $\frac{2}{3}$ 1073 Formed by heating *o* toluidine with glycerin, *o* nitro toluene, and H_2SO_4 (Skraup, *M* 2, 153) Strongly refractive liquid, smelling like quinoline, not solidified in a mixture of solid CO_2 and ether, v sl sol water, sol alcohol and ether Oxidised by $KMnO_4$ to pyridine dicarboxylic (quinolinic) acid

Salts — $B'HCl \frac{1}{2} aq$ very soluble transparent prisms — $B'_2H_2PtCl_2 \cdot 2aq$ orange prisms — $B'H_2SO_4$ prisms, sl sol cold, v sol hot, water — $B'_2C_2H_5(NO_2)_2OH$ [200°] Pale yellow leaflets

Methylo iodide $B'MeI$ Crystalline powder

(B 4) — Methyl - quinoline tetrahydride $C_{10}H_{11}N$ Oil Obtained by reducing (B 4) methyl quinoline (Ziegler, *B* 21, 866)

Nitrosamine C_9H_9MeNO Formed by treating (B 4) methyl quinoline tetrahydride with $NaNO_2$ and $HOAc$ (Ziegler) Oil Slowly converted by alcoholic HCl into an isomeric nitroso derivative, which crystallises from benzene in lustrous steel blue crystals [140°], and may be reduced to an amido (B 4) methyl quinoline tetrahydride, giving a hydrochloride melting at 166°

(B 2, 4) Di-methyl-quinoline $C_{11}H_{11}N$ s.s.
 $\begin{array}{c} CMe\ CH \\ | \\ C_9H_9N \end{array}$ (268° cor) SG $\frac{1}{4}$ 10665

Formed by heating (1, 3, 4) *m* xylidine with glycerin, nitrobenzene, and H_2SO_4 (Behrend, *B* 17, 2716) Colourless oil Yields a sulphonic acid melting at 166° — $B'H_2PtCl_6$ yellow needles — $B'H_2SO_4$ minute needles Its bichromate forms long yellow needles

(B 1, 2) or (B 2, 3) Di methyl-quinoline
 $\begin{array}{c} CMe\ CMe \\ | \qquad \qquad | \\ C_9H_9N \text{ or } C_9H_9N \end{array}$ (274°)

Obtained by heating a mixture of *o* xylidine $C_8H_7Me(NH_2)$ [1 2 4], nitrobenzene, glycerin, and H_2SO_4 (Berend, *B* 17, 1149) — $B'_2H_2PtCl_6 \cdot aq$ minute yellow needles — $B'H_2SO_4$ aq glistening prisms — $B'H \cdot Cr_2O_3$ slightly soluble orange prisms

(B 1, 4) Di-methyl-quinoline $C_{11}H_{11}N$ s.s.
 $\begin{array}{c} CH\ CMe \\ | \qquad \qquad | \\ C_9H_9N \end{array}$ [5°] (265°) at 736 mm SG

$\frac{1}{4}$ 10752 (B), $\frac{2}{3}$ 1070 Formed by warming the sulphate of *p* xylidine $C_8H_7Me(NH_2)$ [1 4 2] with nitrobenzene and glycerin (Berend, *B* 18, 3165, Lellmann a Alt, *A* 237, 308) Oxidised by dilute HNO_3 at 170° to (B 4) methyl quinoline (B 1) carboxylic acid — $B'_2H_2PtCl_6$ yellow needles — $B'H_2Cr_2O_3$ [149°] Orange needles

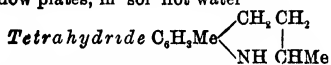
Tetra hydride $C_{11}H_{13}N$ (271°) Formed by reduction with zinc and HCl (Berend, *B* 18, 3165) Pleasant smelling liquid — $B'HCl$ slender needles (from alcohol) or six sided plates (from water)

(B 4, Py 8) Di-methyl-quinoline $C_{11}H_{11}N$ s.s.
 $\begin{array}{c} CH\ CH\ C\ CH\ CH \\ | \qquad \qquad | \\ CH_3CMe\ CN\ CMe \end{array}$ *o* Methyl quinoline s.s.

(252°) Formed by heating a mixture of *o* toluidine, paraldehyde, and HClAq at 100° (Doebner & Miller, *B* 16, 2469) Formed also by heating its carboxylic acid with KOH (Panajotoff, *B* 20, 40) Colourless liquid, v sol alcohol and ether, sl sol water Volatile with steam — *Platino chloride* $B'H_2PtCl_5 \cdot 2aq$ dark yellow needles — $B'_2H_2Cr_2O_7$, orange yellow needles, sol hot, sl sol cold, water On oxidation with chromic acid it yields an acid $C_{11}H_9NO_2$ identical with that obtained from *o* amido benzoic acid and aldehyde (Von Miller & R Meyer, *B* 23, 2260)

Methylo iodide $B'MeI$ [221°] Prepared by heating the base with MeI at 100° (Möller, *A* 242, 309) Lemon yellow needles (from indole) Alkalis liberate an unstable base, which on heating yields the original *o* methyl quin aldimine It also yields the salts $(B'MeCl)_2PtCl_5$, crystallising in yellow needles, v sl sol cold water, $B'MeAuCl_4$, and $(B'Me)_2Cr_2O_7$, crystallising from water in orange plates

Ethylo iodide $B'EtI$ [229°] From the base and EtI at 140° (Möller) Lemon yellow needles (from alcohol) Yields the salts $(B'Et)_2PtCl_5$, crystallising in orange yellow needles, and $B'EtAuCl_4$, crystallising in lemon-yellow plates, m sol hot water



Tetrahydro methyl quinaldine (261°) Formed by reducing (*B* 4, *Py* 3) di methyl quinoline (Doebner & Miller, *B* 16, 2469) Colourless liquid Gives a red colouration with $FeCl_3$ — $B'_2H_2PtCl_5$, concentric red needles

(*B* 3, *Py* 3) Di-methyl-quinoline $C_{11}H_{11}N$ + e
CH CH C CH CH

m *Methyl quinaldine*
CMe CH C N CMe

[61°] (264°) Formed by heating a mixture of *m* toluidine, paraldehyde, and HCl (Doebner & Miller, *B* 16, 2471) Long colourless needles v sol alcohol, ether, and benzene, sl sol water Oxidised by chromic acid to methyl quinoline *m*-carboxylic acid [284°] (Miller & Rast, *B* 23, 2263, 2493) — $B'_2H_2Cr_2O_7$, thick orange needles, nearly insol cold water — $B'_2H_2PtCl_5$, small yellow needles

(*B* 2, *Py* 3) Di-methyl-quinoline

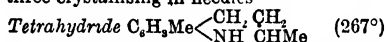
CMe CH C CH CH

p *Methyl quinaldine* [60°]
CH CH C N CMe

(*D* a M), [55°] (Jacobsen & Reimer, *B* 16, 2603) (266°) (*D* a M), (260°) (*J* a R) Formed by heating a mixture of *p* toluidine, paraldehyde, and HCl (Doebner & Miller, *B* 16, 2470) Large colourless trimetric prisms, smell like anise seed V sol alcohol, ether, and benzene, sl sol hot water Oxidised by chromic acid to methyl quinoline *p* carboxylic acid [256°] (Miller & Schunck, *B* 23, 2263) Heated with phthalic anhydride and $ZnCl_2$ at 200° it yields methyl-quinophthalone or methyl-quinoline yellow $C_{11}H_9N C_6O_5 C_6H_5$, [203°] which crystallises from alcohol in golden needles (*J* a R) — $B'_2H_2PtCl_5$, nearly colourless slender needles, sl sol. hot water — $B'_2H_2Cr_2O_7$, long yellow needles, m sol hot water — The hydrochloride, nitrate, sulphate, and acetate, are easily soluble crystalline salts

Methylo iodide $B'MeI$. [237°] Formed

from the base and MeI in the cold (Möller, *A* 242, 311) Lemon yellow needles, sl sol cold alcohol, insol ether, v sol water Yields the salts $(B'Me)_2PtCl_5$, $B'MeAuCl_4$, and $(B'Me)_2Cr_2O_7$, all three crystallising in needles



Formed by reducing the base with tin and HCl (Doebner & Miller, *B* 16, 2471) Liquid, sl sol water, v sol alcohol and ether Its aqueous solution is coloured blood red by $FeCl_3$

(*B* 4, *Py* 1) Di-methyl-quinoline

CH CH C CMe CH

(274° i V) Formed by
CH CMe C N CH

distilling oxy (*B* 4, *Py* 1) di methyl quinoline with zinc dust (Knorr, *A* 245, 369) Light yellow oil with pungent taste and smell, v sol alcohol and ether, almost insol water — $B'_2H_2PtCl_5 \cdot aq$ [220°] — $B'HAuCl_4$ [181°]

(*B* 3, *Py* 1) Di-methyl-quinoline

CH CH C CMe CH

(c 283° i V) Formed

CMe CH C N CH

by distilling (*Py* 3) oxy (*B* 3, *Py* 1) di methyl quinoline with zinc dust (Knorr, *A* 245, 371) Oil, volatile with steam — $B'_2H_2PtCl_5 \cdot 2aq$ [227°] Reddish brown prisms

(*B* 2, *Py* 1) Di methyl quinoline

CMe CH C CMe CH

CH CH C N CH

(γ) *Methyl p toluquinoline*

(280° i V) Formed by distilling (*Py* 3) oxy

(*B* 2, *Py* 1) di methyl quinoline with zinc dust (Knorr, *A* 245, 366) Pungent oil, v sol alcohol and ether Oxidised by chromic acid to a methyl quinoline carboxylic acid (Von Miller & Daniel, *B* 23, 2261) — $B'_2H_2PtCl_5 \cdot 2aq$ needles, decomposing at 231° — $B'HAuCl_4$ needles, decomposing near 192° — $B'_2H_2Cr_2O_7$ needles, decomposing near 150° — $B'_2C_6H_5(NO_2)_3OH$ needles, decomposing near 230°

(*Py* 2, 3) Di-methyl-quinoline

C₆H₅ < CH CMe

N CMe [66°] (261° uncor) at 729 mm

Formed by adding tiglic aldehyde (1 mol) to a mixture of aniline hydrochloride (1 mols) and HCl heated to 85° (Rohde, *B* 20, 1912, 22, 268) Formed also in small quantity by heating a mixture of acetic and propionic aldehydes with aniline and HClAq at 100° Tables or plates, v sl sol water, v sol ether, v e sol alcohol Chromic acid mixture oxidises it to (*Py* 3)-methyl quinoline (*Py* 2) carboxylic acid On condensation with benzil, by heating at 100° for 24 hours, it yields a base [176°], of which the solutions of the salts exhibit green fluorescence This base changes on heating for some time at 180° to a polymeride [240°]

Salts — $B'HCl \cdot 2aq$ radiating needles, v e sol water and alcohol — $B'_2H_2PtCl_5 \cdot 2aq$ orange needles, turning black at 280° — $B'HO$, prisms, v e sol water and alcohol — $B'_2H_2SO_4 \cdot aq$ [235°] slender radiating needles, v e sol water, m sol alcohol — $B'_2H_2Cr_2O_7$, long orange needles, m sol hot water, turns brown at 150° *Picrate*: [225°]

Methylo iodide $B'MeI \cdot aq$ [218°] Sick-like groups of needles (from alcohol), m sol. water and alcohol

(*Py* 1, 2) Di-methyl-quinoline

C₆H₅ < CMe CMe

N CH [68°] (290° i V) at 737 mm.

formed by distilling (Py 8, 2, 1) oxy di methyl-quinoline with zinc-dust (Knorr, A 245, 362). On oxidation by chromic acid mixture it yields Py 2)-methyl quinoline (Py 1)-carboxylic acid (Santz, B 23, 2257) — $B'H_2PtCl_4$ 2aq [240°] — $3'HauCl_4$ [177°] Plates — $B'H_2Cr_2O_7$, red needles — Picrate $B'O_4H_4(NO_2)_3OH$ [c 205°] Yellow needles

Methylo-iodide $B'MeI$ [191°] Light-yellow needles (from alcohol)

(Py 1, 3) Di-methyl-quinoline

$C_8H_8 \begin{smallmatrix} < CMe & CH \\ & N & CMe \end{smallmatrix}$ (264° uncor) SG 1.061

Formation — 1 By saturating a mixture of paraldehyde (120 g) and acetone (200 g) with gaseous HCl, allowing the mixture to stand for a day or two and then pouring it slowly into a solution of aniline (200 g) in conc HClAq (400 g) in this reaction the ketone $CH_3CHCHCOCH_3$ is doubtless first formed (O Beyer, J pr [2] 32, 125, 33, 401) — 2 Together with methane, by heating acetone with aniline hydrochloride at 180°, the yield being 3 p.c. of the theoretical (Riehm, A 238, 3, B 18, 3296) — 3 Together with the bases C_8H_8N CMe₂ (228°), V D 66 4 (calc 36 5), and $C_8H_8N(?)$ (275°), by heating aniline with acetone and a dehydrating agent such as $ZnCl_2$ or P_2O_5 — 4 By heating aniline with mesityl oxide at 180° (Engler a Riehm, B 18, 2245) — 5 By heating acetanilide with acetone, the yield being 8 p.c. of the theoretical — 6 From acetone and benzanilide — 7 By the action of NaOHAq on a mixture of o amido acetophenone and acetone (O Fischer, B 19, 1037) — 8 By dissolving the anilide of methylene di methyl diketone (acetyl acetone) in conc H_2SO_4 , warming, diluting with cold water, and neutralising with NH_3 (Combes, C R 106, 142)

Properties — Oil, with sweetish smell, somewhat like quinoline, volatile with steam. Not affected by nitrous acid

Reactions — 1 CrO_3 oxidises it to (Py 3)-methyl quinoline (Py 1) carboxylic acid — 2 By heating with phthalic anhydride and a little $ZnCl_2$ at 200° it is converted into a 'phthalone' $C_8H_8N C_2O_2 C_6H_5$, which crystallises from alcohol in orange needles [238°] (Beyer, J pr [2] 33, 407)

Salts — $B'HCl$ slender, stellate flat needles (by sublimation), v sol water and alcohol — $B'H_2PtCl_4$ [229°] — $B'H_2PtCl_4$ 2aq — $B'H_2ZnCl_4$ 1½aq — $B'H_2SO_4$ [225°–228°] Concentric needles, v sol water, sl sol alcohol — $B'H_2Cr_2O_7$, [172°] Orange needles, v sol hot, v sl sol cold, water — $B'O_4H_4(NO_2)_3OH$ [190°] Needles (from acetone)

Methylo-iodide $B'MeI$ [226°]

Ethylo-iodide $B'EtI$ [314°]. Needles (from alcohol)

Di-methyl-quinoline $C_8H_8(C_2H_5Me_2N)$. [65°] (267°) at 713 mm Occurs among the products of the action of aniline on a mixture of isobutyric aldehyde, methylal, and HCl (Von Miller a Kinkel, B 20, 1937) Monoclinic prisms — $B'H_2PtCl_4$ 2aq [268°] Groups of needles (from HClAq) — $B'O_4H_4(NO_2)_3OH$ [220°] Aggregates of tables (from hot alcohol)

(Py 1, 4) Di-methyl-quinoline tetrahydride

$C_8H_8 \begin{smallmatrix} < CHMe & CH_2 \\ & NMe & CH_2 \end{smallmatrix}$ (255° i.v.). Formed by adding Na to a boiling alcoholic solution of oxy-

(Py 1, 4)-di-methyl-quinoline (methyl-lepidone) (Knorr a Klotz, B 19, 3302) Colourless liquid

(Py 3 4) Di-methyl-quinoline tetrahydride

$C_8H_8 \begin{smallmatrix} < CH_2 & CH_2 \\ & NMe & CHMe \end{smallmatrix}$ (248°) at 710 mm Formed by heating (Py 3) methyl-quinoline tetrahydride with MeI (Doeberner a Miller, B 16, 2468) Formed also by reducing the methylo iodide of (Py 3) methyl quinoline with tin and HCl (Feer a Königs, B 18, 2388) Colourless liquid On heating with benzotrichloride it gives a green dye stuff — $B'H_2PtCl_4$ sparingly soluble red granular pp

Methylo-iodide $B'MeI$. [205°] Colourless needles, v e sol water and hot alcohol, insol ether (Moller, A 242, 318) Not acted on by alkalis, but converted by moist Ag_2O into the ammonium base, which is split up by distillation into MeOH and (Py 3, 4) di-methyl quinoline Yields the salts $(B'Me)_2PtCl_4$ crystallising from water in small brick-red crystals, $B'MeAuCl_4$ crystallising in lemon yellow needles, and $(B'Me)_2Cr_2O_7$ crystallising in small hexagonal plates

Di methyl-quinoline $C_{11}H_{11}N$ (245°) Occurs in commercial quinaldine (Einhorn, B 18, 3144) — $B'H_2PtCl_4$ [238°]

(Py 1, 2, 4) Tri-methyl-quinoline dihydride

$C_{12}H_{12}N$ s.e. $C_8H_8 \begin{smallmatrix} < CMe & CMe \\ & NMe & CH_2 \end{smallmatrix}$ (244° i.v.) at

746 mm Formed by heating methyl indole (methyl ketole) $C_8H_8 \begin{smallmatrix} < CH & CH \\ & NH & \end{smallmatrix}$ CMe with MeI and

a little MeOH for 20 hours at 100° (Fischer a Steche, B 20, 818, A 242, 353). Formed by heating (In 1, 2) di methyl indole with MeOH and MeI in sealed tubes at 100° (Fischer a Steche, A 242, 364, Wolff, B 21, 125) Formed also by heating indole with MeI, MeOH, and Na CO₃ at 130° (Giamcian a Zatti, B 22, 1980, Zatti a Ferrarini, B 23, 2302, Fischer a Meyer, B 23, 2631) Formed likewise by heating scatole (1 pt) with MeI (2½ pts) and MeOH (1 pt) for 12 hours at 130° (F a M) Oil, turning red in air, miscible with alcohol, ether, benzene, and chloroform Nitrous acid gives a reddish brown oil, becoming crystalline $FeCl_3$ added to its solution in HClAq ppts golden yellow plates or needles of a compound v sol. hot water, v sl sol conc HClAq

Salts — The hydrochloride is v sol water — Platinochloride [c 212°] — $B'H_2SO_4$ plates — $B'HI$ [253°] Prisms (from alcohol) — $B'O_4H_4(NO_2)_3OH$ [148°] Golden-yellow needles (from alcohol)

(Py 1, 2, 4) Tri-methyl-quinoline tetrahydride

$C_8H_8 \begin{smallmatrix} < CHMe & CHMe \\ & NMe & CH_2 \end{smallmatrix}$ (?) (239° i.v.) at 749 mm

Formed by reducing the dihydride with zinc and HClAq (Fischer a Steche, A 242, 356) Colourless liquid, sl sol water, v sol ether, alcohol, and benzene $FeCl_3$ gives a brown amorphous pp in its solution in HClAq — Picrate [162°] (F a S), [164°] (Giamcian a Zatti, B 22, 1981), yellow plates (from alcohol) — Platinochloride bright-red crystals

Methylo-iodide $B'MeI$ [251°]. Plates or needles (from alcohol), m sol water

(Py 1, 3, 4) Tri-methyl-quinoline tetrahydride $C_8H_8 \begin{smallmatrix} < CHMe & CH_2 \\ & NMe & CHMe \end{smallmatrix}$ Hydro-iodide

B'HI [215°] Formed from (*Py* 1, 3)-di-methyl-quinoline by reducing in alcoholic solution with Na and heating the product with MeI and MeOH at 100° (Fischer & Meyer, *B* 23, 2681).

(*Py* 1, 2, 4)-Tri-methyl-quinoline tetrahydride $C_8H_8 \begin{smallmatrix} CHMe \\ CHMe \\ CHMe \\ CHMe \end{smallmatrix}$. Hydro-iodide

B'HL [205°] Formed like the preceding from (*Py* 1, 2)-di-methyl-quinoline (F. A. M.). Crystalline

(*B* 1, 2, 4)-Tri-methyl-quinoline $C_{12}H_{12}N \begin{smallmatrix} CHMe \\ CHMe \\ CHMe \\ CHMe \end{smallmatrix}$ (43°) (286° uncor)

Formed by heating ψ cumidine [63°] with glycerin, nitrobenzene, and H_2SO_4 (Berend, *B* 18, 376) White prisms — $B'HNO_3$, sparingly soluble needles — $B'H_2SO_4$, white prisms — $B'_2H_2PtCl_6$ 2aq very sparingly soluble orange-red needles

(*B* 2, 4, *Py* 3) Tri-methyl-quinoline $C_{12}H_{12}N \begin{smallmatrix} CHMe \\ CHMe \\ CHMe \\ CHMe \end{smallmatrix}$ *op Di methyl quin-*

aldine [46°] (260°) at 719 mm Formed by heating *m*-xylidine with paraldehyde and conc HClAq on the water-bath (Panajotoff, *B* 20, 32) Small plates (from alcohol) or monoclinic prisms (from petroleum-ether), insol water, v e sol alcohol Readily volatile with steam Yields on oxidation (*B* 4, *Py* 8)-di-methyl-quinoline (*B* 2) carboxylic acid Forms, with chloral, a compound $C_7H_7O_2Na$, melting at 108°, and crystallising in yellowish needles, v sol alcohol

Salts — $B'HCi$ white needles — $B'_2H_2PtCl_6$ 2aq orange-yellow needles, sl sol water — $B'H_2SO_4$ aq white needles, v sol water and alcohol, sl sol ether — $B'HNO_3$, transparent triclinic prisms — $B'_2H_2Cr_2O_7$, orange yellow needles — $B'_2H_2(NO_3)_2 \cdot OH$ [185°] Yellow needles

Methylo-iodide $B'MeI$ aq Yellow needles, v sol water

Tetrahydride $C_8H_8 \begin{smallmatrix} CH_2 \\ CH_2 \\ CH_2 \\ CH_2 \end{smallmatrix}$

(200°–250°) Formed by reducing the base with tin and HCl (P) — $B'_2H_2PtCl_6$, orange plates

(*B* 1 or 3, 2, *Py* 8) Tri-methyl-quinoline $CMe CMe C CH CH$ or $CMe CH C CH CH$

$CH OH C N CMe$ or $CMe CH C N CMe$ Formed from *o*-xylidine, aldehyde, and HCl (Merz, *B* 17, 1158) Monoclinic crystals — $B'_2H_2PtCl_6$, minute needles

(*B* 2, *Py* 1, 3)-Tri-methyl-quinoline $CMe CH C Me CH$

$CH CH C N CMe$ [64°] (P), [40°] (O); (278°) (P), (281°) (O) Formed by saturating a mixture of acetone (40 g) and paraldehyde (30 g) with HCl at 0°, allowing to stand 12 hours in a freezing mixture, pouring into a solution of *p*-toluidine (65 g) in fuming HClAq (180 g) and heating on the water bath for 7 hours (Pätzinger, *J pr* [2] 88, 41) Formed also by dissolving the *p*-toluide of methylene di-methyl diketone $CH_2 CO CH_2 C(NC_2H_5)_2 CH_2$ in conc H_2SO_4 , warming, diluting, and neutralising with NH_3 (Combes, *C R* 106, 145) White needles, containing water of crystallisation which is given off in a desiccator, the anhydrous base crystallising in tables and absorbing water (1 mol) from the air Sl sol water, v sol alcohol, ether, and petroleum-ether. Volatile with steam.

It has a bitter, acid taste, and an irritating vapour

Salts — $B'HCi$ 2aq white needles, v e sol hot water and alcohol [260°] — $B'_2H_2PtCl_6$ 2aq yellow needles, v sl sol water, almost insol alcohol [220°] (O) — $B'_2H_2SO_4$ [222°] White needles or rhombohedra (containing aq), m sol cold alcohol — $B'_2H_2Cr_2O_7$, Yellowish red needles, v sl sol cold water — $B'_2H_2(NO_3)_2 \cdot OH$ [201°] Greenish-yellow needles (from hot acetone)

Methylo-iodide $B'MeI$ [226°] Needles (containing aq), sol water, alcohol, and ether

(*B* 2, *Py* 2, 3) Tri-methyl-quinoline $CMe CH C CH CMe$ [87°] (285°) Formed

from tiglic aldehyde (1 mol), *p*-toluidine (4 mols) and HCl (8 mols) (Von Miller & Ohler, *B* 23, 2268) White nodules (from ligroin), insol water, m sol ligroin, v sol ether Yields on oxidation (*Py* 2, 8) di-methyl-quinoline (*B* 2)-carboxylic acid [270°]

Salts — $B'_2H_2PtCl_6$ 2aq laminae — The chromate crystallises in orange laminae, v sl sol water — The hydrochloride and sulphate are v e sol water — Picrate [212°] Straw coloured laminae

(*B* 4, *Py* 1, 3) Tri-methyl-quinoline

$CH CH C CMe CH$ $CH CMe C N CMe$ (280°) Formed by the action of H_2SO_4 on the *o*-toluide of methylene dimethyl diketone $CH_2 CO CH_2 C(NC_2H_5)_2 CH_2$ (Combes, *C R* 106, 145) Oil — Platinichloride [257°]

(*B* 4, *Py* 3, 4)-Tri-methyl-quinoline tetrahydride $CH CH C CH_2 CH_2$ (244°) Formed by the action of MeI on (*B* 4, *Py* 3) di-methyl-quinoline tetrahydride (Doebner & Miller, *B* 16, 2470) Colourless liquid

(*Py* 1, 2, 3) Tri-methyl-quinoline

$C_8H_8 \begin{smallmatrix} CMe \\ CMe \\ CMe \\ CMe \end{smallmatrix}$ [65°] (285°) Formed by heating the amide of di-methyl ethylidene diketone $CH_2 CO CHMe C(NC_2H_5)_2 CH_2$ with H_2SO_4 , and separated by passing NH_3 through the diluted product (Combes, *C R* 106, 144) — $B'_2H_2PtCl_6$ [215°]

Tri-methyl-quinoline $C_{12}H_{12}N$ (270°–280°) A by product in the preparation of quinaldine from paraldehyde, aniline, and HCl (Doebner & Miller, *B* 18, 3352) — $B'_2H_2PtCl_6$ 2aq pale-yellow needles

(*B* 1, 2, 4, *Py* 8)-Tetra-methyl-quinoline

$CMe CH C CH CH$ $CH CMe C N CMe$ [c 20°] (300°) Formed by heating solid ψ cumidine [63°] with paraldehyde and HClAq at 105° (Doebner & Miller, *B* 17, 1710) Crystalline, v sol alcohol and ether, insol water — $B'_2H_2Cr_2O_7$, long yellow needles

(*B* 2, 4, *Py* 1, 3)-Tetra-methyl-quinoline

$CMe CH C Me CH$ $CH CMe C N CMe$ [84°] (285° uncor).

Formed from acetone and *m*-xylidine (Levin & Riehm, *B* 19, 1894) White plates (from ether) — $B'H_2SO_4$, [235°–242°]; needles, v sol water, m. sol alcohol — $B'HCi$: white needles (by sublimation). — $B'_2H_2PtCl_6$ — $B'_2H_2Cr_2O_7$, orange needles, sl sol water

Tetramethylquinoline $C_{12}H_{12}N$ (265°–273°). Occurs in crude quinaldine, obtained from par

aldehyde, aniline, and HClAq (Einhorn, *B* 18, 3145) On oxidation with CrO_3/Cl_2 it yields an aldehyde $\text{C}_8\text{H}_7\text{NO}$, crystallising from water in needles (containing 8aq), melting at 74° when hydrated and 102° when anhydrous, it is oxidised by ammoniacal AgNO_3 to an acid crystallising in needles [224°]

Salt $-\text{B}'\text{H}_2\text{PtCl}_4$

References—AMIDO-, CHLORO-, NITRO-, and OXY-, METHYL-QUINOLINES, and METHYL-QUINOLYL

(γ) **METHYL-ISOQUINOLINE** $\text{C}_{10}\text{H}_9\text{N}$ e e
 $\text{C}_8\text{H}_7\text{N} \begin{smallmatrix} \text{CMe} \text{CH} \\ \text{CH N} \end{smallmatrix}$ (256° uncor) Formed by the distillation of either of the compounds $\text{CMe}_2\text{CO} > \text{NH}$ and $\text{CMe}_2\text{CO} > \text{NMe}$ with zinc-dust (Le Blanc, *B* 21, 2300) Liquid, not solidified at -75° $-\text{B}'\text{H}_2\text{PtCl}_4$ [253 5°]. Brownish red crystals $-\text{B}'\text{C}_6\text{H}_5(\text{NO}_2)_2\text{OH}$ [195°] Slender needles

(α)-Methyl-isoquinoline $\text{C}_8\text{H}_7\text{N} \begin{smallmatrix} \text{CH CH} \\ \text{CMe N} \end{smallmatrix}$

One of the products got by distilling papaveroline over zinc dust (Krauss, *M* 11, 361 Oil Volatile with steam $-\text{B}'\text{H}_2\text{PtCl}_4$ 13aq [229°] Orange yellow pp $-\text{Picrate}$ [210°] Light yellow pp

Derivative $-\nu$ CHLORO-METHYL-ISOQUINOLINE
METHYL-QUINOLINE-ACRYLIC ACID ν
METHYL-QUINOLYL-ACRYLIC ACID

(Py 3)-**METHYL-QUINOLINE** (B 2) **CARB- OXYLIC ACID** $\text{C}_{11}\text{H}_9\text{NO}_2$ e e
 $\text{CO}_2\text{H} \text{C} \begin{smallmatrix} \text{CH CH CH CH} \\ \text{CH CH C N CMe} \end{smallmatrix}$ *Quinaldine p-carb- orylic acid* [259°] Formed by the reaction of α mixture of paraldehyde, p amido benzoic acid, and HClAq (Doebner & Miller, *B* 17, 939) Formed also by oxidation of the corresponding di-methyl quinoline (Schunck, *B* 23, 2263) White needles, which may be sublimed Sol hot alcohol, ν sl sol water $-\text{HA}'\text{HCl}$ aq long slender needles or small prisms $-(\text{HA}')_2\text{H}_2\text{PtCl}_4$ 4aq monoclinic tables $-(\text{HA}')_2\text{H}_2\text{CrO}_4$ red needles, sol hot water $-\text{CaA}'_2$ 2aq feathery crystals $-\text{CuA}'_2$ 6aq small plates $-\text{AgA}'$ sparingly soluble crystalline powder $-\text{PbA}'_2$ prisms

(Py 3)-Methyl-quinoline (B 3) **carboxylic acid** $\text{CH CH C CH CH} \begin{smallmatrix} \text{CH CH C N CMe} \end{smallmatrix}$ *Quinaldine m-carboxylic acid* [285°] Formed by heating α mixture of paraldehyde, m amido benzoic acid, and HClAq (Doebner & Miller, *B* 17, 941) Formed also by the oxidation of the corresponding aldehyde by Ag_2O (Eckhardt, *B* 22, 281) Formed also by oxidation of the corresponding di methyl quinoline (Rust, *B* 23, 2262), and from the corresponding amido methyl quinoline by Sandmeyer's reaction (Rust, *B* 23, 3485) Long silky needles which may be sublimed Sol alcohol, nearly insol water $-\text{HA}'\text{HCl}$ aq small tables, sl sol cold water $-(\text{HA}')_2\text{H}_2\text{PtCl}_4$ monoclinic prisms $-(\text{HA}')_2\text{H}_2\text{CrO}_4$ yellow needles, ν sol hot, sl sol cold, water $-\text{CaA}'_2$ 2aq sparingly soluble prisms $-\text{CuA}'_2$ 8aq green tables $-\text{AgA}'$ crystalline pp

(Py 3)-Methyl-quinoline (B 4)-**carboxylic acid** $\text{CH CH} \begin{smallmatrix} \text{CH CH} \\ \text{CH C(CO}_2\text{H) C N CMe} \end{smallmatrix}$ *Quinaldine o-carboxylic acid* [151°] Formed by heating α mixture of amido benzoic acid (25 g), paraldehyde (18 g), and HClAq at 100° (Doebner

& Miller, *B* 17, 943) Formed also by oxidation of the corresponding di-methyl quinoline (R. Meyer, *B* 23, 2259) Colourless needles (containing 3aq), ν sol hot water and alcohol. Split up by heat into CO_2 and quinaldine. $-\text{HA}'\text{HCl}$ concentric tables, ν sol water $-(\text{HA}')_2\text{H}_2\text{PtCl}_4$ 2aq large red prisms, sol hot, sl sol cold, water $-\text{CuA}'_2$ 13aq small dark-green needles $-\text{AgA}'$ amorphous pp, changing into slender needles when heated with water for a long time

(B 4)-Methyl-quinoline (B 1)-**carboxylic acid** $\text{CH C(CO}_2\text{H)} \begin{smallmatrix} \text{CH C(CO}_2\text{H)} \\ \text{CH C(CH}_3\text{)} \end{smallmatrix} > \text{C}_8\text{H}_7\text{N}$ *o-Methyl-quinoline ana-carboxylic acid* [286°] Prepared by heating the corresponding di methyl-quinoline with dilute HNO_3 at 170° (Lellmann & Alt, *A* 237, 310) White powder Yields (B 4) methyl-quinoline on distillation with lime A solution of its ammonium salt gives a dirty green pp with FeSO_4 , a light green pp with CuSO_4 , and a white pp with lead acetate

Salts $-\text{HA}'\text{HCl}$ aq silky needles $-(\text{HA}')_2\text{H}_2\text{PtCl}_4$ 6aq yellow needles $-(\text{HA}')_2\text{H}_2\text{PtCl}_4$ 2aq needles $-(\text{HA}')_2\text{H}_2\text{PtCl}_4$ $-\text{CaA}'_2$ crystalline pp

(Py 3)-Methyl-quinoline (Py 1) **carboxylic acid** $\text{C}_8\text{H}_7\text{N} \begin{smallmatrix} \text{C(CO}_2\text{H) CH} \\ \text{N} \end{smallmatrix} \begin{smallmatrix} \text{CH} \\ \text{CMe} \end{smallmatrix}$ *Amlurtronic acid*. [242°]

Formation -1 By boiling α mixture of aniline and pyruvic acid with water, or by boiling anilpyruvic acid with water In this reaction $\text{CH}_3\text{CHCHCOCO}_2\text{H}$ may perhaps be assumed as an intermediate acid (Böttinger, *B* 14, 90, 133, *A* 191, 321) -2 By oxidising (Py 1, 3) di methyl quinoline with CrO_3 (Beyer, *J pr* [2] 33, 411, Seltz, *B* 23, 2257) -3 By heating isatin with acetone and dilute ($5 p c$) NaOHAq (Pfitzinger, *J pr* [2] 38, 582)

Properties $-\text{Lamins}$ or needles (containing aq), sol water and alcohol, ν e sol dilute acids Tastes bitter

Reactions -1 Yields quinaldine on distillation with lime (Kusel, *B* 19, 2249) -2 Oxidised by KMnO_4 to pyridine tricarboxylic acid -3 Bromine in chloroform forms an oily addition-product, which gives up all its bromine on treatment with boiling water (Böttinger, *B* 16, 2357)

Salts $-\text{HA}'\text{HCl}$ aq needles, loses its HCl when treated with cold water $-(\text{HA}')_2\text{H}_2\text{PtCl}_4$ 2aq $-\text{HA}'\text{HBr}$ 2aq $-\text{BaA}'_2$ aq $-\text{AgA}'$ small plates.

(Py 2) Methyl quinoline (Py 3)-**carboxylic acid** $\text{C}_8\text{H}_7\text{N} \begin{smallmatrix} \text{CH CMe} \\ \text{N C CO}_2\text{H} \end{smallmatrix}$ [144°] Formed by oxidation of (Py 2, 3)-methyl ethyl-quinoline (derived from aniline and propionic aldehyde) with CrO_3 and H_2SO_4 (Doebner & Miller, *B* 17, 1715, 18, 1641) Long silky needles or monoclinic prisms (from ether alcohol), sol hot water and alcohol Split up at 160° into CO_2 and (Py 2) methyl-quinoline $-\text{CuA}'_2$ sl sol water.

(Py 3) Methyl quinoline (Py 2) **carboxylic acid** $\text{C}_8\text{H}_7\text{N} \begin{smallmatrix} \text{CH C CO}_2\text{H} \\ \text{N CMe} \end{smallmatrix}$ *Quinaldine carboxylic acid* [234°] Formed by saponification of its ether, which is produced by the action of o -amido-benzoic aldehyde on an alkaline aqueous solution of acetoacetic ether (Friedländer & Göhring, *B* 16, 1836, *B* 19, 37) Formed also by oxidising (Py 2, 3)-di methyl quinoline with

chromic acid (Rohde, *B* 22, 287) Colourless needles, sl sol water, m sol alcohol. Split up on fusion into CO_2 and quinaldine.

Ethyl ether EtA'. [71°] Long white needles, insol water. May be distilled.—(EtA'), H_2PtCl_6 2aq sparingly soluble yellow needles.

Methyl iodide of the ethyl ether EtA'MeI [205°] Orange needles, sl sol cold water and alcohol, v sol hot water. Ag_2O converts it into the oxide ($\text{EtA}'\text{Me}$), O, an amorphous pp which, when freshly prepared, is sol benzene and chloroform, insol water, alcohol, ether, and ligroin, after a time it becomes insol benzene and chloroform. The oxide decomposes when heated between 180° and 240° HCl converts it into $\text{EtA}'\text{MeCl}$, which yields ($\text{EtA}'\text{MeCl}$), PtCl_4 , crystallising in golden plates [217°]

(*Py* 1) Methyl-quinoline (*B* 2) carboxylic acid $\text{CO}_2\text{H} \cdot \text{C} \cdot \text{CH} \cdot \text{C} \cdot \text{Me} \cdot \text{CH} \cdot \text{HC} \cdot \text{CH} \cdot \text{C} \cdot \text{N} \cdot \text{CH}$ *Lepidine p carboxylic acid* [250°–270°] Obtained by oxidation of the corresponding di methyl quinoline by chromic acid mixture (K Daniel, *B* 23, 2264) Stellate groups of needles Yields lepidine on distillation with soda lime

Methyl-quinoline carboxylic acid $\text{C}_{11}\text{H}_9\text{NO}_2$ $\text{C}_6\text{H}_5(\text{C}_5\text{HMeN} \cdot \text{CO}_2\text{H})$ [190°] Formed by oxidising the fraction 275°–285° (? di methyl quinoline) of the product of the action of aniline on a mixture of isobutyric aldehyde, methylal, and HCl (Von Miller a Kinkelin, *B* 20, 1940) Small crystals, m sol water and alcohol, gives off CO_2 on fusion

(*Py* 4) Methyl quinoline tetrahydride (*B* 3)-carboxylic acid

$\text{CH} \cdot \text{CH} \cdot \text{C} \cdot \text{CH}_2 \cdot \text{CH}_2 \cdot \text{CO}_2\text{H} \cdot \text{C} \cdot \text{CH} \cdot \text{C} \cdot \text{NMe} \cdot \text{CH}_2$ *Karoline m-carboxylic acid* [164°] Obtained by heating quinoline tetrahydride (*B* 3) carboxylic acid with MeI at 150° (Fischer a Korner, *B* 17, 765) Needles, sl sol water, v sol alcohol.

(*Py* 4) Methyl quinoline tetrahydride

(*Py* 1) carboxylic acid $\text{C}_6\text{H}_4 \cdot \text{CH}(\text{CO}_2\text{H}) \cdot \text{CH}_2 \cdot \text{N}(\text{CH}_3) \cdot \text{CH}_2$

Methyl tetrahydro cinchoninic acid Formed by heating the hydrochloride of quinoline tetrahydride (*Py* 1) carboxylic acid with MeOH and MeI at 100°, and decomposing the resulting salt with Ag_2O (Weidel, *M* 3, 66) Crystallises from cold alcohol in large prisms (containing 2aq), v sol water and alcohol, nearly insol ether. Reduces warm ammoniacal AgNO_3 . Decomposes on distillation, yielding an anhydride $\text{C}_{22}\text{H}_{19}\text{N}_2\text{O}$, which is a colourless oil (298° at 744 mm), insol ether, alcohol, and dilute acids, and is converted by heating with HClAq at 150° into MeCl and quinoline tetrahydride (*Py* 1) carboxylic acid (Weidel a Hazura, *M* 5, 643) On heating the anhydride with aqueous KOH it is converted into 'homohydrocinchoninic acid' $\text{C}_{11}\text{H}_{11}\text{NO}_2$, which crystallises in nearly leaflets [125°], insol water, sol alcohol and ether, and forms a hydrochloride $\text{C}_{11}\text{H}_{11}\text{NO}_2\text{HCl}$, crystallising in trimetric prisms ($a b c = 93 \cdot 1 \cdot 2 \cdot 07$), and a methyl iodide $\text{C}_{11}\text{H}_{11}\text{NO}_2\text{MeI}$ aq, forming monoclinic crystals, converted by Ag_2O into $\text{C}_{11}\text{H}_{11}\text{MeNO}_2$ aq, crystallising in glittering prisms, v sol water

Salts—The metallic salts are extremely deliquescent.— $\text{HA}'\text{HCl}$ aq large monoclinic

crystals—(HA'), H_2PtCl_6 large orange crystals.

— $\text{HA}'\text{Hlaq}$ large monoclinic crystals

(*B* 2, *Py* 3) Di-methyl-quinoline (*Py* 1)-carboxylic acid $\text{CMe} \cdot \text{OH} \cdot \text{C} \cdot \text{C}(\text{CO}_2\text{H}) \cdot \text{CH} \cdot \text{OH} \cdot \text{CH} \cdot \text{C} \cdot \text{N} \cdot \text{CMe}$

[262°] Formed by the action of aqueous NaOH on a mixture of *p* methyl-satin and acetone (Pfitzinger, *J pr* [2] 38, 584) Shining plates (from water)

(*B* 4, *Py* 3)-Dimethyl-quinoline (*B* 2)-carboxylic acid

$\text{CO}_2\text{H} \cdot \text{C} \cdot \text{CH} \cdot \text{C} \cdot \text{CH} \cdot \text{CH} \cdot \text{CH} \cdot \text{CMe} \cdot \text{C} \cdot \text{N} \cdot \text{CMe}$

Formed by oxidising (*B* 2, 4, *Py* 3) trimethyl-quinoline with CrO_3 and dilute H_2SO_4 (Panajotoff, *B* 20, 38) Needles (by sublimation), sl sol water and cold alcohol. On distillation with lime it gives (*B* 4, *Py* 3)-di-methyl-quinoline

Salts—(HA'), H_2PtCl_6 4aq orange needles.— BaA'_2 white needles, v sol water— AgA'_2 amorphous precipitate becoming crystalline— $\text{C}_6\text{H}_5(\text{NO}_2)\text{OH}$ [221°] Yellow needles, sl sol water, v sol alcohol

(*Py* 2, 3) Di methyl-quinoline (*B* 2)-carboxylic acid $\text{CO}_2\text{H} \cdot \text{C} \cdot \text{CH} \cdot \text{C} \cdot \text{CH} \cdot \text{CMe} \cdot \text{HC} \cdot \text{CH} \cdot \text{C} \cdot \text{N} \cdot \text{CMe}$ [270°]

Formed by oxidising (*B* 2, *Py* 2, 3 tri methyl quinoline (Miller a Ohler, *B* 23, 2268) Silky needles, sl sol water and alcohol. Gives on distillation (*Py* 2, 3) di methyl quinoline [63°]

— CuA'_2 aq (*Py* 3)-Methyl-quinoline (*B* 2, *Py* 1) di carboxylic acid $\text{CO}_2\text{H} \cdot \text{C} \cdot \text{CH} \cdot \text{C} \cdot \text{C}(\text{CO}_2\text{H}) \cdot \text{CH} \cdot \text{CH} \cdot \text{CH} \cdot \text{C} \cdot \text{N} \cdot \text{CMe}$

Formed by heating *p* amido benzoic acid (30 g) with pyruvic acid (20 g) and aldehyde (12 g) in alcoholic solution on a water bath (Von Miller a R Meyer, *B* 23, 2262) White powder, insol alcohol and other solvents. Cakes together at 160° Gives quinaldine on distillation with lime— CuA'_2 green crystalline pp

(*Py* 3)-METHYL-QUINOLINE (*B* 3)-CARBOXYLIC ALDEHYDE $\text{C}_{11}\text{H}_9\text{NO}$

$\text{CH} \cdot \text{CH} \cdot \text{C} \cdot \text{CH} \cdot \text{CH} \cdot \text{COH} \cdot \text{C} \cdot \text{CH} \cdot \text{C} \cdot \text{NMe}$ [61°] Prepared by adding KMnO_4 to a dilute solution of sodium methyl quinolyl acryic acid covered with benzene and frequently shaking (Eckhardt, *B* 22, 277) Slender hair like crystals (containing $1\frac{1}{2}$ aq!) Melts at 61° when anhydrous and 73° when hydrated. V sol dilute HClAq , sl sol acetic acid, v sol alcohol, ether, and benzene, m sol ligroin and hot water. Volatile with steam. Reduces ammoniacal AgNO_3 . Gives a crystalline sulphate, and a red colour with phenyl hydrazine acetate. Ag_2O oxidises it to the corresponding carboxylic acid [285°] On heating with methyl-quinoline and ZnCl_2 for 2 hours at 150° it yields a condensation product [69°]

Salts— $\text{B}'\text{HCl}$ pale yellow needles— $\text{B}'\text{H}_2\text{PtCl}_6$ [211°] Triclinic crystals (from alcohol containing HCl)— $\text{B}'\text{C}_6\text{H}_5(\text{NO}_2)\text{OH}$ [182°] Needles, blackening at 174°

Phenyl-hydrazide $\text{C}_{11}\text{H}_9\text{N}_2$ The salt $\text{C}_{11}\text{H}_9\text{N}_2\text{HCl}$ separated as brick-red hair-like crystals on adding a hot solution of phenyl hydrazine hydrochloride to a hot solution of the base in HClAq . In presence of H_2SO_4 the salt ($\text{C}_{11}\text{H}_9\text{N}_2$), (H_2SO_4) 9aq separates in small brick red needles

(Py. 8) Methyl-quinoline (B 2)-carboxylic aldehyde $\text{CHO} \cdot \text{C} \cdot \text{CH} \cdot \text{C} \cdot \text{CH} \cdot \text{CH}$
 $\text{OH} \cdot \text{CH} \cdot \text{C} \cdot \text{N} \cdot \text{CMe}^*$ [106°]

Formed by oxidising the corresponding methyl-quinolyl acrylic acid with KMnO_4 , accompanied by agitation with benzene (Miller & Kinkelin, B. 18, 3237). Needles (from water), m sol ligroin and hot water, v sol alcohol, ether, benzene, and acids. On heating with quinaldine at 150° it forms $\text{C}_{10}\text{H}_7\text{N} \cdot \text{CH} \cdot \text{CH} \cdot \text{C}_6\text{H}_5\text{NMe}$, a yellow insoluble substance melting above 300°.— $\text{B}_2\text{H}_6\text{PtCl}_4 \cdot 2\text{aq}$ orange prisms.

Phenyl hydrazide $\text{C}_{17}\text{H}_{11}\text{N}_3$. [160°]. Golden prisms (from alcohol).

(B 2, 4) Di methyl quinoline (Py 8) carb-oxylic aldehyde $\text{CH} \cdot \text{CMe} \cdot \text{C} \cdot \text{CH} \cdot \text{CH}$
 $\text{C} \cdot \text{CH} \cdot \text{C} \cdot \text{N} \cdot \text{CMe}$ [107°]

Formed by oxidising (B 2, 8) di methyl quinolyl-acrylic acid with KMnO_4 (Panajotoff, B. 28, 1471). Yellowish plates, v sol alcohol and ether, sl sol water. Sol acids, insol alkalis.

Tri-methyl-quinoline carboxylic aldehyde $\text{C}_{13}\text{H}_{11}\text{NO}$ [101.5°]. Formed from tetra methyl-quinoline (derived from paraldehyde, aniline, and HCl) by treatment with CrO_3Cl , followed by water (Einhorn, B. 18, 3144). Crystallises from water in needles (containing 3aq) which melt at 74°. When anhydrous it melts at 101.5°. It reduces ammoniacal AgNO_3 , forming a mirror. It yields an oxim [203°] and a phenyl-hydrazide [207°].

METHYL-QUINOLINE MERCAPTAN *o*. METHYL-QUINOLYL MERCAPTAN

(B 2) METHYL-QUINOLINE (B 4)-SULPHONIC ACID $\text{C}_{10}\text{H}_7\text{NSO}_3$, *o*

$\text{C}(\text{CH}_3) \cdot \text{CH} \cdot \text{CH} \cdot \text{C}(\text{SO}_3\text{H}) > \text{C}_6\text{H}_4\text{N}$ *p*-Toluquinoline sulphonic acid. Formed by boiling a mixture of *p*-toluidine sulphonic acid, nitrobenzene, glycerin, and H_2SO_4 (Fischer & Wittmack, B. 17, 441). Formed also by the sulphonation of (B 2) methyl-quinoline (Herzfeld, B. 17, 1552). Colourless plates, sl sol boiling water— KA' v sol water— BaA'_2 (dried at 130°) amorphous, sl. sol water.

(B 4) Methyl-quinoline (B 1)-sulphonic acid $\text{CH} \cdot \text{C}(\text{SO}_3\text{H}) > \text{C}_6\text{H}_4\text{N}$. Obtained by heating *o*-toluidine sulphonic acid $\text{C}_6\text{H}_4\text{Me}(\text{NH}_2)\text{SO}_3\text{H}$ [124] with glycerin, nitrobenzene, and H_2SO_4 (Herzfeld, B. 17, 904). Formed also by sulphonating (B 4) methyl quinoline (Herzfeld, B. 17, 1550). Flat prisms (from water), v sol water— KA' long soluble tables— BaA'_2 soluble trimetric tables.

(B 4) Methyl-quinoline (B 2)-sulphonic acid $\text{SO}_3\text{H} \cdot \text{C} \cdot \text{CH} > \text{C}_6\text{H}_4\text{N}$. Formed by heating *o*-toluidine sulphonic acid $\text{C}_6\text{H}_4\text{Me}(\text{NH}_2)\text{SO}_3\text{H}$ [125] with glycerin, nitrobenzene, and H_2SO_4 (Herzfeld, B. 17, 903). Colourless sparingly soluble prisms.

(Py 8) Methyl-quinoline (B. 2)-sulphonic acid $\text{SO}_3\text{H} \cdot \text{C} \cdot \text{CH} \cdot \text{C} \cdot \text{CH} \cdot \text{CH}$
 $\text{CH} \cdot \text{CH} \cdot \text{C} \cdot \text{N} \cdot \text{CMe}^*$ Quinaldine *p*-sulphonic acid. Formed in small quantity in the sulphonation of quinaldine. Prepared by heating amido-benzene-*p*-sulphonic acid (100 pts) with paraldehyde (80 pts) and HCl (100 pts) for 2 hours on the water-bath (Doebner & Miller, B. 17, 1708). Small monoclinic crystals, v sol. hot

water. On fusion with KOH it yields oxy-methyl-quinoline [213°].

(Py 8) Methyl-quinoline (B. 4)-sulphonic acid $\text{CH} \cdot \text{CH} > \text{C} \cdot \text{CH} \cdot \text{CH}$
 $\text{CH} \cdot \text{C}(\text{SO}_3\text{H}) \cdot \text{C} \cdot \text{N} \cdot \text{CMe}^*$ Quinaldine *o*-sulphonic acid. Formed, in small quantity, in sulphonating quinaldine (Doebner & Miller, B. 17, 1708). Long flat triclinic prisms, v sol. hot water. On fusion with potash it yields oxy-methyl quinoline [74°].

(Py. 8) Methyl-quinoline (B 1 or 8)-sulphonic acid $\text{CH} \cdot \text{C}(\text{SO}_3\text{H}) \cdot \text{C} \cdot \text{CH} \cdot \text{CH}$
 $\text{CH} \cdot \text{CH} > \text{C} \cdot \text{N} \cdot \text{CMe}$ or

$\text{CH} \cdot \text{CH} > \text{C} \cdot \text{CH} \cdot \text{CH}$
 $\text{C}(\text{SO}_3\text{H}) \cdot \text{CH} \cdot \text{C} \cdot \text{N} \cdot \text{CMe}$ Quinaldine *m*-sulphonic acid. The chief product of the sulphonation of quinaldine (Doebner & Miller, B. 17, 1708). Glistening monoclinic prisms, sol hot, sl sol cold, water. On fusion with potash it gives oxy methyl quinoline [234°]. Distillation with KCy yields the nitrile of the corresponding carboxylic acid (Richard, B. 23, 3488).

(Py 1) Methyl-quinoline sulphonic acid $\text{C}_6\text{H}_4(\text{SO}_3\text{H}) < \text{CMe} \cdot \text{CH}$
 $\text{N} \cdot \text{CH}$ Lepidine sulphonic acid.

Formed by heating 'homohydrocinchononic acid' $\text{C}_{16}\text{H}_{19}\text{NSO}_3$ with H_2SO_4 at 180° (Weidela Hazura, M. 5, 652). Crystallises from water in thin plates (containing aq), nearly insol cold, v. sol. hot water.

(Py 1) Methyl quinoline (B 2) sulphonic acid $\text{SO}_3\text{H} \cdot \text{C} \cdot \text{CH} \cdot \text{C} \cdot \text{Me} \cdot \text{CH}$
 $\text{HC} \cdot \text{CH} \cdot \text{C} \cdot \text{N} = \text{CH}$ Formed by heating lepidine (1 pt) with H_2SO_4 (8 pts) at 300° (Busch & Koemigs, B. 23, 2680). Silky needles (containing 2aq), v sol hot water— AgA' aq. white jelly, becoming crystalline.

(Py 1, 3) Di-methyl-quinoline sulphonic acid $\text{C}_{11}\text{H}_{11}\text{NSO}_3$, *o* $\text{C}_6\text{H}_4(\text{SO}_3\text{H}) < \text{CMe} \cdot \text{CH}$
 $\text{N} \cdot \text{CMe}$ Formed by sulphonating the corresponding di methyl-quinoline (Beyer, J. pr. [2] 33, 407). Tables or flat needles, not melting below 303°. On fusion with potash it yields oxy di methyl quinoline [44°].

(B 1, 4) Di-methyl-quinoline (B 2) sulphonic acid $\text{SO}_3\text{H} \cdot \text{C} \cdot \text{CMe} > \text{C}_6\text{H}_4\text{N}$. *p*-Xyloquinoline sulphonic acid. Formed by heating (B 1, 4)-di-methyl-quinoline with fuming H_2SO_4 . Formed also from xyldine sulphonic acid by heating with nitrobenzene, glycerin, and H_2SO_4 (Nöbling & Frühling, B. 21, 3157)— KA' needles or plates, v. sol water— BaA'_2 aq plates, v. sol. hot water— $\text{BaA}'_2 \cdot 2\text{aq}$.

(B 1, 4)-Di-methyl-quinoline (B. 8)-sulphonic acid $\text{SO}_3\text{H} \cdot \text{C} \cdot \text{CMe} > \text{C}_6\text{H}_4\text{N}$. Formed by heating xyldine sulphonic acid (derived from *p*-xylene sulphonic acid) with nitrobenzene, glycerin, and H_2SO_4 (Nöbling & Frühling, B. 21, 3156). Short white plates, sl sol cold water, v. sol hot water and dilute acetic acid— KA' aq v. e sol. water— BaA'_2 aq. needles, v. sol. hot water.

(B 2, 4)-Di-methyl-quinoline sulphonic acid $\text{C}_6\text{HMe}(\text{SO}_3\text{H})(\text{C}_6\text{H}_4\text{N})$. Formed by heating the corresponding di-methyl-quinoline with fuming H_2SO_4 at 165° (Berend, B. 17, 2716). Minute needles (from alcohol-ligron).

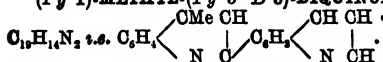
(B 2, 4, Py 3)-Tri-methyl-quinoline (B 1)-sulphonic acid. Formed from the corresponding tri-methyl-quinoline by heating with H_2SO_4 at 125° (Panajotoff, B 20, 36) Small yellow needles (from dilute HCl aq), not melting at 260°. Insol cold water — BaA, 384g silky needles

METHYL-QUINOLINIC ACID v METHYL-PYRIDINE DICARBOXYLIC ACID

METHYL-QUINOLINIUM HYDRATE v.

Methylo-hydrate of QUINOLINE

(Py 1)-**METHYL-(Py 3 B 3)-DIQUINOLYL**



[188°]. Obtained by heating flavamine (10 pts), with nitrobenzene (5 pts), glycerin (30 pts), and H_2SO_4 (80 pts) (Fischer, B 19, 1036) Colourless crystals. Strong base. Its salts with mineral acids have a splendid blue fluorescence in dilute solution

Methylo-sodide B'Mel fine yellowish needles, easily soluble in water

Di-(Py 3) methyl-di-quinolyl $C_{22}H_{26}N_4 \cdot n$

$\begin{array}{c} H \text{ OH} \\ \diagdown \quad \diagup \\ \text{Me. N} > C_6H_5 \end{array} \begin{array}{c} \text{CH CH} \\ \diagdown \quad \diagup \\ N \quad \text{OMe} \end{array}$ [207°] (above 360°) Prepared by gradually adding paraldehyde (90 g) to a solution of benzidine (80 g) in conc HCl aq (400 g) at 100°. After heating for ten hours the product is diluted with water, treated with $NaNO_2$, boiled, and pptd by KOH. Hins, A 242, 326) Slender white needles, sol alcohol, benzene, and chloroform, sl sol water and ether. — $B''H_2PtCl_2$ 2aq light yellow pp, sl sol. hot water — $B''H_2HNO_3$ small colourless needles, v sol water, sl sol alcohol — $B''H_2Cr_2O_7$, slender yellow needles, sl sol hot water

Di-methyl-di-quinolyl $C_{22}H_{26}N_4$ [162°] Formed by heating quinaldine with sulphur (Von Miller, B 21, 1928) Crystallises from alcohol in white needles (containing aq) Its picrate crystallises in needles. The platinum-chloride forms needles, v sl sol water

Tetra-methyl-di-quinolyl $C_{22}H_{26}N_4 \cdot n$
 $\begin{array}{c} CH \text{ OMe} \\ \diagdown \quad \diagup \\ \text{OMe N} > C_6H_5 \end{array} \begin{array}{c} \text{CH CH} \\ \diagdown \quad \diagup \\ N \quad \text{OMe} \end{array}$

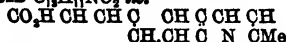
Tetra-methyl diquinolyl [232°] Formed by heating benzidine with acetone and conc HCl aq at 180° (Schestopal, B 20, 2506) White plates, insol water, sl sol ether, v sol alcohol. When heated with benzoic aldehyde and $ZnCl_2$ at 180° it forms a compound crystallising from alcohol in yellow needles

Salts — $B''H_2Cl$. Needles, v sol water and alcohol — $B''H_2SO_4$. Needles (from water) or prisms (from alcohol) — $B''H_2PtCl_2$ needles — $B''H_2Cr_2O_7$, needles, sl sol cold water — Picrate insol water and cold alcohol. — $B''H_2Cl$, 2HCl flesh coloured needles

Methylo-sodide B'Mel, [270°].

Ethylo-sodide B'EtL, [158°].

(Py 3)-**METHYL QUINOLYL-(B 2)-ACRYLIC ACID** $C_{11}H_{11}NO_2 \cdot n$.



Quinaldine-acrylic acid Obtained by heating a mixture of paraldehyde (40 g), *p*-amido-cinnamic acid hydrochloride (50 g.), and conc. HCl (50 g) (Miller & Kunkeln, B 18, 8285) Small concentric needles. Decomposes between 240° and 250°. Sol hot alcohol, sl sol. cold alcohol and water. By an alkaline solution of $KMnO_4$ it is

oxidised to methyl-quinoline carboxylic aldehyde C_9H_9MeCHO

Salts — $A'H_2HCl$ aq easily soluble concentric prisms — $A'H_2HNO_3$ aq needles or prisms, sl sol dilute HNO_3 — $(A'H_2HCl) \cdot PtCl_2$ 2aq, reddish-yellow prisms

(Py 3)-Methyl-quinolyl (B 3)-acrylic acid

$C_{11}H_{11}NO_2 \cdot n$. $CO_2H \text{ CH CH } C \text{ OH } C \text{ N } CMe$ [246°] Obtained by heating *m*-amido-cinnamic acid with paraldehyde and conc HCl aq at 150° (Eckhardt, B 22, 272) Small white monoclinic prisms (from alcohol), tending to become yellow, sl sol ether, chloroform, and petroleum ether, m sol alcohol, benzene, and acetone. When heated above its melting point a small sublimate of slender needles [223°] is obtained. $KMnO_4$ oxidises it to the corresponding methyl-quinoline carboxylic aldehyde. On warming with chloral on the water bath two compounds are formed. One of these compounds $C_{11}H_{11}NO_2 \cdot CH_2CH(OH)CO_2H$ crystallises from alcohol in prisms [201°], which dissolve in nitric acid with violet fluorescence, and which yield the salts $C_{11}H_{11}Cl_2NO_2 \cdot HCl$ [over 300°] and $AgC_{11}H_{11}Cl_2NO_2$ crystallising in slender needles. The other compound $C_{22}H_{26}Cl_2N_4O_2$ crystallises from alcohol in needles [128°] which dissolve in nitric acid with blue fluorescence, and which forms a salt $C_{22}H_{26}Cl_2N_4O_2 \cdot HCl$ [217°]

Salts — $HA'HCl$ aq deliquescent trimetric needles — $(HA') \cdot H_2PtCl_2$ 2aq yellow needles or plates — $HA'HNO_3$ aq needles, sol water — $HA'C_6H_5(NO_2)OH$ aq [152°] Bundles of slender needles, sol alcohol, hot water, and HOAc, sl sol ether — CaA' 3aq slender needles (from water), v sol dilute HCl aq and acetic acid — AgA' 2aq minute crystalline lancelets — AgA' 4aq needles

(Py 3)-Methyl-quinolyl-(B 1?)-acrylic acid $C_{11}H_{11}NO_2$. Formed on one occasion in the preparation of the preceding isomeride (E) Crystallises in yellowish monoclinic plates (containing aq) [184°] and in white crystals (containing $\frac{1}{2}$ EtOH) [204°] Its ammoniacal solution, unlike that of its isomeride, is not pptd. by salts of Ba, Ca, and Mg

(B 2, 4)-Di-methyl-(Py 3)-quinolyl-acrylic acid $\begin{array}{c} CMe \text{ CH } C \text{ CH CH} \\ \diagdown \quad \diagup \\ CH \text{ CMe } C \text{ N } C \text{ CH CH } CO_2H \end{array}$ Formed by boiling the compound of chloral and (B 2, 4, Py 3) tri-methyl-quinoline with aqueous K_2CO_3 (Panajotoff, B 20, 42) Yellowish needles, decomposing at 180°, forming a product melting at 210°

METHYL-QUINOLYL-AMMONIUM IODIDE v. Methylo-sodide of QUINOLINE

(Py 1)-**METHYL-QUINOLYL ETHYL SULPHIDE** $C_9H_9 \begin{array}{c} \text{CMe CH} \\ \diagdown \quad \diagup \\ N \quad \text{CSEt} \end{array}$ Formed by treating methyl-quinolyl mercaptan with NaOEt and EtI (Roos, B 21, 628) Oil, volatile with steam — $B''H_2PtCl_2$ 4aq — $B''HI$ [214°] Long yellow needles

(Py 3)-Methyl-quinolyl ethyl sulphide $C_9H_9 \begin{array}{c} \text{C(SeEt) CH} \\ \diagdown \quad \diagup \\ N = \text{CMe} \end{array}$ [56°]. Formed from sodium methyl-quinolyl mercaptide and EtI (R). White needles, v. sol. alcohol and ether, insol. water

together with benzoyl methyl-indole by heating methyl-indole (methyl ketone) with benzoyl chloride and ZnCl_2 . Also formed by oxidation of the leuco-compound benzylidene-methyl-indole. Amorphous yellow pp, v sol alcohol and ether, or yellowish-red prismatic crystals, sl sol alcohol, v sl sol ether. The amorphous modification is converted into the crystalline by long boiling with water. It dissolves in acids and in alcoholic NaOH or KOH with a red colour. Its salts are red dyestuffs closely resembling rosaniline. By zinc-dust and NH_3 it is reduced to benzylidene-methyl indole [248°]—B'HCl small metallic green crystals, sl sol water (Fischer & Wagner, B 20, 815).

METHYL SALICYLIC ACID v **OXY-TOLUIC ACID** and the *Methyl derivative of o-OXY BENZOIC ACID*

METHYL-SALICYLIC ALDEHYDE v *Methyl derivative of o OXY BENZOIC ALDEHYDE*

DI-METHYL-SELENAZOLE v **SELENIUM, ORGANIC COMPOUNDS.**

METHYL-SELENIDE v **SELENIUM, ORGANIC COMPOUNDS**

METHYL SILICATE $\text{C}_2\text{H}_5\text{SiO}_3$, i.e. $\text{Si}(\text{OMe})_2$, (122°) SG 2 10589 VD 538 (calc 526) Formed by the action of SiCl_4 on dry methyl alcohol (Friedel & Crafts, A Ch [4] 9, 32) Colourless liquid with fragrant ethereal odour, m sol water, the solution depositing gelatinous silica when kept for a month. When heated with SiCl_4 in various proportions it gives rise to the compounds $\text{ClSi}(\text{OMe})_2$, (115°), SG 2 1195, VD 558 (calc 542), $\text{Cl}_2\text{Si}(\text{OMe})_2$, (c 100°), SG 2 1260, VD 566 (calc 557), and $\text{Cl}_3\text{Si}(\text{OMe})_2$, (c 84°), VD 566 (calc 573).

Hexa-methyl disilicate $\text{Me}_6\text{Si}_2\text{O}_7$, (202°) SG 2 1144 VD 919 (calc 893) Formed when, in the preparation of Me_4SiO_4 , the methyl alcohol is not quite dry. Formed also by heating Me_4SiO_4 (2 mols) with water (1 mol) and methyl alcohol.

METHYL-STIBINE v **ANTIMONY, Compounds with organic radicals**, vol 1 298

METHYL-STILBAZOL v **STYRYL METHYL-STYRIDINE.**

METHYL-STILBENE v. **s-PHENYL-TOLYL-ETHYLENE**

Di-methyl-stilbene v **Di-TOLYL-ETHYLENE**

Tetra-methyl-stilbene v **Di-XYLYL-ETHYLENE**

METHYL-STYCHONINE v **STYCHONINE**

METHYL-STYRENE v **TOLYL-ACETYLENE**

METHYL STYRYL KETONE v. **STYRYL**

METHYL KETONE

METHYL-SUCCINIC ACID v **PYROTARTARIC ACID**

u-Di-methyl-succinic acid $\text{C}_6\text{H}_8\text{O}_4$, i.e. $\text{CO}_2\text{H} \cdot \text{CH}_2 \cdot \text{CMe}_2 \cdot \text{CO}_2\text{H}$ *Isodipic acid* *Isobutane dicarboxylic acid* Mol w. 146 [140°] H C v 671,400 H Cp 671,700 H F 237,800 (Stohmann, Kleber, & Langbein, J pr [2] 40, 212)

Formation.—1 From its imide, which is formed, together with di-methyl malonic acid, by oxidising mesitylic acid $\text{C}_6\text{H}_3\text{NO}_2$ with KMnO_4 in acid solution (Pinner, B 15, 582)—2 By boiling with HClAq the isobutane tricarboxylic ether $\text{CO}_2\text{Et} \cdot \text{CMe}_2 \cdot \text{CH}(\text{CO}_2\text{Et})_2$ obtained by the action of α -bromo isobutyric ether on sodio malonic ether (Leuckart, B 18, 2350, Bischoff, B 23, 1943). Formed also by heating

the corresponding isobutane tricarboxylic acid (Barnstein, A 242, 133)—3 By oxidising with chromic acid mixture the terpene $\text{C}_{10}\text{H}_{16}$, obtained from copaiba balsam (Levy & Englander, A 242, 192)—4 By oxidising tropilene C_8H_8 with nitric acid (S G 138) (Ladenburg, A 217, 139)—5 By heating its nitrile (obtained from isobutylene bromide and KC_y) with HClAq at 150° (Hell & Rothberg, B 22, 1740)

Properties.—Thick colourless prisms (from benzene), crystallising from water in efflorescent monoclinic forms, $a b c = 2.029 : 1.191$; $\alpha = 118^\circ 36'$, $\beta = 95^\circ 16'$, $\gamma = 101^\circ$ V sol water, alcohol, ether, and acetone, v sl sol chloroform and ligroin. At about 185° it splits up into CO_2 and its anhydride.

Salts—**KHA** 2 $\frac{1}{2}$ aq small prisms (P)—**KHA** 5aq (B)—**NaHA** 3 $\frac{1}{2}$ aq Monoclinic prisms, $a b c = 1.8365 : 1.41801$, $\beta = 90^\circ 43'$ (L & E)—**NaA** 11aq needles, v sol water—**NH₄HA**—(NH $_4$) $_2$ A—**BaA** 2aq (B)—**BaA** 2 $\frac{1}{2}$ aq monoclinic plates, $a b c = 1.601 : 1.1790$, $\beta = 97^\circ 26'$ Sl sol hot, m sol cold, water, insol alcohol (L & E)—**CaA** 7aq minute plates, sl sol water, insol alcohol—**CdA** 6aq—**CuA** 2aq—**PbA** 7aq—**Ag₂A**: white pp, v sl sol water

Methyl ether Me_2A (200°) SG $\frac{1}{2}$ 10568

Ethyl ether EtA . (215°) SG $\frac{1}{2}$ 10134 (B), II 9976 (L & E)

Anhydride $\text{CMe}_2\text{CO} \cdot \text{CO} \cdot \text{O}$ [29°] (218°) (B), (220°) (L & E)

Chloride $\text{C}_2\text{H}_5\text{Me}_2(\text{C}_2\text{O}_2\text{Cl}_2)$ (201°) (B), (193°) (L & E) Formed by heating the acid with PCl_5 at 125° Reacts with phenyl hydrazide, forming the compound $\text{CMe}_2\text{CO} \cdot \text{CO} \cdot \text{N} \cdot \text{NHPh}$ [182°], which yields a nitrosamine [76°]

Imide $\text{CMe}_2\text{CO} \cdot \text{CO} \cdot \text{NH}$ [106°] Formed from the chloride and NH_3 Plates (from ether) Yields $\text{C}_2\text{H}_5\text{KNO}_2$ 2 $\frac{1}{2}$ aq, crystallising in prisms, v. sol alcohol

Nitrile $\text{CN} \cdot \text{CH}_2 \cdot \text{CMe}_2 \cdot \text{CN}$ *Isobutylenes cyanide* (219°) Formed by leaving a mixture of isobutylene bromide and alcoholic KC_y to stand for two weeks, heating to 140° to expel alcohol, and extracting the residue with ether (Hell & Rothberg, B 22, 1740) Clear liquid, v sol water

Anti-s-di-methyl succinic acid $\text{C}_6\text{H}_8\text{O}_4$, i.e. $\text{CO}_2\text{H} \cdot \text{CHMe} \cdot \text{CHMe} \cdot \text{CO}_2\text{H}$ *Maleonoid di-methyl succinic acid* *Butane dicarboxylic acid* [120°] (O & R, H & R, B & V), [124°] (Z) S 30 at 14°

Formation.—1 Together with the isomeric 'para' or fumaroid acid [194°] by reduction of di-methyl maleic acid (Otto & Rössing, B 20, 2738)—2 Together with the isomeric acid [194°] by heating with HClAq the mixture of their ethers obtained by adding α -bromo-propionic ether to an alcoholic solution of α -cyano propionic ether and NaOEt (Zelinsky, B 21, 8160)—3 A mixture of the ethers of the two isomeric s-di-methyl-succinic acids is also formed, with other bodies, when α -bromo-propionic ether is heated with finely-divided silver (Hell & Rothberg, B 22, 60)—4 The mixture of 'anti-' and 'para' di-methyl succinic acids is also formed by hydro-

lysis of butane tricarboxylic acid derived from sodium methyl malonic ether and a bromo propionic ether (Leuckart, *B* 18, 2346, Bischoff a Voit, *B* 22, 889) —5 From its anhydride, which is formed, together with the anhydride of the fumaroid isomeride, by heating the fumaroid acid [194°] for several hours at 200° (Bischoff a Voit, *B* 23, 641)

Properties — Concentrically grouped prismatic needles, more sol hot than cold water, v sol ether, alcohol, acetone, and chloroform, sl sol CS₂, and benzene, almost insol ligroin. A neutral solution of its ammonium salt gives a yellowish red pp with FeCl₃, a greenish blue pp with cupric sulphate, but no pp with salts of Ba, Zn, Mg, Co, and Ni. Bromine at 130° converts it into di methyl maleic acid [95°]

Salts — CaA'' 2aq crystalline powder, sl sol cold water — BaA'' 3aq thin plates, sl sol water — Ag.A'' white crystalline pp

Methyl ether Me.A'' (200°) From Ag.A'' and MeI at 100° Oil, with pleasant odour (Zehnsky a Krapivin, *B* 22, 646)

Ethyl ether Et.A'' (220°) SG § 1 0218, $\frac{1}{2}$ 1 0072 (Z a K), $\frac{1}{2}$ 1 0315 (B a V) H C 1,296,860 (Ossipoff, *C* R 109, 224) Oil, decomposed by heat into the anhydride and Et.O

Chloride (186°-197°)

Imide $\begin{matrix} \text{CHMe CO} \\ \text{CHMe CO} \end{matrix} > \text{NH}$ [106°] (B a V).

[110°] (Z a K) Obtained by distilling the dry ammonium salt in gaseous NH₃. Stellate groups of prisms (from dilute alcohol) or thin needles (from benzene), v sol water, alcohol, benzene and chloroform, sl sol ether, v sl sol ligroin reconverted into the acid [120°] by alkalis

Anilide CONHPh CHMe CO CONHPh [222°] Formed from the chloride and aniline. Needles (from alcohol)

Phenyl-imide $\begin{matrix} \text{CHMe CO} \\ \text{CHMe CO} \end{matrix} > \text{NPh}$ [146°]

Formed by heating the acid (1 mol) with aniline (2 mols) till the aniline begins to distil. Slender needles, v sol alcohol, ether, and benzene, sl sol water

Anhydride $\begin{matrix} \text{CHMe CO} \\ \text{CHMe CO} \end{matrix} > \text{O}$ [87°] When

formed by heating the 'anti' acid to 200°, it yields only the 'anti' acid again when heated with water, but when formed from the 'para' acid by like treatment the product (a mixture of anhydrides?) yields a mixture of 'para' and 'anti' acids, the amount of the latter increasing with the duration of the heating. Formed also from the 'anti' acid by heating with AcCl. In all cases it melts at 87°. When heated with bromine in chloroform at 90° it yields di methyl-maleic acid

'Para' s-di-methyl-succinic acid C₆H₁₀O₄, i.e. MeCH(CO₂H)CHMeCO₂H *Fumaroid variety* of s di methyl succinic acid *Isocadic acid* *Hydroxyprocionchonic acid* [194°] (O a B), (B a V) (192°) (Z) H F 238,000 H C v 670,700 H C p 671,000 (Stohmann, Kleber, a Langbein, *J* pr [21 40, 212] S 1 at 22°

Formation —1 By heating a bromo propionic acid with reduced silver at 155° (Wishnecus, *B* 2, 720, cf anti-di methyl succinic acid, *Formation* 3) —2 By boiling (aβ) di-methyl-acetyl succinic ether with conc alcoholic KOH (Hardtmuth, *A* 192, 143) —8 By

boiling an aqueous solution of sodium di-methyl-maleate with sodium amalgam (Weidel, *A* 173, 109, *M* 3, 612) —4 From its amide which is formed by the action of NH₃ on an oily product of the action of bromine on a solution of cyanoethine in dilute H₂SO₄ (E von Meyer, *J* pr [2] 26, 358) —5 Together with the isomeride [120°] by the reduction of di methyl maleic anhydride by HIAq at 220° or by sodium amalgam (Otto a Beckurts, *B* 18, 838, Otto a Rössing, *B* 20, 2736) —6 By heating cyano-di methyl succinic acid with HClAq (Zehnsky, *B* 21, 3166) —7 Together with the 'anti' isomeride, by all the modes of formation described above for that acid —8 By heating the 'anti' isomeride for several hours with conc HClAq at 180°-190° (Bischoff a Voit, *B* 23, 643)

Properties — Small triclinic needles (from alcohol), sl sol water, m sol alcohol. On heating at 200° it yields a mixture of its anhydride and that of the 'anti' acid, after several hours' heating it is almost wholly converted into the anhydride of the 'anti' acid. Its neutral solution is ppd by FeCl₃, CuSO₄, BaCl₂, and lead acetate. On treatment with bromine it yields di methyl-maleic acid

Salts — NH₄HA'' (dried at 100°). Monoclinic prisms, v sol water — CaA'' 2aq prisms (Bischoff a Rach, *A* 234, 76) — CaA'' aq (Z) — CaA'' 1½aq (W) Monoclinic needles — SrA'' — BaA'' 4aq — PbA'' — PbA'' ½aq short prisms — CuA'' — Ag₂A''

Methyl ether Me.A'' (199°) Oil (Zehnsky a Krapivin, *B* 22, 650) Yields the acid [192°] on saponification

Ethyl ether Et.A'' (220°) SG § 1 013, $\frac{1}{2}$ 1 002 H C 1,303,570 Oil, with pleasant odour. Yields on saponification a mixture of the acids [120°] and [192°]

Chloride (186°-197°)

Imide $\begin{matrix} \text{MeCH} - \text{CO} \\ \text{CHMe CO} \end{matrix} > \text{NH}$ [78°] From the

ether and NH₃. Crystallises from benzene. Yields only the 'para' acid on saponification

Anilide MeCH(CONHPh)CHMe CONHPh [235°] From the chloride and aniline. Needles, sol ether and HOAc. Yields only the 'para' acid on saponification

Phenyl imide $\begin{matrix} \text{MeCH} - \text{CO} \\ \text{CHMe CO} \end{matrix} > \text{NPh}$

[126°] Formed by heating the acid with aniline. Caustic potash converts it into a mixture of 'anti' and 'para' acids

Anhydride $\begin{matrix} \text{MeCH} - \text{CO} \\ \text{CHMe CO} \end{matrix} > \text{O}$ [88°]

Formed from the acid and AcCl. May be reconverted into the original acid. By heating the 'para' acid to 180°-196° a mixture of anhydrides [87°] is formed, which yields on saponification a mixture of 'anti' and 'para' acids

Derivative — v Di-chloro-di-methyl-succinic acid

Tri-methyl-succinic acid C₆H₁₂O₄, i.e. CO₂H CHMe CMe₂ CO₂H [105°] *Electrical conductivity* Bischoff, *B* 23, 1466 Formed by the hydrolysing action of H₂SO₄ on pentane tricarboxylic ether obtained from sodium methyl-malonic ether and a bromo isobutyric ether (Bischoff a Mintz, *B* 23, 649) Yields an anhydride melting between 67° and 89°

This acid is probably identical with s-di-

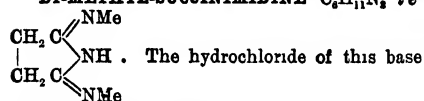
methyl-glutaric acid (Bischoff & Jannanicker, *B.* 23, 3408)

Tetra-methyl-succinic acid $C_8H_{14}O_4$, *see* CO_2H CM_2 , CM_2 , CO_2H *Hexane dicarboxylic acid* Mol w 174 [192] Formed, together with the isomeric tri-methyl-glutaric acid [97], by heating α -bromo isobutyric ether (3 pts) with dry silver powder (2 pts) at 125° for 8 hours. The product is fractionally distilled, and the fraction 200°–250° saponified by heating with $HBrAq$ at 100°. The resulting acids may be separated by steam distillation, tetra methyl-succinic acid alone passing over (Hell, *B* 7, 320, 10, 2229, Auwers & V Meyer, *B* 22, 2014, 3005, 23, 299)

Properties—Short branching crystals, melting at 200° when quickly heated, and 192° when slowly heated, sl sol cold water, m sol hot water, ether, chloroform, and CS_2 , v sol alcohol and benzene, almost insol ligrom

Anhydride [147°] (230.5) Formed by heating the acid alone, with $HClAq$ at 200°, or with $AcCl$ at 100°. Formed also by treating the acid (6 g) with red phosphorus (8 g) and bromine (16 g). Slender needles (from ligrom). May be sublimed. Soon becomes resinous. Nearly insol cold water and cold Na_2CO_3Aq , slowly dissolved by these liquids on heating, being converted into the acid

DI-METHYL-SUCCINIMIDINE $C_6H_{11}N_2$, *see*



is formed by the action of methylamine on succinimido-ether— $BHCl$ glistering prisms, [248°], v sol water, sl sol alcohol (Pinner, *B* 16, 1658)

METHYL-SUCCINURIC ACID *Amide*

NH_2 , CO NMe , CO CH_2 , CH_2 , $CONH_2$ (?) [205°–207°] Formed from methyl succinyl-urea CO $\begin{array}{l} \nearrow NMe \\ \searrow NH \end{array}$ CO CH_2 and alcoholic NH_3 at 100° (Menschutkin, *A* 178, 210) Plates, v sl sol boiling alcohol

DI-METHYL-SULPHAMIC ACID $C_2H_7NSO_3$, *see* NMe_2 , SO_2 , OH [165°] Formed, together with hydrogen di methyl ammonium sulphate NMe_2H , O SO_2 , OH , by boiling its chloride with water (Behrend, *B* 15, 1610, *A* 222, 130) Large six sided plates (from alcohol), v sol water, m sol alcohol, sl sol ether. By boiling with water, alkalis, or dilute nitric acid, it is converted into $(NMe_2H)_2SO_3H$. It expels CO_2 from carbonates

Salts— BaA' , aq plates, v sol water— PbA' , aq— AgA' , aq v e sol water, ppd by adding ether to its alcoholic solution

Ethyl ether EtA' From the chloride and $NaOEt$ Oil

Chloride NMe_2 , SO_2 , Cl (183° at 760 mm., 114° at 75 mm.) Formed by heating dimethylamine hydrochloride (1 mol) with SO_2Cl_2 (1½ mols) on the water-bath, the yield being 50 p c of the theoretical. The product is mixed with water, and the chloride extracted with ether, shaken with aqueous Na_2CO_3 , dried over $CaCl_2$, and distilled (Behrend). Golden yellow oil, partially decomposing on distillation with evolution of HCl . V sol alcohol, ether, benzene, and chloroform, insol water, $HClAq$, and $KOHAg$

Boiling water decomposes it into HCl , H_2SO_4 , NMe_2 , SO_2 , OH , and dimethylamine. Sodium-amalgam reduces it to H_2S , dimethylamine, and H_2SO_4 . Tin and $HClAq$ act in like manner. Zinc dust yields tetra methyl sulphamide

Amide v **DI-METHYL-SULPHAMIDE**

Dimethylamide v **TETRA-METHYL-SULPHAMIDE**

Di-ethyl-amide NMe_2 , SO_2 , NEt_2 (229°) From the chloride and NEt_2H Oil, partially decomposed on distillation

***n*-DI-METHYL-SULPHAMIDE**

NMe_2 , SO_2 , NH_2 [96°] Formed by passing NH_3 into the chloride of di methyl sulphamic acid (Behrend, *B* 15, 1611, *A* 222, 126) Six sided prisms with pyramidal ends, v sol water and alcohol, m sol ether

***s* Di methyl-sulphamide** $NHMe$, SO_2 , $NHMe$ [78°] Prepared by the action of methylamine on SO_2Cl_2 in ethereal solution at 0° (Franchimont, *R T C* 3, 418) Prisms, v e sol water and alcohol, v sl sol benzene. Tastes sweet. Nitric acid converts it into SO , $(NMe_2NO)_2$

Tetra methyl sulphamide SO , $(NMe_2)_2$ [73°] Formed from SO_2Cl_2 and $NHMe_2$ dissolved in chloroform (Behrend, *B* 14, 722, *A* 222, 119) Formed also from NMe_2 , SO_2 , Cl and $NHMe$. Colourless plates (from alcohol), v sol alcohol and ether, v sl sol water, aqueous acids and alkalis. May be sublimed. Dry HCl at 120° decomposes it into NMe_2 , SO_2 , Cl and $NHMe$. Conc HNO_3 yields NMe_2NO_2 (Franchimont, *R T C* 3, 420)

METHYL SULPHATES **Mono-methyl sulphate** CH_3O , SO_2 , OH *Methyl sulphuric acid* Formed by mixing methyl alcohol (1 pt) with H_2SO_4 (2 pts), allowing the hot mixture to cool, diluting with water, adding $BaCO_3$, filtering and evaporating. The barium salt thus obtained is then decomposed by H_2SO_4 (Dumas & Pélégot, *A Ch* [2] 58, 54, 61, 199, *A* 15, 40, Kane, *P M* 7, 397) Formed also from Cl , SO , OH and methyl alcohol (Claesson, *J pr* [2] 19, 240) Liquid, not solidified at -30° , v e sol water, m sol alcohol, miscible with dry ether. Yields Me_2SO , and H_2SO_4 on distillation. When the potassium salt is heated with potassium acetate methyl acetate is formed, methyl ethers of other acids are formed in like manner

Salts— $KMeSO_3$, aq very deliquescent monoclinic tables, $a b c = 742$ 779 1, $\beta = 86^\circ 51'$ (Schabus, *J* 1854, 552)— $Ca(MeSO_3)_2$ very deliquescent octahedra— $Ba(MeSO_3)_2$ 2aq monoclinic tables, $a b c = 824$ 1 907 1, $\beta = 83^\circ 30'$ *S G* 192 2 273— $Pb(MeSO_3)_2$ aq long deliquescent prisms, v sol water— $UO_2(MeSO_3)_2$ aq very deliquescent crystals (Pélégot, *A* 56, 231)

Chloride MeO , SO_2 , Cl (132.5°) at 722 mm. Formed from SO_2Cl_2 (1 mol) and $MeOH$ (1 mol) (Behrend, *J pr* [2] 15, 32) Formed also from $MeOCl$ and SO_2 (Sandmeyer, *B* 19, 861) Pun gent oil, decomposed by water into HCl and MeO , SO_2 , OH

Di-methyl sulphate Me_2SO_4 . Mol w 126 (188°) *S G* 21 1 324 (*D* & *P*)

Formation—1 From Me_2O and SO_3 —2 By dry distillation of $MeHSO_4$ (Dumas & Pélégot, Claesson, *J pr* [2] 19, 244, *B* 13, 1699)—3. By distilling methyl alcohol (1 pt) with conc H_2SO_4 (9 pts), washing the distillate with water,

drying the oily layer with CaCl_2 , and rectifying (Dumas & Péligot, *A Ch* [2] 58, 32)

Properties—Oil, decomposed by boiling water and by alkalis into MeOH and H_2SO_4 . An ethereal solution of NH_3 forms $\text{MeO SO}_2 \text{ONH}_2\text{Me}$. Distillation with fused NaCl forms MeCl and Na_2SO_4 . Distillation with KOBz yields MeOBz and K_2SO_4 . Sodium formate yields methyl formate and sodium sulphate. It forms double compounds with sulpho acetates, sulpho benzoates, and isethionates (Geuther, *A* 218, 288)

METHYL SULPHIDE $(\text{CH}_3)_2\text{S}$ Mol w 62 (41°) (R), (37°) (K) SG ⁴⁰ 845 H.F.p 12,730 H.F.v 11,570 SV 75.6 (Lossen, *A* 254, 71) Formed by passing MeCl into a solution of K_2S in MeOH (Regnault, *A Ch* [2] 71, 391, *A* 34, 26) Prepared by distilling a concentrated solution of MeNaSO_4 (from 500 c.c. MeOH) with aqueous KOH (500 g) that has been previously half saturated with H_2S , the yield being moderate (150 g). Colourless mobile liquid with very unpleasant odour (Klaason, *B* 20, 3407)

Reactions—1 Takes fire when dropped into dry chlorine, but if the temperature be kept low only substitution products $(\text{CH}_3\text{Cl})_2\text{S}$, $(\text{CH}_3\text{Cl})_3\text{S}$, and $(\text{CCl}_3)_2\text{S}$ may be obtained (Riche, *A Ch* [3] 43, 283) The compound $(\text{CCl}_3)_2\text{S}$ boiled at 156°–160° and gave a V D 5.68 (calc 9.41) —2 *Iodoacetic ether* forms $\text{S}(\text{CH}_2\text{CO}_2\text{Et})_2$, tri-methyl-sulphine iodide, and $\text{Me}_3\text{S}(\text{CH}_2\text{CO}_2\text{Et})_2$ (Letts, *Tr E* 28, 618) —3 *Bromoacetic ether* forms $\text{Me}_3\text{SBrCH}_2\text{CO}_2\text{Et}$ which crystallises in pearly scales and yields when treated with moist Ag_2O unstable $\text{Me}_3\text{S}(\text{OH})\text{CH}_2\text{CO}_2\text{Et}$ (Letts)

Combinations— Me_3SBr Crystals (Cahours, *A* 135, 355) When dissolved in water it gives off HBr . When its alcoholic solution is treated with zinc and the product evaporated and mixed with HgCl_2 , there is formed a pp of $(\text{SMe}_2)_2\text{HgCl}_2\text{ZnBr}$ (Patein, *Bl* [2] 50, 201) — Me_3SiI Crystals resembling iodine — Me_3SHgCl , — $(\text{Me}_3\text{S})_2\text{PtCl}_2$ (Lorr, *A* 107, 234) Yellow crystalline powder Melts with decomposition at 218° (Blomstrand, *J pr* [2] 38, 365) — $(\text{Me}_3\text{S})_2\text{PtCl}_2$ [159°] Formed by the action of Me_3S on potassium platinum chloride at 60° (Blomstrand, *J pr* [2] 38, 358) Exists in two allotropic forms, crystallising in lemon yellow monoclinic crystals, and in yellow dimetric plates (containing CHCl_3) — $(\text{Me}_3\text{S})(\text{Et}_3\text{S})\text{PtCl}_2$ Formed from $(\text{Et}_3\text{S})_2\text{PtCl}_2$ and Me_3S — $(\text{Me}_3\text{S})_2\text{PtCl}_2\text{Br}_2$ — $(\text{Me}_3\text{S})_2\text{PtBr}_4$ reddish brown monoclinic crystals (from chloroform) — $(\text{Me}_3\text{S})_2\text{PtBr}_2$ Formed from $(\text{Me}_3\text{S})_2\text{PtSO}_4$ and KBr (B) Yellow monoclinic crystals — $(\text{Me}_3\text{S})_2\text{PtI}_2\text{Cl}_2$ greenish black crystalline powder — $(\text{Me}_3\text{S})_2\text{PtI}_2\text{Br}_2$ — $(\text{Me}_3\text{S})_2\text{PtI}_4$ black crystalline powder — $(\text{Me}_3\text{S})_2\text{PtI}_2$ [172°] Formed from $(\text{Me}_3\text{S})_2\text{Pt}(\text{NO}_3)_2$ and KI (B) Red crystals — $(\text{Me}_3\text{S})_2\text{Pt}(\text{NO}_3)_2$ [156°] Formed from $(\text{Me}_3\text{S})_2\text{PtCl}_2$ and AgNO_3 (B) Small brownish needles — $(\text{Me}_3\text{S})_2\text{Pt}(\text{NO}_3)_2$ Formed from $(\text{Me}_3\text{S})_2\text{PtSO}_4$ and potassium nitrite (B) Small white plates, soluble in chloroform — $(\text{Me}_3\text{S})_2\text{PtSO}_4$ 2aq [91°] Formed from silver sulphate and $(\text{Me}_3\text{S})_2\text{PtCl}_2$ Yellowish crystalline mass, v sol water — $(\text{Me}_3\text{S})_2\text{PtCrO}_4$ reddish-brown pp got by adding K_2CrO_4 to a solution of $(\text{Me}_3\text{S})_2\text{PtSO}_4$. Sl sol water, insol alcohol and chloroform

Methylo-sodide SMe_2I **Tri-methyl-sulphine iodide** Formed, even in the cold, by the

union of Me_3S with MeI (Cahours, *C R* 80, 1817, 81, 1163, *A Ch* [5] 10, 13, *A* 135, 355, 136, 151) Formed also by heating MeI at 100° with ppd As_2S_3 or with Na_2S (Klinger, *B* 15, 881, *A* 252, 357) and by heating Se_2I with MeOH at 140° (Klinger & Maassen, *A* 252, 252) Large prisms (from water), v sol hot water, sl sol alcohol, insol ether Its aqueous solution is partially decomposed on evaporating at 100°, forming Me_3S and iodine Moist Ag_2O yields SMe_2OH , a strongly alkaline base whence the other salts may be prepared by neutralisation with acids An aqueous solution of SMe_2I gives with alcoholic HgCl_2 a pp of Me_3SIHgI which forms pale yellow needles, nearly insol water and ether, sol alcohol (Patein, *Bl* [3] 2, 159) Bromine forms Me_3SIBr , which separates from hot alcohol in orange-red crystals [95°] which in alcoholic solution give with platinum chloride a pp of $(\text{Me}_3\text{SiCl})_2\text{PtCl}_4$ (Dobbin & Masson, *C J* 47, 56) Tri-methyl sulphine di-bromo-iodide is converted by dry NH_3 into $\text{Me}_3\text{SIBr.N.H}_3$, an amorphous light green mass [75°–80°] Chlorine forms Me_3SiCl , which separates from alcohol in yellow crystals [104°], converted by aqueous ammonia into iodide of nitrogen and by gaseous ammonia into $\text{Me}_3\text{SiCl.2NH}_3$, an unstable compound which loses ammonia and absorbs water when exposed to air (D & M) The compound $(\text{Me}_3\text{Si})_2\text{As}_2\text{I}_4$ is formed by heating As_2S_3 with MeI at 100° and crystallises in blue black needles (Klinger & Maassen, *A* 252, 260) The compound $\text{Me}_3\text{SISnI}_2$ crystallises in yellow needles, sl sol cold water The compound $(\text{Me}_3\text{Si})_2\text{CdI}_2$ formed by heating CdS with MeI at 100°, crystallises from water or alcohol in white needles, melting at 185° when slowly heated and 195° when quickly heated, and converted by aqueous CdI_2 into $\text{Me}_3\text{SiCdI}_2$ [168°]

Methylo-chloride Me_3SiCl From the base and HCl Deliquescent prisms Gives the salts $(\text{Me}_3\text{SiCl})_2\text{PtCl}_2$ crystallising from hot water in sparingly soluble orange-yellow prisms and Me_3SAuCl crystallising in thick prisms, v e sol water Me_3SiCl shaken with an ethereal solution of iodine yields reddish black crystals of $\text{Me}_3\text{Si}_2\text{Cl}$, which is probably also formed from Me_3SiI and ICl (Dobbin & Masson) Me_3SiCl is converted by ICl into Me_3SiCl_2 Dry chlorine forms Me_3SiCl_2 , a yellow liquid, solidifying on exposure to air Decomposed by water, alcohol, and ether, into Me_3SiCl and chlorine

Methylo-bromide Me_3SBr Formed from $\text{Me}_3\text{S OH}$ and HBr Also formed from Me_3S and MeBr Prisms, v sol water Converted by iodine in ethereal solution into Me_3SIBr . With ICl it forms $\text{Me}_3\text{SiClBrI}$ as yellow crystals [87°], completely decomposed at 190°

Methylo-sulphhydrate $\text{Me}_3\text{S SH aq}$ From $\text{Me}_3\text{S OH}$ and H_2S (Brown & Blaikie, *J pr* [2] 23, 395)

Methylo-sulphide $(\text{Me}_3\text{S})_2\text{S}$ **Tri-methyl-sulphine sulphide** An aqueous solution of this body may be got by saturating one half of a conc solution of $\text{Me}_3\text{S OH}$ with H_2S , and adding the other half This solution, if allowed to evaporate in dry air or in coal gas, when it reaches a certain strength forms Me_3S , thus $(\text{Me}_3\text{S})_2\text{S} = 3\text{Me}_3\text{S}$ (Crum Brown & Blaikie, *Pr E* 9, 563, *C N* 37, 130) On gently heating a solution of $(\text{Me}_3\text{S})_2\text{S}$ in a sealed tube, Me_3S

separates as an upper layer. The aqueous solution has the characters of an alkaline sulphide, dissolving sulphur (forming $(\text{Me}_2\text{S})_2\text{S}_2$) and Sb_2S_3 . Acids decompose it with evolution of H_2S .

Methylo-thiosulphate $(\text{Me}_2\text{S})_2\text{S}_2\text{O}$, aq. Formed by exposing an aqueous solution of $(\text{Me}_2\text{S})_2\text{S}$ to atmospheric oxidation. Formed also by exposing $(\text{Me}_2\text{S})_2\text{S}_2$ to air. Hygroscopic four-sided prisms, sl sol alcohol. Decolourises a solution of iodine. At 135° it is decomposed into Me_2S and $\text{Me}_2\text{S O SO}$, SMe , a white crystalline mass $[100^\circ]$ which does not act upon iodine but slowly oxidises to sulphate (Crum Brown & Blaikie, *J pr* [2] 23, 395).

Methylo-sulphate $(\text{Me}_2\text{S})_2\text{SO}_3$, aq. Formed from the hydroxide by saturating one half of its solution with SO_2 , and adding the other half (Crum Brown & Blaikie, *Pr E* 9, 712). Crystals. Gives off water of crystallisation at 140° . At 170° it gives off Me_2S , leaving a liquid residue, which solidifies on cooling, and is apparently $(\text{Me}_2\text{S})_2\text{SO}_3\text{Me}$.

Methylo-dithionate $(\text{Me}_2\text{S})_2\text{S}_2\text{O}_6$, 6aq. Formed from $\text{Me}_2\text{S OH}$ and dithionic acid. Deliquescent cubes, insol alcohol. Decomposes at 220° into SO_2 and $(\text{Me}_2\text{S})_2\text{SO}_3$, and the latter then further decomposes into Me_2S and $\text{Me}_2\text{S SO Me}$.

Methylo-metaphosphate $\text{Me}_2\text{S PO}_3$. From AgPO_3 and Me_2SI . Hygroscopic glassy mass, giving off Me_2S on heating.

Methylo-ferrocyanide $(\text{Me}_2\text{S})_2\text{FeCy}_2$, 9aq. From Me_2SI and silver ferrocyanide. Transparent green plates, which lose their water in a desiccator, and then give off Me_2S at 220° .

Methylo-ferricyanide $(\text{Me}_2\text{S})_2\text{FeCy}_2$, 7½aq. From Me_2SI and silver ferricyanide (Crum Brown & Blaikie, *Pr E* 10, 253). Pale orange efflorescent plates.

The **methylo-chromate** and **methylo-iodate** explode at 100° .

Methylo-carbonate $(\text{Me}_2\text{S})_2\text{CO}_3$. Formed from Me_2SI and Ag_2CO_3 , the liquid being evaporated to a syrup and left to crystallise over H_2SO_4 . Deliquescent prisms, with strong alkaline reaction. At 100° it is split up into CO_2 , water, Me_2S , and methyl alcohol.

Methylo-oxalate $(\text{Me}_2\text{S})_2\text{C}_2\text{O}_4$, aq. Formed from $\text{Ag}_2\text{C}_2\text{O}_4$ and Me_2SI . Deliquescent crystals. Split up at 140° into Me_2S and methyl oxalate.

Methylo-acetate $\text{Me}_2\text{S OAc}$. Formed from Me_2SI and AgOAc (Crum Brown & Blaikie, *Pr E* 10, 53, *C N* 39, 51). Split up at 100° into Me_2S and MeOAc .

Methylo-benzoate $\text{Me}_2\text{S OBz}$. From the iodide and AgOBz . Thin plates (from alcohol). Decomposed by heat into Me_2S and MeOBz .

Ethylo-iodide SMe.EtI . **Di-methyl-ethyl-sulphide iodide** $[110^\circ]$. Formed either from Me_2S and EtI or MeEtS and MeI (Klinger & Maassen, *A* 243, 212, 252, 246, *cf* Kruger, *J pr* [2] 14, 193). Hygroscopic crystalline mass, v sol alcohol, insol ether. Yields on distillation Et_2SI and Me_2SI . In alcoholic solution it reacts with AgCy at 90° , forming SMe.EtCyAgCy , a deliquescent crystalline body, v sol alcohol, insol ether, and decomposed by heat into AgCy and SMe.EtCy (Patein, *C R* 106, 861). Forms the following compounds — $(\text{Me}_2\text{SEtI})_2\text{CdI}_2$, $[180^\circ]$, crystallising in small

needles, sl sol water — $\text{Me}_2\text{SEtICdI}_2$, $[99^\circ]$, crystals falling from hot conc CdI_2 aq in long needles. — $\text{Me}_2\text{SEtIHgI}_2$, $[66^\circ]$ (Patein, *Bl* [3] 2, 159).

Ethylo-chloride Me_2SEtCl . The following compounds of this body have been prepared (Klinger & Maassen, *A* 243, 212) — $\text{Me}_2\text{SEtCl}(\text{HgCl}_2)$, $[119^\circ]$, long needles, insol. hot water — $\text{Me}_2\text{SEtCl}(\text{HgCl}_2)_2$, $[200^\circ]$, white crystalline powder, slightly soluble in water — $(\text{Me}_2\text{SEtCl})_2\text{PtCl}_4$, $[c\ 213^\circ]$, small orange crystals, sl sol water, insol alcohol and ether — $\text{Me}_2\text{SEtClAuCl}_2$, $[244^\circ]$, long yellow needles, m sol hot water.

Di-methyl disulphide $\text{C}_2\text{H}_6\text{S}_2$, *re* Me_2S_2 , (117°) (C), $(112^\circ$ at 744 mm) (Pierre, *A* 80, 128) *SG* 1.046 (C) *VD* 3.30 *SV* 100.6 (Lossen, *A* 254, 71). Formed by passing MeCl through an alcoholic solution of K_2S_2 , or by distilling $\text{Ca}(\text{SO}_3\text{Me})_2$ with K_2S_2 (Cahours, *A Ch* [3] 18, 157, *A* 61, 92). Liquid, with intolerable odour of onions, v sl sol water, miscible with alcohol and ether. Burns with blue flame. Chlorine converts it first into crystalline $\text{Me}_2\text{S}_2\text{Cl}_2$, and finally into a liquid mixture of $(\text{CCl}_3)_2\text{S}$ and SCl_2 (Riche, *A* 92, 356). Dilute nitric acid converts it into methyl methane thiosulphonate MeSO_2SMe .

Derivative — *v* **PER-CHLORO-METHYL DI-SULPHIDE**.

Di-methyl trisulphide Me_2S_3 , (170°) at 760 mm *SG* 1.2162, $\frac{1}{2}$ 1.2059, $\frac{1}{3}$ 1.199. Formed together with Me_2S and S from methyl mercaptan and S_2Cl_2 (Klason, *B* 20, 3414). Formed also from MeCl and K_2S_3 (Cahours). Pale yellow liquid, with very disagreeable odour.

Derivative — *v* **HEXA-BROMO DI-METHYL TRI-SULPHIDE**.

METHYL SULPHITE $\text{C}_2\text{H}_5\text{SO}_3$, *re* $\text{SO}(\text{OMe})_2$, (121.5°) *SG* 1.0456 *VD* 3.68 (calc 3.80). Formed by the action of methyl alcohol on S_2Cl_2 or on SOCl_2 (Carius, *A* 110, 209, 111, 93). Colourless liquid, with pleasant odour, miscible with alcohol and ether. It dissolves slowly in water with evolution of SO_2 and formation of MeOH .

METHYL SULPHOCYANIDE $\text{C}_2\text{H}_5\text{NS}$, *re* MeSCy , (133°) *SG* 1.115 (C), $\frac{1}{2}$ 1.069 (Nasini & Scala, *G* 17, 66) *R_n* 33.8 *SV* 78.1 (Lossen, *A* 254, 73) *H F p* — 31,410 *H F v* — 31,990 (Thomsen, *Th*). Obtained by distilling potassium sulphocyanide with calcium methyl sulphate (Cahours, *A Ch* [3] 18, 261, *A* 61, 95). Liquid with alliacious odour, v sl sol water, miscible with alcohol and ether. Boiling nitric acid oxidises it to methane sulphonic acid. Chlorine acts upon it according to the equation — $3\text{MeSCN} + 11\text{Cl}_2 = \text{Cl}_3\text{Cy}_2 + 2\text{CCl}_4 + \text{CCl}_3\text{Cl} + 9\text{HCl}$ (James, *J pr* [2] 85, 462). Cold aqueous KOH does not attack it, but alcoholic potash forms Me_2S , ammonia, KCy , and K_2CO_3 . Alcoholic KSH forms KSCy and Me_2S . When heated at 180° it partially changes to methyl thiocarbimide.

Methyl polysulphocyanide $(\text{CH}_3\text{CNS})_n$ (?) $[188^\circ]$. Prepared by heating methyl sulphocyanide to 180° with a trace of HCl (Hofmann, *B* 13, 1349). Sublimable. Colourless crystals. Sol acetic acid, insol acids and alkalis. By heating with alcoholic NH_3 to 150° it gives a well-crystallising base.

METHYL-SULPHONAMIDES v. METHYL**SULPHAMIDES**

DI-METHYL-SULPHONE $\text{O}_2\text{H}_2\text{SO}_2$ *ie* Me_2SO_2 . Mol w 94 [109°] (238°) R_{∞} 82.09 in a 2.46 aqueous solution (Kanonnikoff, *J R* 15, 451) Formed by oxidising Me_2S with fuming HNO_3 (Saytzeff, *A* 144, 148) or with aqueous KMnO_4 (1.30) (Beckmann, *J pr* [2] 17, 454) Thick needles (from alcohol) Not affected by reducing agents

DI METHYL-SULPHONE DI-CARBOXYLIC ACID $\text{O}_2\text{S}(\text{CH}_2\text{CO}_2\text{H})_2$ *Sulphono di acetic acid* [182°] Formed by oxidation of thio di glycollic (sulphido diacetic) acid in alkaline solution with KMnO_4 (Lovén, *B* 17, 2817) Long trimetric tables V sol water, alcohol, and ether At 200° it splits up into di methyl sulphone and CO_2

Salts — $\text{A}^+\text{Ba}^+\text{aq}$ fine felted sparingly soluble needles — $\text{A}^+\text{Ba}^+\text{aq}$ small prisms

Ethyl ether A^+Et , thick oil

Amide $\text{O}_2\text{S}(\text{CH}_2\text{CONH}_2)_2$ glistening plates, v sol hot water, slightly in cold

METHYL-SULPHONIC ACID v METHANE SULPHONIC ACID

METHYL SULPHOXIDE $(\text{CH}_3)_2\text{SO}$ The nitrate $(\text{CH}_3)_2\text{SOHNO}_3$ is formed by oxidising methyl sulphide with conc HNO_3aq and crystallises in deliquescent needles, whence BaCO_3 sets free the oxide (Saytzeff, *A* 144, 148) Oil, v sol water Solidifies when strongly cooled Reduced by zinc and H_2SO_4 to methyl sulphide Me_2S

METHYL-SULPHURIC ACID v METHYL SULPHATES**METHYL SULPHYDRATE v METHYL MERCAPTAN****METHYL-TARCONIC ACID v NARCOTINE**

DI-METHYL TARTARIC ACID $\text{C}_4\text{H}_6\text{O}_6$ *ie* $\text{CO}_2\text{H CMe}(\text{OH})\text{CMe}(\text{OH})\text{CO}_2\text{H}$ *Di oxy di methyl succinic acid* Formed, together with lactic acid, by the action of zinc and a little HCl aq on pyruvic acid in alcoholic solution (Böttger, *A* 188, 315, *B* 9, 1064, 1621) Syrup — KHA^+ small six sided plates, v sl sol water — KA^+ needles, m sol water — BaA^+ $3\frac{1}{2}$ aq prismatic needles, sl sol water — CaA^+ crystalline pp, nearly insol water

METHYL-TAURINE v METHYL AMIDO ETHANE SULPHONIC ACID**METHYL-TAURO GAMINE v METHYL GUANIDO ETHANE SULPHONIC ACID****METHYL TEREPHTHALIC ACID v TOLUENE DICARBOXYLIC ACID**

Di methyl terephthalic acid *v* XYLENE DICARBOXYLIC ACID

METHYL-TELLURIDE v TELLURIUM, ORGANIC COMPOUNDS

METHYL TETRADECYL KETONE $\text{C}_{15}\text{H}_{32}\text{O}$ *ie* $\text{CH}_3\text{CO C}_{13}\text{H}_{27}$ [43°] (231°) Formed by distilling a mixture of barium pentadecanoate and barium acetate (Krafft, *B* 15, 1707) Yields myristic and acetic acids on oxidation

DI-METHYL THETINE $\text{C}_4\text{H}_8\text{SO}_4$ *ie*

$\text{CH}_3\text{C}(\text{SM}_2)\text{C}(\text{O})\text{O}$ *Anhydride of the methylohydroxide of the methyl derivative of thio glycollic acid* Obtained by decomposing its hydrobromide with moist Ag_2O , or its sulphate with

baryta (Crum Brown & Letts, *Tr E* 28, 571, *B* 6, 1384, 7, 695) Crystals (containing aq) Hygroscopic, but gives up the water over H_2SO_4 V sol water, m sol alcohol Neutral in reaction Decomposes when heated into CO_2 and $(\text{Me}_2\text{S})_2\text{CO}$, the methylo-carbonate of di methyl sulphide Yields di methyl sulphone when oxidised by KMnO_4 The following compounds may be styled its salts — $\text{Me}_2\text{SBr CH}_2\text{CO}_2\text{H}$ Formed from bromo-acetic acid and Me_2S Large deliquescent rectangular plates (from alcohol) Acid to litmus With lead oxide it forms $\text{C}_4\text{H}_8\text{SO}_4\cdot 2\text{PbBr}_2$ Decomposed by heat or by boiling alcohol into SMe_2Br , methyl bromide, and $\text{S}(\text{CH}_2\text{CO}_2\text{H})_2$ (Letts, *Tr E* 28, 591) Yields methane sulphonic acid on oxidation (Letts, *Tr E* 28, 601) — $(\text{C}_4\text{H}_8\text{SO Br})_2\text{PtBr}_2$ (?) dark red crystals — $\text{Me}_2\text{SOCl CH}_2\text{CO}_2\text{H}$ Formed from the sulphate and BaCl_2 Deliquescent crystals, v sol water, v sl sol alcohol — $(\text{C}_4\text{H}_8\text{SO}_4)_2\text{HI}$ (?) — $\text{Me}_2\text{SOCl CH}_2\text{CO}_2\text{H}$ Formed by leaving di methyl thetine in contact with dilute HIAq Crystals, insol water, sol alcohol and ether — $\text{SO}_4(\text{SMe CH}_2\text{CO}_2\text{H})_2$ *Di methyl thetine sulphate* Formed from $\text{Me}_2\text{SBr CH}_2\text{CO}_2\text{H}$ and silver sulphate Crystalline mass, sol water, v sl sol alcohol Acid to litmus When heated over 140° it fuses and splits up into CO_2 and $(\text{Me}_2\text{S})\text{SO}_4$, the methylo sulphate of di methyl sulphide — $\text{Me}_2\text{S}(\text{NO}_2)_2\text{CH}_2\text{CO}_2\text{H}$ From $\text{Me}_2\text{SBr CH}_2\text{CO}_2\text{H}$ and AgNO_3 Colourless crystals with acid reaction Yields di methyl sulphone when oxidised by conc HNO_3

METHYL-THIALDINE $\text{C}_4\text{H}_7\text{NS}$ [79°] Obtained on treating crude thio acetic aldehyde with an aqueous solution of methylamine (W Markwald, *B* 19, 2378) Long needles (from alcohol), insol water, sl sol cold alcohol, v e sol hot alcohol and ether Its solutions possess a feeble alkaline reaction Volatilises in steam with partial decomposition With acids it forms very soluble salts

μ - METHYL - THIAZOLE $\text{C}_3\text{H}_5\text{NS}$ *ie* $\text{S CMe} \gg \text{N}$ (128° cor) Formed by boiling CH CH_2 (128° cor) Formed by boiling thioacetamide $\text{CH}_3\text{CS NH}_2$ with chloro acetal (Hantzsch *A* 250, 270), or with di chloro di ethyl oxide (Hantzsch, *B* 21, 943) Mobile liquid, miscible with water Its hydrochloride and hydrobromide form hygroscopic needles. With mercuric chloride it forms compounds melting at 112° and 154° $\text{B}^+\text{H PtCl}_6$ [199°] Hexagonal plates or flat needles (from water) — Picrate [146] Yellow needles, sl sol water, v sol alcohol and benzene

(a)-Methyl-thiazole $\text{S} \begin{array}{c} \text{---CH} \\ \text{CH CMe} \end{array} \gg \text{N}$ (132°) Obtained by distilling oxy methyl thiazole (from chloro acetone and metallic sulphocyanides) with zinc dust (Hantzsch *A* Arapides, *B* 21, 942), *A* 249, 24) Formed also by boiling with alcohol the diazo compound derived from amido methyl thiazole (from chloro acetone and thio urea) (Popp, *A* 250, 277) Colourless liquid, sinking under water, but slowly dissolving, v sol alcohol and ether The hydrochloride is deliquescent. It forms two compounds with HgCl_2 , melting at 119° and 148° The aurochloride melts at 185°, and the picrate at 174° — $\text{B}^+\text{H}_2\text{PtCl}_6$ [204°] Orange prisms, m sol water, sl sol alcohol.

$\alpha\mu$ Di-methyl-thiazole C_4H_6NS $\epsilon\epsilon$
 $S \text{ CMe} \searrow N$ (145° cor) SG 15 1 0601 Formed from thioacetamide and chloro-acetone (Hantzsch, B 21, 943, A 250, 265) Colourless liquid, more sol cold than hot water Reduced in alcoholic solution by sodium to ethylamine and propyl mercaptan (Schatzmann, A 261, 1) — $B'H_2PtCl_6$ [216°] Prisms, m sol water — $B'HCl(HgCl_2)_4aq$ [110°] White needles, v e sol water — $B'(HgCl_2)_2$ [177°] Sl sol cold water, v e sol dilute $HClAq$ — *Picrate* [138°] *Methylo-sodide* $B'MeI$ Pointed crystals, v sol water

$\beta\mu$ Di-methyl-thiazole $S \text{ CMe} \searrow N$ (150° cor) Formed by condensing thioacetamide with (α)-chloro propionic aldehyde (Hubacher, A 259, 240) Oil, v sl sol water, v sol alcohol and ether Volatile with steam — $B'H_2PtCl_6$ [202°] — *Picrate* [167°]

Tri-methyl-thiazole C_5H_8NS $\epsilon\epsilon$
 $S \text{ CMe} \searrow N$ (167°) SG 15 1 013 Formed by heating thioacetamide with chloro methyl ethyl ketone at 100° (Poubelle, A 259, 258) Liquid, m sol cold, insol hot, water — $B'HCl$ [174°] — $B'H_2PtCl_6$ [233°] — *Aurochloride* [156°] — *Picrate* [133°] *Mercuric double chloride* [119°]

References — METHYL-AMIDO METHYL-THIAZOLE, METHYL-IMIDO-DI-METHYL-THIAZOLE, OXY METHYL THIAZOLE, and TOLYL AMIDO METHYL-THIAZOLE

METHYL-THIAZOLE CARBOXYLIC ACID

$C_4H_5NSO_2$ $\epsilon\epsilon$ $S \text{ CMe} \searrow N$ [145°] Formed by heating the dicarboxylic acid [169°] at 171° (Roubleff, A 259, 271) Small needles or prisms, v sol cold, v e sol hot, water, m sol alcohol, sl sol ether, almost insol benzene

Methyl-thiazole carboxylic acid

$CO_2H \text{ CMe} \searrow N$ [257°] Formed by saponifying its ether with alcoholic potash (Wohmann, A 259, 299) Pearly plates (from water) or needles (from alcohol), sl sol ether and hot water, almost insol benzene

***Ethyl ether* EtA'** [28°] (233° i V) at 726 mm Obtained from amido methyl thiazole carboxylic ether by diazotisation, conversion into chloro methyl thiazole carboxylic ether [51°] and reduction of this by zinc dust and $HOAc$ Flat prisms, volatile with steam

Methyl-thiazole dicarboxylic acid

$S \text{ C(CH}_3)_2 \searrow N$ [169°] Formed by condensing thioacetamide with chloro oxalacetic ether and saponifying with alcoholic soda (Roubleff, A 259, 268) Long white needles, v sol cold water, sl sol ether and benzene — BaA'' 2aq needles. — HgA'' 3½aq crystalline pp

Di-methyl thiazole carboxylic acid

$S \text{ CMe} \searrow N$ [227°] Formed by saponifying its ether (R) Silky needles or small prisms, sl sol hot water, m sol alcohol and ether May be sublimed Yields on distillation with lime di-methyl thiazole — AgA' white needles, sol hot water — $HA'HCl$ plates

***Ethyl ether* EtA'** [51°] (242° cor)

Formed from thioacetamide and chloro acetoacetic ether (Hantzsch, A 250, 269) Needles (from ether), insol water, v sol alcohol and ether

METHYL-THIAZYL-PROPIONIC ETHER

$S \text{ CMe} \searrow N$ Formed by condensing $CH \text{ O CHMe CO}_2Et$ bromo methyl-acetoacetic ether with thioacetamide in the cold (Roubleff, A 259, 262) Thick oil When saponified and heated with lime it yields methyl ethyl thiazole

($\alpha\alpha'$)-METHYL-THIENYL-GLYOXYLIC ACID

$C_8H_8SO_2$ $\epsilon\epsilon$ $S \text{ C(CH}_3)_2 \searrow N$ [80°] Formed by oxidising ($\alpha\alpha'$) methyl thienyl methyl ketone with alkaline $KMnO_4$ (Ruffi, B 20, 1747) — CaA' 2aq needles — BaA' 2aq needles — AgA' ($\alpha\beta$) - Methyl - thienyl - glyoxylic acid $SC_6H_4Me \text{ COCO}_2H$ [142°] Formed by oxidising (a) methyl (b) thienyl methyl ketone with an aqueous solution of $KMnO_4$ and KOH (Ruffi, B 20, 1748) Needles (from water), subliming even in the cold

Phenyl hydrazide

$SC_6H_4Me \text{ C(N,HPH) CO}_2H$ [141°] Crystalline *Oxim* $SC_6H_4Me \text{ C(NO}_2\text{H) CO}_2H$ [104°]

($\alpha\alpha'$)-Di-methyl-(β)-thienyl-glyoxylic acid

$S \text{ CMe} \searrow N$ Formed by oxidation of ($\alpha\alpha'$) di-methyl (β) thienyl methyl ketone with alkaline $KMnO_4$ (Ruffi) Oil, slowly solidifying Yields leuco thiophene green when heated with di-methyl aniline and $ZnCl_2$ — AgA'

METHYL THIENYL KETONE v THIENYL

METHYL KETONE

(a) METHYL-THIENYL METHYL KETONE

C_8H_8SO $\epsilon\epsilon$ $S \text{ C(CH}_3)_2 \searrow N$ [25°] (233° cor) Formed by the action of $AcCl$ on (a) methyl thiophene in presence of $AlCl_3$ (Demuth, B 18, 3025, 19, 1859, Ernst, B 19, 3275) Large tables On oxidation with $KMnO_4$ it yields thiophene dicarboxylic acid Fuming HNO_3 forms a nitro derivative [121°] *Oxim* $C_8H_8S(NO_2H)$ [125°] Needles (from dilute alcohol)

***Phenyl hydrazide* $C_8H_8S(N_2,HPH)$** [128°] Needles (from alcohol)

(β) - Methyl - thienyl methyl ketone

$C_8H_8SMe \text{ CO CH}_3$ (216° cor) Formed from (β) methyl thiophene and $AcCl$ in presence of $AlCl_3$ (Demuth, B 18, 3025)

($\alpha\beta'$) - Di-methyl - thienyl methyl ketone

C_8H_8SO $\epsilon\epsilon$ $S \text{ CMe} \searrow N$ or $S \text{ CMe} \searrow N$ (227°) Formed by the action of $AcCl$ on ($\alpha\beta'$) di-methyl thiophene dissolved in ligroin in presence of $AlCl_3$ (Zelninsky, B 20, 2019) Liquid Coloured red by isatin and H_2SO_4

Oxim $C_8H_8S(NO_2H)$ [70°] Needles

***Phenyl hydrazide* $C_8H_8S(N_2,HPH)$** [70°]

Di-methyl-thienyl methyl ketone

$SC_6H_4Me \text{ CO CH}_3$ (224°) SG 15 1 091 Formed from the di-methyl thiophene of coal tar by treatment with $AcCl$ and $AlCl_3$ (Messinger, B 18, 2301) Liquid Gives a red colour with isatin and H_2SO_4 Oxidised by alkaline $KMnO_4$ to thiophene dicarboxylic acid

Oxim $C_8H_8S(NO_2H)$ [65°] Needles

TETRA-METHYL-THIO-ANILINE v TETRA

METHYL-DI-AMIDO-DI-PHENYL SULPHIDE

METHYL-THIOCARBAMINE CYAMIDE v.

CARBIMIDO METHYL THIO UREA

METHYL-THIOCARBAMINE-ETHYL-CY-AMIDE v ETHYL CARBIMIDO METHYL THIO UREA.**DI-METHYL-THIOCARBAZIC ACID**

C_2H_5NS , \pm NMe , $NHCS$ SH [112°] Formed from di methyl-hydrazine and CS_2 (Renouf, *B* 13, 2172) Colourless plates

METHYL-THIOCARBIMIDE CH_3NCS Mol w 73 [34°] (119°) VD 2.42 (calc 2.53) SG $\frac{4}{3}$ 1.069 R₂ 35.75 (Nasini & Scala, *G* 17, 66) HFp -24.520 HFv -25.100 HO (gas) 392,000 (Thomsen, *Th* 4, 197) Formed by the action of $AgNO_3$ or $HgCl_2$ on the product of the union of CS_2 on methylamine (Hofmann, *B* 1, 172) Formed also by heating pure methyl sulphocyanide for some time at 180°-185° (Hofmann, *B* 18, 2196) Pungent crystals Reacts with sodium cyanamide and alkyl iodides, forming methyl-alkyl cyano tho-ureas Thus $CN.HNa$ and MeI give $CH_3NHCSNCyCH_3$ [c 195°], while allyl iodide yields the compound $NHMeCSNCyC_2H_5$ [78°], propyl iodide forms $NHMeCSNCyC_3H_7$ [91°], and benzyl chloride forms $NHMeCSNCyCH_2Ph$ [173°] (Hecht, *B* 23, 1658)

METHYL THIOCARBONATES

Methyl dithiocarbonic acid $C_2H_5S_2O$ \pm CH_3OCS SH *Methyl xanthic acid* *Methyl xanthoquinic acid* *Xantho methylic acid* The potassium salt of this acid is formed by adding CS_2 to a solution of KOH in methyl alcohol (Dumas & Peligot, *A Ch* [2] 24, 55, Desains, *A Ch* [3] 20, 504) It crystallises in silky fibres, SG $\frac{15}{8}$ 1.6878 (Clarke, *B* 11, 1505) Iodine converts it into $(CH_3OCS)_2S_2$ With EtI it yields $MeOCSSEt$ (184°) - PbA_2

Methyl dithiocarbonate $CH_3OCSSCH_3$ (163°) (S), (171°) (C) SG $\frac{15}{8}$ 1.176 (S), $\frac{15}{8}$ 1.143 (O) Formed from CH_3CS SK and MeI (Salomon, *J pr* [2] 8, 117) Formed also together with CO and S by heating the compound $(CH_3OCS)_2S_2$ (*v supra*) (Cahours, *A Ch* [3] 19, 158)

Methyl trithiocarbonate Me_3CS_3 (200°-205°) SG $\frac{15}{8}$ 1.159 Formed by distilling a mixture of concentrated solutions of K_2CS_3 and $Ca(SO_4Me)$ (Cahours, *A Ch* [3] 19, 163) Yellow liquid with pungent odour, nearly insol water, miscible with alcohol and ether Combines with bromine forming red crystals of $Me_3CS_3Br_2$ (Berend, *A* 128, 833)

METHYL THIO-COUMARILIC ACID $C_6H_5(CH_3)O CO SH$

Ethyl ether $C_6H_5(CH_3)O CO SEt$ [92°], glistening yellow needles, v sol ether, sl sol alcohol Formed by heating methyl-coumarilic-ethyl-ether with P_2S_5 By treatment with alcoholic KOH it is reconverted into methyl coumarilic acid (Hantzsch, *B* 19, 2400)

METHYL-THIOFORMALDINE $C_2H_5S_2N$ \pm $(CH_2)_2S_2NMe$ [65°] (c 185°) Formed from an aqueous solution of formic aldehyde by successive addition of H_2S and methylamine (Wohl, *B* 19, 2346) Needles (from ether) with unpleasant smell, insol water, sol acids and alcohol, v sol ether Volatile with steam On boiling it is converted into a substance melting at 130°-140° - B/HCl [188°] White needles, v sol water Its solution is ppd by $AgNO_3$, $HgCl_2$, and platinum chloride.

Methylo-iodide $BMeI$ [161°-163°]. Slender needles, v sol water Gives rise to $B/MeCl$ and $(B/MeCl)_2PtCl_4$

METHYL-THIOHYDANTOIN $C_2H_5N_2SO$ \pm

$NH C \begin{smallmatrix} S CH_3 \\ NMe CO \end{smallmatrix} (?)$ Formed by warming methyl thio urea with chloro-acetic acid and water (Andreasch, *M* 6, 840) Thick prisms or needles (from water), sol hot water and alcohol Boiling $KOHAq$ yields thioglycolic acid Nitrous acid forms a nitrosamine $C_2H_5(NO)N_2SO$ which is an orange red powder, sol hot water

(a) Di methyl-thiohydantoin $C_4H_9N_2SO$ \pm

$NMe C \begin{smallmatrix} S CH_3 \\ NMe CO \end{smallmatrix} [71°]$ Formed by heating di methyl thio urea with chloro acetic acid in aqueous solution (Andreasch, *M* 8, 408) Long colourless prisms, v sol water, alcohol, and ether Smells like nicotine Hot aqueous alkalis convert it into thioglycolic acid Nitrous acid forms an isonitroso compound $C_4H_9N_2SO_2$ [220°]

(B) Di-methyl-thiohydantoin

$NH C \begin{smallmatrix} S CMe_2 \\ NH CO \end{smallmatrix} [114°]$ Formed from di-argentic thiohydantoin and MeI (Andreasch, *M* 8, 416) Thin plates, v sol water, sl sol alcohol Oxidised by $KClO_3$ and HCl to urea and other products

DI-METHYL-THIONINE C_4H_9NS \pm

$N \begin{smallmatrix} C_2H_5 \\ C_2H_5 \end{smallmatrix} \begin{smallmatrix} S \\ S \end{smallmatrix} \begin{smallmatrix} NHMe \\ NMe \end{smallmatrix}$ Obtained by the action of

$FeCl_3$ upon methyl *p* phenylene diamine in the presence of H_2S and HCl The free base is a crystalline powder, sl sol ether and alcohol, insol water The blue alcoholic solution has a strong red fluorescence The hydrochloride is easily sol water with a blue colour and reddish-brown fluorescence The hydriodide (B/HI) is a dark-blue powder, sol hot water and alcohol, sl sol cold water, dyes silk blue The free base by boiling with water is converted into methyl-thionoline with evolution of NH_3Me By treating the product with 70 p.c. H_2SO_4 , a second molecule of NH_3Me is split off and thionol

$N \begin{smallmatrix} C_2H_5 \\ C_2H_5 \end{smallmatrix} \begin{smallmatrix} OH \\ S \end{smallmatrix} \begin{smallmatrix} OH \\ O \end{smallmatrix}$ is formed (Bernthsen & Goske, *B* 20, 931)

METHYL-THIONOLINE $N \begin{smallmatrix} C_2H_5 \\ C_2H_5 \end{smallmatrix} \begin{smallmatrix} S \\ S \end{smallmatrix} \begin{smallmatrix} NHMe \\ O \end{smallmatrix}$

Formed by boiling di-methyl thionine with water, methyl-amine being evolved By treatment with 70 p.c. H_2SO_4 , a second molecule of methyl-amine is split off, and there is formed

thionol $N \begin{smallmatrix} C_2H_5 \\ C_2H_5 \end{smallmatrix} \begin{smallmatrix} OH \\ S \end{smallmatrix} \begin{smallmatrix} OH \\ O \end{smallmatrix}$ (Bernthsen & Goske,

B 20, 932)**Di-methyl-thionoline v METHYLENE VIOLET.****METHYL-THIOPARABANIC ACID**

$C_6H_5N_2SO_2$ \pm $CS \begin{smallmatrix} NMe CO \\ NH CO \end{smallmatrix}$ *Oxalyl methyl-thio urea* [105°] Prepared by passing cyanogen into an alcoholic solution of methyl thio urea and boiling the ppd CSN_2H_2MeCy with $ocno$.

HClAq (Andreasch, *B* 14, 1447, *M* 2, 277) Yellow plates, sol water, alcohol, and ether May be sublimed Converted by warming with aqueous AgNO₃ into methyl parabanic acid

Di-methyl-thioparabanic acid C₈H₈N SO₂ 2 e

CS <NMe CO
NMe CO> *Thiocholestrophane Oxalyl-di methyl-thio-urea* [113°] Prepared by passing cyanogen into an alcoholic solution of *s* di-methyl-thio urea and boiling the ppd cyanide with HClAq (Andreasch) Yellow monoclinic tables, sl sol cold water, v e sol alcohol, and ether Split up by boiling alkalis into di methyl-thio urea and oxalic acid On heating with BaCO₃ and water it gives di methyl oxamide and CSO Boiling aqueous AgNO₃ forms cholestrophane

(a) METHYL-THIOPHENE C₈H₈S 2 e

S <CMe CH
CH CH *Thiotolene* (111°) Occurs in coal tar (V Meyer, *B* 18, 3009) Formed by the action of sodium on a mixture of (a) iodo thio phene and MeBr (V Meyer a Kreis, *B* 17, 1562, Egli, *B* 18, 544) Formed also by the action of P₂S₅ on acetyl propionic (levulic) acid (Kues, *B* 19, 556) Oil Forms a tri bromo-derivative [87°]

(β)-Methyl-thiophene S <CH CMe
CH CH *Thio*

tolene Occurs, together with the preceding isomeride, in crude toluene from which it may be separated by shaking with H₂SO₄, and passing steam through the boiling acid diluted with 20 p c water The mixture of methyl thiophenes so obtained boils at 113 cor and has a S G $\frac{13}{4}$ = 1.0194 (Meyer a Kreis, *B* 17, 787, Schulze, *B* 17, 2853) Formed by distilling sodium pyrotartrate with P₂S₅ (Volhard a Erdmann, *B* 18, 455) Oil Oxidised by alkaline KMnO₄ to (β) thiophenic acid Gives a tri bromo-derivative [34°], and a bromo di nitro derivative [125°]

Di-methyl-thiophene C₈H₈S *Thioxene* Crude thioxene is obtained in considerable quantity by passing steam through sulphuric acid used in purifying xylene diluted with 20 p c of water and heated to boiling (Schulze, *B* 17, 2853)

(aβ') - Di-methyl-thiophene S <CMe CH
CH CMe

m Thioxene (138° cor) S G $\frac{22}{5}$ 9956 V D 4.02 (calc 3.9) Formed by distilling β acetyl-isobutyric acid with P₂S₅ (Zelinsky, *B* 20, 2018) Gives on oxidation methyl-thiophene carboxylic acid and thiophene dicarboxylic acids

Di methyl thiophene SC₂H₅Me₂ (139° cor) S G $\frac{31}{4}$ 9777 Formed from iodo (β) methyl-thiophene, MeI and sodium (Demuth, *B* 19, 1856) Liquid

(aa') - Di-methyl-thiophene S <CMe CH
CMe CH

Thioxene (137° cor) S G $\frac{13}{5}$ 9755 Occurs in coal tar Prepared from crude thioxene by conversion into iodo di methyl thiophene and reducing this body with zinc dust and alcoholic NaOH (Messenger, *B* 18, 565, 1606) Formed by heating acetonil acetone (di methyl ethylene diketone) CH₃COCH₂CH₂COCH₃ (3 pts) with powdered P₂S₅ (2 pts) for an hour at 140°-150°, the yield is 50-60 p c of the theoretical (Paal, *B* 18, 2252) Formed also from (a) iodo (a') methyl-thiophene by leaving it for some weeks in contact with sodium and MeI (Ruffi, *B* 20, 1747)

Colourless mobile liquid of slight odour Dissolves sulphur considerably Gives a red colour with isatin and H₂SO₄, a violet with phenanthraquinone and H₂SO₄, and a reddish-brown with phenyl glyoxylic acid and H₂SO₄ By KMnO₄ it is oxidised to thiotolene carboxylic acid [142°] The di bromo-derivative melts at [47°-50°], and the tri bromo derivative at [144°] It also forms a bromo-derivative [194°], a second di bromo-derivative [46°], a per-bromo-derivative C₈Br₈S [114°], and oily iodo and nitro-derivatives (Messinger) With phenanthraquinone, HOAc, and H₂SO₄ (Laubenheimer's reagent) it gives a reddish-violet colouration

(ββ') Di methyl-thiophene S <CH CMe
CH CMe

(145°) S G $\frac{23}{4}$ 1.0078 Formed by distilling sodium-*s*-di methyl succinate with P S₅ (Zelinsky, *B* 21, 1836) Yellow oil With a trace of isatin in conc H SO₄ it gives an emerald green colour On oxidation it yields an acid crystallising in needles [139°], sl sol cold water

(aβ) - Di-methyl-thiophene S <CMe CMe
CH CH

om Thioxene (137° cor) S G $\frac{21}{5}$ 9938 Formed by distilling β acetyl *n* butyric acid with P S₅ (Paal a Fuschel, *B* 20, 2559, Grunewald, *B* 20, 2585) Colourless, strongly refracting oil In the indophenine reaction it gives a bluish violet colour Laubenheimer's reagent yields a reddish violet colour KMnO₄ oxidises it to methyl thiophene carboxylic acid and thiophene (ββ') di carboxylic acid

Tri-methyl-thiophene C₉H₁₀S 2 e

S <CMe CMe (162°) Formed by distilling S <CH CMe CH CMe CH₃ CO CHMe CHMe CO₂H with P₂S₅ (Zelinsky, *B* 20, 2025)

Tetra-methyl thiophene C₉H₁₂S 2 e

S <CMe CMe (184° uncor) S G $\frac{31}{4}$ 9442 S <CMe CMe Formed from tri methyl thiophene by treatment of this substance (12 g) dissolved in petroleum ether with iodine (48.5 g) and HgO (21 g), distilling with steam, and allowing the iodo tri methyl thiophene which comes over to stand with MeI and sodium (Zelinsky, *B* 21, 1837) Oil Does not give the indophenine reaction

References — DI-BROMO METHYL THIOPHENE AND IODO DI METHYL THIOPHENE

METHYL-THIOPHENE CARBOXYLIC ACID

C₈H₈SO₂ 2 e S <CMe CH
CH CCO₂H or

S <C(CO₂H) CH
CH=CMe *Thiotolene carboxylic acid*

[119°] Formed by oxidising the corresponding di methyl thiophene with alkaline KMnO₄ in the cold (Zelinsky, *B* 20, 2020) Needles, sl sol cold water, v sol ether May be sublimed — CaA', 2/1aq plates — AgA'

(β)-Methyl thiophene (a)-carboxylic acid

S <C(CO₂H) CMe
CH=CMe *c Thiotolene acid* [144°]

Obtained by saponifying its ether, which is formed by the action of ClCO₂Et and sodium amalgam upon iodo (β) methyl thiophene (Levi, *B* 19, 656) Formed also by oxidising (β)-methyl thienyl methyl ketone with alkaline KMnO₄ (Demuth, *B* 19, 680), and by boiling the amide with alcoholic potash Needles (from water), v sol hot water and alcohol Not at-

tacked by KMnO_4 . — CaA'_2 4aq plates — BaA'_2 5aq small plates, sol water — AgA'

Chloride $\text{C}_6\text{H}_4\text{MeS COCl}$ (219°) Liquid, smelling like benzoyl chloride

Amide $\text{C}_6\text{H}_4\text{MeS CONH}_2$ [119°] (Z), [123°] (Levi) Formed by the action of Cl CONH_2 on (B) methyl thiophene in presence of AlCl_3 (Zelinsky, B 20, 2024, Gattermann, A 244, 58) Needles (from water), v sol water

(a) Methyl thiophene (α')-carboxylic acid
 $\text{S} \begin{smallmatrix} \text{C}(\text{CO}_2\text{H})\text{CH} \\ \text{C}(\text{CH}_3)=\text{CH} \end{smallmatrix}$ Methyl thiophene acid

Thiotolemaic acid [142°] (P), [187°] (L) Formed by oxidation of thioxene (from acetylacetone) with alkaline KMnO_4 (Paal, B 18, 2253) Obtained also from its ether, which is produced by heating iodo (a) methyl thiophene with ClCO_2Et and sodium amalgam (Levi, B 19, 656) White needles Somewhat volatile with steam Begins to sublime at c 120° in long needles V e sol alcohol, ether, and boiling water, sl sol cold water Gives no colour reaction with isatin and H_2SO_4 Oxidised by alkaline KMnO_4 to the corresponding thiophene dicarboxylic acid — CaA'_2 3½aq — AgA' white crystalline powder

Methyl-thiophene carboxylic acid

$\text{S} \begin{smallmatrix} \text{CMe CCO}_2\text{H} \\ \text{CH CH} \end{smallmatrix}$ (?) [134 5°] Obtained by oxidising (aB) di methyl thiophene with an alkaline 1 p c solution of KMnO_4 (Grunewald, B 20, 2586)

Di-methyl-thiophene carboxylic acid $\text{C}_6\text{H}_4\text{SO}_2$
 $\text{e S} \begin{smallmatrix} \text{C}(\text{CO}_2\text{H})\text{CCH}_3 \\ \text{C}(\text{CH}_3)\text{CH} \end{smallmatrix}$ [172°] Obtained from its amide by hydrolysis (Zelinsky a Gattermann, A 244, 59) Needles (from dilute alcohol) — AgA'

Imide $\text{C}_6\text{H}_4\text{MeS CONH}_2$ [116°] Formed by acting on di methyl thiophene with Cl CONH_2 in presence of AlCl_3 Colourless needles (from water)

Tri-methyl-thiophene carboxylic acid
 $\text{C}_6\text{H}_3\text{SO}_2$ e $\text{S} \begin{smallmatrix} \text{C}(\text{CO}_2\text{H})\text{CCH}_3 \\ \text{C}(\text{CH}_3)\text{CCH}_3 \end{smallmatrix}$ [208°] Obtained by hydrolysis of its amide (Gattermann a Zelinsky, A 244, 60) Needles (from alcohol)

Amide $\text{C}_6\text{H}_4\text{MeS CONH}_2$ [147°] Formed by the action of Cl CONH_2 upon (aB') tri methyl thiophene in presence of AlCl_3 Crystallises from water

(B) **METHYL-THIOPHENE SULPHONIC ACID** $\text{C}_6\text{H}_4\text{MeS SO}_3\text{H}$ Formed by warming (B) methyl thiophene with fuming H_2SO_4 (Muhlert, B 19, 1620) Syrup, turning red on exposure to air — KA' ½aq — ZnA' 3½aq — PbA' (at 110°)

Chloride $\text{C}_6\text{H}_4\text{MeS SO}_2\text{Cl}$ Oil
Amide $\text{C}_6\text{H}_4\text{MeS SO}_2\text{NH}_2$ [80°] Crystalline nodules (from ether)

METHYL-THIOPHENIC ACID v **METHYL THIOPHENE CARBOXYLIC ACID**

METHYL-THIO-DIPHENYLAMINE v **METHYL IMIDO DI PHENYL SULPHIDE**

METHYL DITHIOPHOSPHATES

Di-methyl di thio-phosphoric acid $\text{C}_6\text{H}_4\text{PO}_3\text{S}_2$ e $\text{Me}_2\text{HPO}_3\text{S}_2$ Formed, together with $\text{Me}_2\text{PO}_3\text{S}_2$, by the action of P_2S_5 (1 pt) on methyl alcohol (5 pts) in the cold (Kowalewski, A 119, 303) Thick liquid, soluble in water Decomposes below 100° — PbA'_2 prisms (from alcohol) Melts below 100°

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Tri-methyl dithiophosphate $\text{Me}_3\text{PO}_3\text{S}_2$ Formed as above Liquid, v sl sol water

METHYL-THIO-PHTHALIMIDINE $\text{C}_6\text{H}_4\text{NS}$
 $\text{e C}_6\text{H}_4 \begin{smallmatrix} \text{C}(\text{NMe}) \\ \text{CH}_2 \end{smallmatrix} \text{S}$ From thiophthalimide and MeI (Way a Gabriel, B 23, 2483) It is converted by conc HClAq at 190° into thio phthalide, and by oxidation into o di cyano di benzyl di sulphide $(\text{C}_6\text{H}_4\text{CyCH}_2)_2\text{S}_2$ [124°] — $\text{B'HCl} - \text{B}'_2\text{H}_4\text{PtCl}_6 - \text{B}'\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OH}$ yellow needles

METHYL-THIO-UREA $\text{C}_6\text{H}_4\text{N}_2\text{S}$ e NH_2CSNHMe [118°] Formed from methyl thiocarbimide and NH_3 (Andreasch, M 2, 277) Prisms, v sol water and alcohol, sl sol ether — B'Hl large plates, v e sol water and alcohol Melts below 100° Decomposed by Ag_2O into AgI and methyl-cyanamide (Berntsen a Klinger, B 11, 492)

Di-methyl-thio-urea $\text{C}_6\text{H}_4\text{N}_2\text{S}_2$ e $\text{CS}(\text{NHMe})_2$ [51 5°] Formed from methyl thiocarbimide and methylamine (Traumann, A 249, 49, Hecht, B 23, 286, cf Andreasch, M 2, 277) Transparent very hygroscopic plates V sol water, alcohol, and acetone, sl sol ether and benzene, v sl sol. light petroleum

METHYL-THYMO-ACRYLIC ACID v *Methyl-derivative of OXY METHYL PROPYL CINNAMIC ACID*

DI-METHYL-TOLANE v **DI-TOLYL ACETYL**

ENE

DI-METHYL-TOLENYL-AMIDINE

Hydrochloride $\text{C}_{10}\text{H}_{11}\text{N}_2\text{HCl}$ e $(\text{NHMe C}(\text{C}_6\text{H}_4\text{Me})\text{NHMe})\text{HCl}$ [200°] Formed by adding the hydrochloride of p tolenyl imido ether ($\text{EtO C}(\text{C}_6\text{H}_4\text{Me})\text{NH})\text{HCl}$ to an alcohol solution of methylamine (Glock, B 21, 2654) Long silky needles (from water), v sol water and alcohol — $\text{B}'_2\text{H}_4\text{PtCl}_6$ 2aq [95°] Dimetric crystals

u Di-methyl-tolenyl-amidine Hydrochloride $(\text{NMe C}(\text{C}_6\text{H}_4\text{Me})\text{NH})\text{HCl}$ Formed from the hydrochloride of p tolenyl imido ether and dimethylamine (G) Short prisms

METHYL-TOLINDOLE v **DI-METHYL INDOLE**

METHYL-TOLISATIN v **DI-METHYL ISATIN**

DI-METHYL-TOLUBUTYLAMINE v

METHYL BUTYL PHENYL DI-METHYL AMINE

METHYL o-TOLUIDINE $\text{C}_6\text{H}_4\text{N}$ e $[2] \text{C}_6\text{H}_4\text{MeNHMe}$ o *Tolyl methyl amine* (207°) SG 15 973 Prepared by the action of tin and HClAq on the nitro-amine which is obtained from the crude product of the action of methyl alcohol and HCl on o-toluidine (Monnet, Reverdin, a Nolting, B 11, 2278) Obtained also by heating o-toluidine hydrobromide (or hydriodide) with 5 p c more than an equivalent quantity of methyl alcohol for 8 hours at 150°, the yield being 46 p c of the theoretical (Reinhardt a Staedel, B 16, 29) It is also produced by distilling o tolyl amido acetic acid (Widman, J pr [2] 38, 303) Colourless oil — $\text{B}'_2\text{H}_4\text{PtCl}_6$

Acetyl derivative $\text{C}_6\text{H}_4\text{MeNAcMe}$ [56°] (251°) (R a S), (o 260°) (M, R, a N)

Nitrosamine $\text{C}_6\text{H}_4\text{MeN(NO)Me}$ Oil Converted by alcoholic HCl into the isomeric p-nitroso o-methyl-toluidine

[5 1 2] $\text{NO C}_6\text{H}_4\text{MeNHMe}$ or $\text{C}_6\text{H}_4\text{Me} \begin{smallmatrix} \text{NHMe} \\ \text{N} \end{smallmatrix} \text{O}$, which crystallises in green plates, [151°], sol. benzene On boiling with dilute aqueous NaOH

D D

it is converted into nitroso cresol and methylamine Potassium permanganate oxidises it to nitro methyl-toluidine [184°] Its hydrochloride $C_8H_9N.OH_2Cl$, aq crystallises in yellow cubes [110°], v sl sol conc HCl aq (Kock, *A* 243, 808)

Methyl-*m*-toluidine $[31]C_8H_9Me.NHMe$ (207°) Formed by the action of MeI on *m*-toluidine, and purified by means of the acetyl derivative (Nörling, *B* 11, 2279) Oil — $B'_2H_2PtCl_6$

Acetyl derivative $C_8H_9Me.NAcMe$ [66°] (c 250°) Sol hot water, alcohol, and ether

Methyl-*p*-toluidine $[41]C_8H_9Me.NHMe$ (208°) Formed by passing $MeCl$ into boiling *p*-toluidine The ethereal extract of the crude product is freed from *p*-toluidine by ppn with H_2SO_4 , evaporated, and mixed with Ac_2O The resulting mixture of di methyl toluidine and acetyl-methyl toluidine may then be fractionally distilled, and the acetyl derivative saponified by HCl aq or $NaOH$ aq (Thomsen, *B* 10, 1582) Oil — $B'_2H_2PtCl_6$

Acetyl derivative $C_8H_9Me.NAcMe$ [83°] (283°) Plates (from ether alcohol), sl sol. water, v sol alcohol and ether

Propionyl derivative C_8H_9NO i.e. $C_8H_9Me.N(C_2H_5O)Me$ (266°–269°) Formed from methyl *p*-toluidine and propionic anhydride (Norton a Livermore, *B* 20, 2270) Converted by dilute nitric acid into tri nitro methyl *p*-toluidine [130°]

Nitrosamine $C_8H_9Me.NMe(NO)$ [54°] Insol water, v sol alcohol and ether

Di-methyl *o*-toluidine $C_8H_9N_2O$ i.e. $[21]C_8H_9Me.NMe_2$ (183°) Formed by distilling its methyl hydroxide (Thomsen, *B* 10, 1586, Nörling, *B* 11, 2279) Prepared by heating *o*-toluidine hydrobromide (or hydriodide) with a little (5 p.c.) more than two equivalents of methyl alcohol for 8 hours at 150°, the yield being 93 p.c. of the theoretical (Reinhardt a Staedel, *B* 16, 29) Oil Converted by HNO_3 (S G 16) into di nitro *o*-tolyl methyl nitramine (Van Romburgh, *R T C* 3, 395) — $B'_2H_2PtCl_6$ — $B'_2H_2FeCy_4$ white needles — $B'_2H_2FeCy_4$ 4 $\frac{1}{2}$ aq yellow unstable crystals (Wurster a Roser, *B* 12, 1826)

Methylo iodide $C_8H_9Me.NMe_2I$ Formed from *o*-toluidine and MeI (Thomsen) It is also a product obtained by heating di methyl aniline methylo-iodide at 220°–230° (Hofmann, *B* 10, 1585) Needles

Di-methyl *m*-toluidine $[31]C_8H_9Me.NMe_2$ (208°) (*N*, *R* a *S*), (215°) (Wurster a Riedel, *B* 12, 1797) Formed from *m*-toluidine and MeI , or by distilling its methylo iodide (Nörling) When dissolved in H_2SO_4 and treated with HNO_3 (S G 15), it yields $C_8H_9(NO_2)_2Me.NMe_2NO_2$ (Van Romburgh, *R T C* 3, 418) — $B'_2H_2PtCl_6$ — $B'_2H_2FeCy_4$ 2 aq white needles — $B'_2H_2FeCy_4$ 1 $\frac{1}{2}$ aq (Wurster a Roser, *B* 12, 1826)

Methylo iodide $B'MeI$ Yields ($B'MeCl$), $PtCl_6$, yellow cubes

Di-methyl-*p*-toluidine $[41]C_8H_9Me.NMe_2$ (208°) (*R* a *S*, *H*), (210°) (Van Romburgh) S G 988 Formed, together with other products, by heating dimethylaniline methylo-iodide at 220°–230° (Hofmann, *B* 5, 707) Formed also by heating its methylo iodide with water and PbO and distilling the resulting $C_8H_9Me.NMe_2OH$ (Hubner,

Tolle, a Athenstädt, *A* 224, 387, cf Thomsen, *B* 10, 1586) Prepared by heating *p*-toluidine hydrobromide (or hydro iodide) with rather more (5 p.c.) than two equivalents of methyl alcohol for eight hours at 150°, the yield being 95 p.c. of the theoretical (Reinhardt a Staedel, *B* 16, 29) Oil Converted by fuming HNO_3 and H_2SO_4 into $C_8H_9(NO_2)_2Me.NMe_2NO_2$ (Van Romburgh, *R T C* 3, 404) — $B'_2H_2PtCl_6$ — $B'_2H_2FeCy_4$ aq white powder — $B'_2H_2FeCy_4$ 2 $\frac{1}{2}$ aq yellow crystals (Wurster a Roser, *B* 12, 1826)

Methylo-iodide $C_8H_9Me.NMe_2I$ Formed from *p*-toluidine and MeI Converted by moist Ag_2O into $C_8H_9Me.NMe_2OH$ Yields ($C_8H_9Me.NMe_2Cl$), $PtCl_6$

References — Bromo-*DI*-METHYL-TOLUIDINE and Nitro METHYL-TOLUIDINE

DI-METHYL-*o*-TOLUIDINE SULPHONIC ACID $C_8H_9Me.(NMe_2)SO_3H$ Formed by sulphonating di methyl *o*-toluidine (Michler a Sampaio, *B* 14, 2167) Large prisms, v sol hot water, insol alcohol — BaA' , soluble plates — CaA' , small nodules — ZnA_2 easily soluble needles

METHYL-TOLUQUINOLINE v *DI* METHYL QUINOLINE

METHYL-TOLUQUINOXALINE v. *DI* METHYL QUINOXALINE

METHYL-TOLYLENE-DIAMINE v *TOLYL* ENF *DI* *THYL* DIAMINE

METHYL TOLYL KETONE v *TOLYL* METHYL KETONE

METHYL-TOLYL NITROSAMINE v *Nitrosamine* of METHYL TOLUIDINE

DI METHYL-TOLYL PHOSPHINE v *TOLYL* *DI* METHYL PHOSPHINE

METHYL TRIDECYL KETONE $C_{13}H_{27}O$ i.e. $CH_3.CO.C_{12}H_{25}$, [39°] (294°) S G (liquid) 22 8182 Formed by distilling a mixture of barium myristate and barium acetate (Krafft, *B* 12, 1669, 15, 1724) Yields acetic and tridecic acids on oxidation

DI - METHYL - TRIDECYL - PYRIDINE $C_{20}H_{33}N$ i.e. $C_5NH_4Me_2(C_{12}H_{25})$ (216° at 13 mm) Formed by distilling its dicarboxylic acid with soda lime (Krafft a Mai, *B* 22, 1758) Oil — $B'_2H_2PtCl_6$ yellow plates

DI METHYL-TRIDECYL-PYRIDINE DI-CARBOXYLIC ACID $C_8NMe_2(C_{12}H_{25})(CO_2H)_2$ Formed by saponifying its ether with alcoholic potash — $H_2A'HCl$ crystalline powder, v e sol alcohol

Ethyl ether EtA' (265° at 10 mm) Formed from its dihydride by treatment of the alcoholic solution with nitrous acid (Krafft a Mai, *B* 22, 1758) Yellowish oil — $Et_2A'HCl$ needles

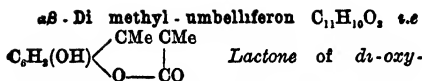
Di methyl-tridecyl pyridine dihydride di carboxylic ether $C_8NMe_2(C_{12}H_{25})_2H_2(CO_2Et)_2$ [60°] Formed by heating a mixture of myristic aldehyde, alcoholic NH_3 , and acetoacetic ether (Krafft a Mai, *B* 22, 1757) Hard crystalline crusts

METHYL-TROPIDINE v *TROPIDINE*

METHYL-TROPINE v *TROPINE*

METHYL-UMBELLIC ACID v *DI-OXY-PHENYL-CROTONIC ACID*

METHYLUMBELLIFERON v *Lactone* of *DI-OXY-PHENYL CROTONIC ACID*



phenyl pentenone acid [256°] Formed by the action of H_2SO_4 on a mixture of methyl acetoacetic ether and resorcin (Pechmann a Duisberg, *B* 16, 2127) Colourless needles of high refractive power Its dilute alkaline and H_2SO_4 solutions have a blue fluorescence

Isomeride v Lactone of Di oxy tolyl crotonic acid.

METHYL UNDECYL KETONE v METHYL HENDFCYL KETONF

METHYL URACIL v Di oxy methyl pyrimidine

METHYL - URAMIDO - ACETIC ACID v METHYL HYDANTOIC ACID

m - METHYL - URAMIDO BENZOIC ACID $C_9H_7N_3O_5$ *s.e.* $HMeN$ CO NH C_6H_5 CO_2H Formed by the action of methylamine upon cyanic acid oxamido benzoic acid NC CO NH C_6H_5 CO_2H (Gress, *B* 18, 2415) White needles, sol alcohol, v sl sol water — $A'Ag$ white plates

METHYL URAMINE An old name for METHYL GUANIDINE

METHYL UREA $C_2H_5N_2O$ *s.e.* NH . CO $NHMe$ [102°] Formed from methyl cyanate and ammonia, or by evaporating a mixture of potassium cyanate and methylamine sulphate (Wurtz, *C* 32, 414) It is also a product of the action of HCl and $KClO_4$ on caffeine (Fischer, *A* 215, 257) It may be prepared by boiling its acetyl derivative with conc HNO_3 (Hofmann, *B* 14, 2734)

Properties — Deliquescent prisms, v e sol water and alcohol Its aqueous solution is neutral to litmus, and yields with nitric acid a pp of the nitrate $C_2H_5N_2OHNO_3$ [128°], which is converted by fuming nitric acid into methylamine and ammonium nitrate (Franchimont, *R T C* 3, 220)

Nitrosamine NH_2CO $N(NO)Me$ [124°] Formed by adding $NaNO_2$ to a solution of methyl urea nitrate containing ice (Von Bruning, *B* 21, 1810, *A* 253, 6) Yellow plates or tables (from ether), v sol hot water, alcohol, and ether Decomposed by long boiling with water Exhibits Liebermann's reaction On reducing with zinc dust and $HOAc$ it yields methyl - semi - carbazide NH_2CO $NHMeNH_2$ whence, by heating with conc $HClAq$ for 6 hours at 100°, methyl hydrazine is got

Acetyl derivative $NHAc$ CO $NHMe$ [180°] Formed from methyl urea and Ac_2O Formed by the action of boiling dilute $NaOH$ on a mixture of bromoacetamide (CH_2CO $NHBr$) and acetamide This mixture is prepared by adding aqueous (10 p.c) $NaOH$ to acetamide (10 pts) mixed with bromine (13.5 pts) until the colour changes from red to yellow (Hofmann, *B* 14, 2725, 15, 408) In this reaction methyl cyanate is perhaps first formed and then unites with acetamide Monoclinic prisms (Haushofer, *J* 1882, 365), sol alcohol, ether, and hot water Split up by alkalis into NH_3 , CO_2 , methylamine, and acetic acid Boiling conc $HClAq$ forms acetic acid and methyl urea

s Di-methyl-urea $C_4H_{10}N_2O$ *s.e.* $CO(NHMe)_2$ [102.5°] (269°) Formed by the action of methylamine on methyl cyanate (Wurtz, *Rép*

chim pure, 1862, 199) According to Wurtz, the same body when prepared by the action of cold water on methyl cyanate melts at 99.5° and boils between 273° and 288° *s* Di methyl-urea crystallises easily, is v sol water and alcohol, and forms a hygroscopic nitrate $C_4H_{10}N_2OHNO_3$ [c 65°] which is decomposed by fuming HNO_3 , forming methylamine (Franchimont, *R T C* 3, 222)

Cyano-acetyl derivative $C_5H_8N_2O_2$ *s.e.* $NHMe$ CO NMe CO CH_2Cy Formed from *s* di-methyl urea and cyano acetyl chloride (Mulder, *B* 12, 466) Crystals, not melted below 260° Gives with bromine water the compound $CO \begin{cases} NMe CO \\ NMe CO \end{cases} CBr_2$ Conc HNO_3 forms, on warming, two purple-red compounds

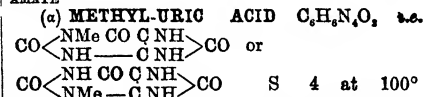
u Di methyl urea NMe_2CO NH_2 [180°]. Formed by evaporating the mixed solutions of potassium cyanate and dimethylamine sulphate (Franchimont, *R T C* 2, 122, 3, 222) Large hard crystals with sweet taste, sl sol alcohol and ether Boiling Ac_2O forms NMe_2Ac and cyanuric acid Aldehyde in the cold slowly forms $CH_3CH(NHCO$ $NMe_2)_2$ [160°] Chloral forms $CCl_3CH(OH)NHCO$ NMe_2 [156°] and $C_2H_5Cl_2N_2O_2$ [74°] (Van der Zande, *R T C* 3, 222) Nitrate $B'HNO_3$ [101°] Very large crystals Converted by fuming HNO_3 into dimethyl nitramine — Oxalate $B'H_2C_2O_4$ *aq* Decomposes at 105° — Picrate $B'C_6H_3(NO_2)_3OH$ [130°] Decomposes on fusion (Van der Zande)

Tri-methyl-urea $C_4H_{11}N_3O$ *s.e.* NMe_2CO $NHMe$ [76°] (232° cor) Formed by mixing ethereal solutions of methyl cyanate and dimethylamine (Franchimont, *R T C* 3, 226) Crystallises well from ether, v sol water and alcohol, sol benzene Fuming HNO_3 forms di methyl nitro amine (NMe_2NO_2) and methylamine

Tetra methyl urea $C_4H_{12}N_4O$ *s.e.* $CO(NMe_2)_2$ (177° v) $SG \pm 972$ Prepared by passing dimethylamine into a solution of di methyl chloro formamide $ClCONMe_2$ (derived from $COCl_2$ and NMe_3) in benzene (Michler a Escherich, *B* 12, 1162, Franchimont, *R T C* 3, 228) Oil, v sol alcohol and ether With conc HNO_3 (*S* *G* 1.5) it yields NMe_2NO_2 and dimethylamine

Reference — CHLORO METHYL UREA

METHYL URETHANE v METHYL CARBAMATE



Formed by heating at 150° acid lead urate with MeI diluted with ether (Hill, *B* 9, 370, 1090, *Am S* [3] 12, 428) Thin prisms, sl sol boiling water, insol alcohol and ether Sol conc H_2SO_4 and reppd by water Oxidised by alkali $KMnO_4$ to methyl allantoin $C_4H_8N_4O_3$ [225°] $KClO_4$ and HCl oxidise it to urea and methyl-alloxan By heating with $HClAq$ it is split up into glycoecoll, methylamine, CO_2 , and NH_3 (Fischer, *B* 17, 1776)

Salts — K_2A'' 3aq Ppd by adding alcohol to aqueous solution — KHA'' 3aq — NaA'' 3aq — $NaHA''$ 3aq — CaA'' 3aq — BaA'' 4aq — BaA'' 3aq; tufts of delicate needles

Methyl uric acid $C_5H_7MeN_3O_6$. Possibly identical with the preceding. Prepared by fusing urea (3 mols) with methyl amido acetic acid (1 mol), extracting the fusion with water, ppg by ammoniacal $AgNO_3$ and magnesium mixture, digesting the pp with an alkaline sulphide, filtering, acidifying, and evaporating to a small bulk (Horbaczewski, *M* 6, 859). Formed also by heating methyl hydantoin with amyl allophanate or with biuret (Horbaczewski, *M* 8, 584). Micaceous scales, sol boiling water and alkalis, sl sol cold water, dilute acids, alcohol, and ether. Gives the murexide reaction. On heating with $HClAq$ at 170° it gives a product (probably methylamine) which gives the carbamide reaction.

(β)-Methyl-uric acid $C_5H_7MeN_3O_6$ $\pm e$
 $CO \begin{matrix} \text{NH} & CO & C & NMe \\ \diagdown & & \diagup & \\ & C & & NH \end{matrix} CO$ Tri oxy methyl
 purin S 05 at 100°

Formation — 1 By heating the di methyl derivative of chloro di oxy methyl purin with $HClAq$ at 130° (Fischer, *B* 17, 332) — 2 By heating di chloro oxy methyl purin with $HClAq$ at 140° (Fischer, *B* 17, 1777) — 3 Occurs in small quantity, together with a much larger quantity of (α)-di methyl uric acid, by heating neutral lead urate with MeI at 100° (F).

Properties — Slender crystals. Sol alkalis. Reduces ammoniacal $AgNO_3$ in the cold. Gives the murexide reaction. On heating with PCl_5 it yields di chloro-oxy methyl purin. On oxidation with HNO_3 or with $KClO_3$ and HCl it gives methyl-urea and alloxan. Split up by heating with $HClAq$ into CO_2 , methylamine, NH_3 , and glycecoll.

(α)-Di-methyl-uric acid $C_5H_7Me_2N_3O_6$ $\pm e$
 $C_5H_7Me_2N_3O_6$, $C_5H_7Me_2N_3O_6$ [above 340°] S 5 at 100° , 06 at 20° . Prepared by heating neutral lead urate with MeI in ether for 20 hours at 165° (Mabery a Hill, *Am* 2, 306, *P Am* 4, 15, 256, *B* 11, 1329, Fischer, *B* 17, 1779). Slender monoclinic prisms (containing aq), sl sol water, sol conc H_2SO_4 and $HClAq$, insol alcohol and ether. When heated with conc $HClAq$ at 170° it is resolved into CO_2 , ammonia, NH_3 , and glycecoll. Nitric acid oxidises it, forming methyl alloxan and methyl urea. HCl and $KClO_3$ form methyl urea, methyl alloxan, and a small quantity of a compound $C_5H_7N_3O_6$, [160°]. The ammonium salt is decomposed on boiling its solution with separation of the free acid (difference from (α)-methyl uric acid).

Salts — K_2A' 4aq silky needles, v sol water — KHA' 11aq needles, sol water — Na_2A' 44aq needles, sol water — $NaHA'$ 2aq small needles, sol water — BaA' 3aq flat prisms, m sol hot water — BaH_2A' 23aq

(β)-Di-methyl-uric acid
 $CO \begin{matrix} \text{NH} & CO & C & NMe \\ \diagdown & & \diagup & \\ & C & & NH \end{matrix} CO$ Tri oxy-methyl-
 purin Formed by the action of H_2SO_4 at 140° on the di ethoxy-compound obtained by treating di-chloro-oxy di methyl purin with alcoholic $NaOH$. Formed also by heating di chloro oxy-di-methyl purin with fuming HCl at 130° (Fischer, *B* 17, 337, 1779). Colourless crystal line powder, v sl sol water, alcohol, and ether. Melts at a very high temperature.

Reactions — On heating with HCl it is split up into saccosine, methylamine, CO_2 , and NH_3 , PCl_5 converts it back into di chloro oxy-di-methyl purin. By $K_2Cr_2O_7$ and H_2SO_4 it is oxidised to cholestrophane. By oxidation with HNO_3 or $KClO_3$ and HCl it chiefly gives a body $C_5H_7N_3O_6$, which forms large colourless crystals [174°], v sol water, and decomposed on boiling with baryta water into mesoxalic acid, urea, and probably di methyl urea.

Tri-methyl-uric acid $C_5H_9N_3O_6$ $\pm e$.
 $MeN-CO \begin{matrix} \text{OC} & C & NMe \\ \diagdown & & \diagup \\ & C & \\ & \text{NH} & \end{matrix} CO$ or $HN-CO \begin{matrix} \text{OC} & C & NMe \\ \diagdown & & \diagup \\ & C & \\ & MeN & \end{matrix} CO$

[345°] Formed by heating the lead salt of (β)-di methyl uric acid with methyl iodide (Fischer, *B* 17, 1782). Fine white needles. Sublimable. Sol hot water, sl sol alcohol and chloroform. Dissolves in alkalis. Gives the murexide reaction strongly. Is very unstable towards acids. $C_5H_9N_3O_6$ fine white needles.

Tetra-methyl-uric acid $C_5H_9N_3O_6$ $\pm e$
 $CO \begin{matrix} NMe & CO & C & NMe \\ \diagdown & & \diagup \\ & C & \\ & NMe & \end{matrix} CO$ [218°] Formed by heating the silver salt of tri methyl uric acid with methyl iodide (Fischer, *B* 17, 1784). Distils undecomposed. Fine white needles. V sol hot water, less sol cold, sl sol ether. Has no acid properties. It is readily decomposed by alkalis, with evolution of methyl amine. It gives the murexide reaction.

METHYL-URIC ACID $C_5H_7O_6$ $\pm e$.
 $CO_2H \begin{matrix} CH & CHMe \\ \diagdown & & \diagup \\ & C & \\ & Me & \end{matrix} CO$ or $CO_2H \begin{matrix} CH & CH \\ \diagdown & & \diagup \\ & C & \\ & Me & \end{matrix} CO$

[98°] Formed by heating methyl methionine acid to 250° as long as CO_2 is evolved (Fittig, *A* 250, 205). Needles (from water) or prisms (from other solvents), m sol hot, v sl sol cold, water, v sol cold, v e sol hot alcohol, v sol petroleum ether, $CHCl_3$, benzene, ether, and $HOAc$, volatile with steam. Yields on distillation CO_2 and oxy di methyl pentamethenyl hydride.

Salts — $(C_5H_7O_6)_2Ba$ 4aq trimetric plates, $a b c = 0.9937 \times 1.49021 = A'$, Ca 4aq pearly plates, insol alcohol — AAg small prisms (from water).

Ethyl ether $A'Et$ (219°) Colourless oil, heavier than water, volatile with steam.

METHYL-VALERIC ACID v $Hfpxic$ acid
METHYL-VALEBOLACTONE v *Lactone* of Ox_1 hexoic acid

METHYL-VANILLIN v *Di methyl derivative* of **PROTocatechuic aldehyde**

DI-METHYL-DI-VANILLIN v *Tetra methyl derivative* of **TETRA OXY DI PHENYL DICARBOXYLIC ALDEHYDE**

DI-METHYL-VINYL-AMINE *Methylhydroxide* C_5H_9NO $\pm e$ $NMe_2(C_2H_5)OH$ *Neurine*. Formed by the action of moist oxide of silver on di methyl bromo vinyl methyl bromide $NMe_2(CH_2CH_2Br)Br$, the product of the union of trimethylamine with ethylene bromide (Hofmann, *C R* 47, 558, *Chem Gaz* 1858, 434, Baeyer, *A* 140, 311). It is also a product of the decomposition of protagon by baryta (Liebreich, *B* 2, 12), and of the putrefaction of flesh (Brieger, *B* 16, 1190, 1406, 17, 516, 1137). It is extremely soluble in water, and is extracted from the solution by ether with great difficulty (Marino, *G* 13, 441). Its solution is strongly alkaline, and fumes with HCl . On evaporating

its solution it decomposes with evolution of NMe_3 . Its aqueous solution is poisonous, and antagonistic to atropine as regards the heart and glandular system (Cervello, *Arch Ital Biol* 7, 232)

Methylo chloride $\text{*NMe}_3(\text{C}_2\text{H}_5)_2\text{Cl}$ Very deliquescent needles — $(\text{NMe}_3(\text{C}_2\text{H}_5)_2\text{Cl})_2\text{PtCl}_2$, yellow crystals, readily changing to neurine platinumchloride $(\text{NMe}_3(\text{C}_2\text{H}_5\text{OH})\text{Cl})_2\text{PtCl}_2$ — $\text{NMe}_3(\text{C}_2\text{H}_5)_2\text{AuCl}$, yellow needles

METHYL VINYL KETONE CARBOXYLIC ACID $\text{CH}_3\text{COCHCHCO}_2\text{H}$ β -Acetyl acrylic acid [125°] Formed by the action of aqueous sodium carbonate on β bromo acetyl propionic acid $\text{CH}_3\text{COCHBrCH}_2\text{CO}_2\text{H}$ in the cold (Wolf, *B* 20, 425) Plates, v sol alcohol and ether, m sol cold water — CaA_2 nodules, v sol water — ZnA^+ , amorphous — AgA^+ , stellate groups of needles

Phenyl hydrazide
 $\text{PhNHNCMeCH}_2\text{CH}_2\text{CO}_2\text{H}$ [157°] Formed from the acid and phenyl hydrazine (Decker, *B* 21, 2937) or by saponifying its ether Lemon-yellow needles, m sol hot water, v sol alcohol and ether

Ethyl ether *EtA **Phenyl hydrazide**
 $\text{PhNHNCMeCHCHCO}_2\text{Et}$ [117°] Formed from bromo acetyl propionic ether and phenyl hydrazine (Bender, *B* 21, 2493) Yellow spangles, sl sol cold, v sol hot alcohol On reduction with tin and HClAq it yields methyl indyl acetic acid

Isomeride v TETRIC ACID

METHYL-VIOLET A violet dye obtained by the oxidation of di methyl aniline It consists of a mixture of hexa and penta methyl tri amido tri phenyl carbinols (q) (Fischer, *B* 16, 706, 2904)

DI-METHYL-XANTHINE v THEOBROMINE and THEOPHYLLINE

Tri-methyl-xanthine v CAFFEINE

METHYL-XYLENE v MESITYLENE and ψ CUMENE

METHYL-XYLIDINE $\text{C}_6\text{H}_5\text{N}^+\text{H}^-$
 $\text{C}_6\text{H}_5\text{Me}_2\text{NMe}$ One of the products obtained by heating di methyl aniline hydro iodide at 225° (Hofmann, *B* 5, 712) Oil

Di methyl-xyldine $\text{C}_6\text{H}_5\text{N}^+\text{H}^-$
 $\text{C}_6\text{H}_5\text{Me}_2\text{NMe}_2$ (196°) SG 9293 Obtained by heating methyl xyldine with MeI (Hofmann, *B* 5, 712) Oil

Di-methyl-xyldine $\text{C}_6\text{H}_5\text{Me}_2\text{NMe}$ (203°) Obtained by methylation of crude xyldine (H) — B^+MeI

Di methyl-xyldine $\text{C}_6\text{H}_5\text{Me}_2\text{NMe}$ [87°] A by product in the preparation of methyl aniline (Sesemann, *B* 6, 445) Long needles (from ligron) — B^+EtBr

Di-methyl-xyldine [134] $\text{C}_6\text{H}_5\text{Me}_2\text{NMe}$ (204°) Formed by heating xyldine hydrobromide with MeOH (Baur a Staedel, *B* 16, 32) Oil — $\text{B}^+\text{H}_2\text{PtCl}_2$, small yellow crystals

METHYL XYLAL KETONE v XYLAL METHYL KETONE

METHYSTICIN $\text{C}_{11}\text{H}_{14}\text{O}_3$ (P), $\text{C}_{11}\text{H}_{16}\text{O}_3$ (D) [189°] (D), [137°] (P) Extracted by means of alcohol from Kawa root, the root of *Piper methysticum* (Gobley, *J Ph* [3] 37, 19, O'Rorke, *C R* 50, 498, Nölting a Kopp, *Monat Scient* [8] 4, 9, 20, Czuent, *An* 1, 150, Davidoff, *J R*

19, 522, Pomeranz, *M* 9, 863, 10, 785) White tasteless prismatic needles, sl sol hot water, ether, and ligroin, v sol hot alcohol, benzene, and chloroform Dissolves in NaOHAq , but on heating the solution methystic acid is formed On fusion with KOH protocatechuic acid is formed Boiling with KOH forms piperonal Alkaline KMnO_4 oxidises it to piperonylic acid Acetyl derivative $\text{C}_{11}\text{H}_{14}\text{Ac}_2\text{O}_3$ [123°] Benzoyl derivative $\text{C}_{11}\text{H}_{14}\text{Bz}_2\text{O}_3$ [148°] Methystic acid $\text{C}_{11}\text{H}_{12}\text{O}_3$ [180°] When 10g of methysticin are warmed with 6 pc NaOHAq 5g of methystic acid are formed. Methystic acid when heated at 180° evolves CO_2 , and the product, after cooling, melts at 93° KMnO_4 oxidises methystic acid to piperonal [37°] and piperonylic acid $\text{C}_8\text{H}_8\text{O}_3$ [227°] Dilute mineral acids convert it into methysticol. Ac_2O has no action

Methysticol $\text{C}_{11}\text{H}_{12}\text{O}_3$ $\text{CH}_2\text{<O>C}_6\text{H}_4(\text{C}_2\text{H}_5\text{O})$ $\left[\begin{smallmatrix} 1 \\ 2 \end{smallmatrix} \begin{smallmatrix} 4 \end{smallmatrix} \right]$ [94°] Formed by warming methysticin or methystic acid with dilute HClAq or H_2SO_4 (Pomeranz, *M* 10, 792) Prisms, insol alkalis, v sol alcohol and ether Forms a phenyl hydrazide melting at 143° According to Davidoff methysticin is $\text{C}_{10}\text{H}_{12}\text{O}_3$ and is decomposed by alcoholic NH_3 into 'methysticin hydrate' $\text{C}_8\text{H}_{10}\text{O}_3$ and a compound $\text{C}_8\text{H}_{11}\text{NO}_2$ which crystallises from water in yellow needles The 'methysticin hydrate' is also formed by the action of alcoholic KOH on methysticin It melts at 159°, forms the crystalline salts $\text{K}_2\text{C}_8\text{H}_9\text{O}_3$ and $\text{Ba}(\text{C}_8\text{H}_9\text{O}_3)_2$, the ether $\text{EtC}_8\text{H}_9\text{O}_3$ [100°] and the benzoyl derivative $\text{C}_8\text{H}_9\text{Bz}_2\text{O}_3$ [122°]

METINULIN v INULIN

MICROCOSMIC SALT Sodium ammonium-hydrogen phosphate, $\text{Na}(\text{NH}_4)\text{HPO}_4 \cdot 4\text{H}_2\text{O}$, v. PHOSPHORIC ACIDS AND PHOSPHATES

MILK Milk is the secretion of the mammary glands, to the naked eye it appears to be a white fluid, which in thin layers has a bluish tint Its reaction is alkaline in herbivorous animals and in woman, acid in carnivorous animals Its specific gravity varies greatly, averaging between 1.028 and 1.035 On microscopic examination milk is found to be an emulsion, consisting of a clear fluid which has a light straw colour, in which are suspended numerous minute fat globules of varying size, each enclosed in an envelope of casein Numerous particles of casein and nuclei are also seen (Kehrer, *Arch Gynaek* 2, 1) The clear fluid in which these float contains proteids, lactose, and mineral salts in solution During the first few days after parturition the milk contains a larger proportion of solid constituents, and is found microscopically to contain, besides the ordinary fat globules, a number of the secreting epithelium cells of the mammary gland of which the protoplasm has undergone fatty degeneration, this secretion has received the name *colostrum* The name *uterine milk* is given to the whitish secretion of the uterine glands which is poured out from them during the early months of pregnancy in certain animals

The following tables adapted from Charles's *Physiological Chemistry*, p 383, after Gorup Besanez, Liebermann, Gautier, &c, give quantitative statements of the constituents which occur in the milk of the commoner animals. —

Constituents	Woman Milk	(Colo- strum)	Ass	Ow	Goat	Sheep	Mare
Water	87.65	84.08	90.70	88.56	86.78	83.31	82.84
Solids	12.35	15.92	9.30	13.44	13.24	16.69	17.16
Proteids	8.07	8.23	1.70	4.08	4.23	5.73	1.64
Fats	8.81	5.78	1.55	4.03	4.48	6.05	6.87
Milk sugar	5.01	6.51	5.80	4.60	3.91	3.96	8.65
Mineral salts	0.17	0.35	0.50	0.73	0.62	0.68	

The Ash of Milk in 100 parts

	Woman's Milk (Wildenstein)	Cow's Milk (Weber)	(Haidlen)
Sodium . .	4.21	6.88	8.27
Potassium . .	31.59	24.71	15.42
Chlorine . .	19.06	14.39	16.96
Calcium . .	18.78	17.31	
Magnesium . .	0.87	1.90	56.52
Phosphoric acid	19.00	29.13	
Sulphuric acid	2.64	1.15	
Ferric oxide	0.10	0.33	0.62
Silica	trace	0.09	

The most abundant salts are thus the phosphates. The excess of potassium over sodium salts is the opposite to what obtains in blood plasma and most other fluids of the body. (See also Bunge, *Dissert* Dorpat, 1874, *Zeit Physiol Chem* 13, 399). The iron in milk appears to be combined with nuclein (Bunge).

Condensed Milk (Cane Sugar added).

Water	29 to 24
Solids	71 to 76
Casein	16 to 18
Milk sugar	8 to 18
Cane sugar	27 to 29
Ash	2 to 2.5
P ₂ O ₅	0.2 to 0.7

Swiss Condensed Milk (benzoic acid added—Flieschmann)

Water . .	52.31	Milk sugar	17.43
Fat . .	18.09	Ash	2.78
Proteid . .	12.13	Benzoic acid	1.74

Gases of milk at 0° 41 metre pressure (Pflüger)

Carbonic acid	7.6 in 100 vols of milk
Oxygen	0.1
Nitrogen	0.7

Certain preparations (Koumiss, Kephir) are now made from mare's and cow's milk, in which the alcoholic fermentation is allowed to take place, and which are valuable stimulants in cases of disease, especially of disordered digestion (for references on this subject v Maly's *Jahrsber* 14, 187).

The proteids of milk—The most abundant is casein. This is coagulated by rennet, and the more soluble precursor in the milk of the curd should be more accurately called *caseinogen* (v PROTEIDS). Albumin (*lact-albumin*) which is very similar to serum albumin is also present. Other proteids have also been described in milk, but their presence is doubtful, thus peptone is described by Schmidt Mulheim (*Pf* 28, 287), by Dogiel (*Zeit Physiol Chem* 9, 591), and by Biel (*Maly's Jahrsber* 15, 198), hemi albumose by

J Schmidt (*Maly's Jahrsber* 14, 175). These proteids appear to be primary albumoses formed by the acidification and boiling employed to separate the casein and albumin (Neumeister, *Zeit Biol* 24, 271). True peptone is always absent (Neumeister, *loc cit*, Sebelien, *Zeit Physiol Chem* 13, 135, Halliburton, *Journal of Physiol* 11, 449). Whey proteid (Hammarsten), the lacto-protein of some observers, is a by product of the action of rennet on caseinogen. Two other proteids, lacto globulin, which is identical with paraglobulin (J Sebelien, *Maly's Jahrsber* 15, 184), and lacto syntonn, which resembles acid albumin in its properties (J Biel, *loc cit*), are also stated to occur in small quantities, but the evidence that they exist is not conclusive (Halliburton, *loc cit*). In koumiss the proteids are stated to consist of albumin, syntonn, and peptone (A Dochmann, *Maly's Jahrsber* 11, 190). In colostrum casein is absent, or nearly so, being replaced by globulin (Sebelien).

The fats of milk—When milk stands, some of the fatty globules rise to the surface, and some of these are possibly freed from their proteid casing (Müller), thus forms the layer of cream. Butter is the name generally given to the fat itself, which is separated from the milk globules by mechanical agitation (churning), a certain amount of casein is, however, generally present in butter, and about one third of the fat of the milk remains in the butter milk. The fats present are olein, much palmitin, much less stearin, and about two per cent of the total fats consist of the triglycerides of butyric, caproic, and caprylic acids, with traces of myristic and arachidic acids. Cow's butter contains about 68 per cent palmitin and stearin, 30 per cent olein, and 2 per cent the other fats just mentioned. Rancidity is due to the development of free fatty acids, acrolein, &c., and, according to Hagemann, of lactic acid from the lactose admixed with fat in the butter.

Lactose or milk sugar (C₁₂H₂₂O₁₁) may be obtained from milk by evaporation in rhombic crystals containing one molecule of water of crystallisation, which is given off at 140°. It dissolves in 6 parts of cold and 2.5 parts of boiling water. It is not so sweet as cane sugar, its specific rotation is (α)_D = 59.3°. Dilute acids and hydrolytic ferments convert lactose into galactose, which is a glucose, and capable of undergoing the alcoholic fermentation, and which yields mucic acid when treated with nitric acid. Lactose reduces alkaline solutions of copper salts, and when oxidised yields mucic, saccharic, tartaric, and oxalic acid. Lactose does not itself undergo the alcoholic fermentation, but when much yeast is added fermentation occurs after some time, manite being formed. In presence of casein, &c., it undergoes the lactic acid fermentation, and it is the occurrence of this to which the souring of milk is due.

Preservation of milk.—Many different antiseptics have been suggested for the preservation of milk, e.g. salicylic acid, ether, benzene, chloroform, borax, boroglyceride, &c. Preservation is, however, usually effected by evaporating the milk to a thick syrup and adding considerable quantities of cane sugar. (For recent papers on this subject v A Mayer, *Maly's Jahrsber* 12, 168; P Biedert, Jacobi, Dietzsch, *ibid* 169,

O Loew, *ibid* 171, Morgen & Fleischmann, *ibid* 13, 174, A Baginsky, 175)

Alterations in milk—Many drugs taken in medicinal doses appear in the milk, *eg* salts of arsenic, antimony, lead, mercury (?), zinc, &c, also organic acids and alkaloïds. In various diseases the proportion of solids to water and of the different solid constituents is altered, thus in osteo malacia the lime salts are increased. Blood and pus may occur in milk, a blue colour sometimes observed is due to the growth of a chromogenic bacterium (*vibrio*), which, however, appears to be different from that which produces the blue colour in pus (J Reiset, *C R* 96, 682, 745). In tubercular diseases of cattle there is risk of contamination with tubercle bacilli, milk may also act as the carrier of other infections (scarlet fever, typhoid, &c)

ANALYSIS OF MILK

Qualitative—The specific gravity, which is increased by the removal of the cream, and reaction are ascertained in the usual way, the milk may then be examined microscopically. Casein may be precipitated by acetic or hydrochloric acid, by saturation with magnesium sulphate, or by rennet coagulation. The precipitate contains also the fat which may be separated from the casein by ether. The filtrate contains the sugar, salts, and the other proteids of the milk, which may be identified by their usual tests. The butter is best obtained from milk by adding half its bulk of 10 p.c. caustic potash solution, and then shaking the mixture rapidly with twice its volume of ether, on evaporating the ethereal extract the fat is left as a residue. The milk globules may be separated from the rest of the milk by filtration through porcelain under pressure.

Quantitative—**Solids** To 10 grains of dry sand add 5 c.c. of milk, and dry at 100° to constant weight. The increase in weight gives the solids, in 5 c.c. of milk. Below 10.5 p.c. of solids in cow's milk indicates dilution.

Butter This may be estimated by weighing the amount of residue from an ethereal extract of milk to which an equal volume of 10 p.c. caustic potash has been added. The normal minimum for fats is about 2.75 p.c. (Cameron). There are also various optical methods which depend on the degree of opacity of milk, which varies directly as the number of globules present, galactoscopes have been invented by Donné and Vogel. (On the lacto butyrometer, an instrument for estimating the fat by the ether method, *v* Tollens & Schmidt, *Maly's Jahrsber* 8, 140, Kehrer, *ibid* 11, 179, F Soxhlet, 180, Egger, 181, M Schmöger, 182.) **Cream** This may be roughly estimated by allowing the cream to rise in a narrow graduated vessel, the thickness of the layer being read off at the end of 24 hours. Good milk should yield 10 to 15 p.c. of cream. **Proteids** Pfeiffer's method is a good one, the casein is precipitated by acetic or hydrochloric acid, and freed from fat by ether, after filtering off the casein the albumin is precipitated in the filtrate by boiling, and after that has been removed the albumose is precipitated by tannin. It must, however, be remembered, as stated above, that albumose is an artificial product. Each of these precipitates is collected, dried, and weighed

(*Maly's Jahrsber* 13, 170, *v* also Parr, *ibid* 15, 185) J Sebelien (*Zeit Physiol Chem* 13, 135) estimates the proteids by the amount of nitrogen in the precipitates produced by various reagents. **Lactose** The casein, fat, and albumin having been removed, the milk sugar is converted into dextrose by boiling with sulphuric acid, and this is estimated by Fehling's solution or by the polarimetric method.

[In addition to the papers already quoted abstracts of the most important of recent papers on milk analysis will be found in *Maly's Jahrsbericht* as follows: J Forster, 11, 177, Giunti, 178; Weiske & Kennepohl, 187, de Leon, 12, 151; Knaach, 165, Stenberg, 161, Emmerich, 165, Fleischmann & Morgen, 166, Jorgensen, 167, F Hofmann, 177, Liebermann, 13, 168, C H Wolff, 169, Schrodtt & Haussen, 14, 180, *v* also Schmidt Mulheim, *Pf* 30, 602.] *Fleischmann, Das Molkerwesen*, and Tatlock, *The Produce of the Dairry*, Glasgow, 1888, should be also consulted. W D H

MILK SUGAR *v* SUGAR

MILLET OIL Appears to contain an acid $C_{11}H_{22}O_2$. On fusion with potash oil of millet yields acetic and lauric acids. On distillation it yields an acid $C_8H_{16}O_2$ or $C_9H_{18}O_2$ (216°–220°). Oxidation by $KMnO_4$ forms an hexoic acid $C_8H_{16}O_2$, and two oxy acids $C_{11}H_{20}(OH)_2O_2$ [60°], and oxy hexoic acid [108°], which yields an acetyl derivative $C_8H_{11}(OAc) CO_2H$ [71°] (Kaasner, *Ar Ph* [3] 25, 1081).

MILLON'S REAGENT A solution of mercuric nitrate containing nitrous acid, prepared by dissolving mercury (1 pt) in HNO_3 (1 pt) diluted with water (4 pts). It gives a red coagulum when warmed with albumen (Millon, *A* 72, 349).

MINERAL ACIDS A term sometimes applied to acids formed of elements other than carbon. The term is chiefly applied to the stronger commonly used acids, sulphuric, nitric, and hydrochloric.

MINERALOGICAL CHEMISTRY Mineralogical chemistry is that part of chemistry which relates to the mineral products of nature, it is also a part of mineralogy, the science which treats, not only of the chemical characters of these products, but also of their physical and geometrical characters, of their modes and places of occurrence, and of their classification.

Mineral products, as they are found in nature, frequently retain much the same character over a considerable extent of country and sometimes for a considerable depth, to such a product the term *rock* is applied, and to each kind of rock is given a special name. Mineral products are either simple or mixed. When simple they cannot be resolved, by mechanical or optical means, into matter presenting different characters, simple mineral products are briefly designated as *minerals*. Rocks are almost always mixed products. Generally the presence of different substances is obvious to the unaided eye, and the constituents can be readily separated from each other by mechanical means. From the rock termed *granite*, for example, may be got at least three kinds of matter: firstly, a substance yielding rhomboidal fragments when broken (*felspar*), secondly, a glassy substance yielding only irregular fractures

(quartz); and thirdly, a substance which may readily be split into very thin plates (*mica*). By no mechanical process can these substances be resolved into others having different characters, hence they are termed *minerals*, and as they present manifold differences from each other they have received distinctive names. Some times the composite nature of a rock is not evident until a slice, made sufficiently thin to be transparent, is examined with a microscope. Or, again, a rock, as for example *marble*, may consist wholly of a single kind of matter.

All simple minerals having the same essential characters are said to belong to the same *species*, those belonging to the same species may likewise have one or more unessential, yet more or less important, characters in common, and are then said to constitute a *variety* of that species.

It is the province of the mineralogist to discover what characters are presented by minerals, to name the characters, to determine their relative importance, to name the species and their varieties, to classify the species, and so on. Such a study of minerals must have been in progress since the earliest times. Even before 800 B.C. the study was so far advanced that minerals were classified as *metals* (i.e. those having a metallic lustre), *stones*, and *earths*, the characters then recognised as pertaining to *stones*, and used in their discrimination, were succinctly referred to by Theophrastus as follows—

‘There are in stones of different kinds many peculiar qualities, of which colour, transparency, brightness, density, hardness, and the like are frequent, though other more remarkable properties are not so common. But besides these qualities there are others such as their acting upon other bodies, or being subject or not subject to be acted upon by them. Some are fusible, others will never liquefy in the fire, some may be calcined, others are incombustible to which it may be added that in the action of fire on them they show also many other differences. Some, as amber have an attractive quality. Others serve for the trial of metals, as the Lydian stone. But the most known and general properties of stones are their several fitnesses for the various kinds of work, some of them are proper for engraving on, others may be shaped by the turner’s tools, others may be cut or sawn, some also there are which no iron instruments will touch, and others which can with difficulty, or scarcely at all, be cut by them.’

In this way it was possible even for the Ancients, without any knowledge of either crystalline form or chemical composition, to determine more or less satisfactorily the species met with in the mineral kingdom.

The definition and classification of species by means of what have been termed ‘external characters’ (such as crystalline form, cleavage, density, hardness, colour, lustre, &c.)—i.e. characters which can be determined without involving a total change of the characters of the mineral—had been found so far satisfactory that the external characters were used both for the definition and classification of the species by the best mineralogists even so lately as the early part of the present century. The most important of the external characters is now recognised to be the crystalline form. It was shown in the year 1772 that all the crystalline forms of the same substance, whether the substance is natural or artificial, are so closely related to each other that they are derivable by simple laws from a single form, and it has

since been proved that the whole series of crystalline forms of the same substance can be defined by means of numerical values which fix the shape of one of its forms, hence the limitation of each species to specimens which can be referred to one fundamental crystalline form became a necessity (v. CRYSTALLISATION, vol. II p. 278).

The recognition of chemical elements and chemical compounds, and of the constancy of the proportion of the elements in each compound, the setting forth of the atomic theory and chemical formulae, and the development of analytical methods, had great influence on the advance of mineralogy. When the specimens belonging to a species which had been defined by means of external characters were subjected to chemical analysis, it was discovered that in many cases there was a large variation in the proportion, and even in the nature, of the constituent chemical elements. For a time it was supposed that, while some substances have a great tendency to crystallise, others are comparatively inert, that, for example, the crystalline form of dolomite (carbonate of calcium and magnesium) is due entirely to the crystallising power of the carbonate of calcium, and that the carbonate of magnesium is either a mere inactive inclusion, or so feeble a crystalliser as to be completely overpowered by the carbonate of calcium. The explanation of these variations of chemical composition within species, as defined by means of the external characters, was eventually furnished by the principle of *Isomorphism* (q.v.), and it then became possible to define a species by means of a combination of only two characters—chemical composition and crystalline form. In other words, it was found that all specimens which can be referred to the same fundamental crystalline form and to the same chemical type have all other essential characters in common, though they may differ in the less important ones, such as colour or transparency. For example, if the forms of crystals can be referred to the same fundamental figure, the differences of crystalline development are themselves unessential, hence the substances popularly known as *dogtooth spar* and *needle spar*, which can be crystallographically referred to the same rhombohedron, and chemically are carbonate of calcium, are regarded as varieties of the species *calcite*, which includes all specimens having these two fundamental characters in common. Similarly, colour is another unessential character the specimens belonging to the species *fluor*, for instance, show a remarkable variation in colour. In some cases a variety characterised by a particular colour receives a distinctive name, *emerald* and *aquamarine*, for example, are respectively dark green and light bluish green varieties of the species *beryl*. It is very seldom, however, that varieties are really worthy of being distinguished by special names, and in almost every species the transition from one variety to another is so gradual that it is often impracticable to decide on the single variety to which a specimen may be satisfactorily referred. On the other hand there are specimens which, though chemically carbonate of calcium, can all be referred to a single fundamental crystalline form quite distinct from that of

calcite They are therefore regarded as belonging to a distinct species and are grouped together under the name of *aragonite*. That such a separation is a legitimate one is shown by the fact that the two species, as thus defined, differ in other important characters, such as cleavage, hardness, specific gravity, optical constants, &c. Hence the chemical compound calcium carbonate is said to be *dimorphous*, but, as already indicated, the difference between the two kinds of substance is much more than one of form. Titanic oxide, again, is met with in nature as three different kinds of substance, each presenting its own set of characters, they are necessarily to be regarded as distinct species, and have received the specific names *rutile*, *anatase*, and *brookite*. titanic oxide is said to be *trimorphous*. The variety of chemical composition among specimens regarded as belonging to a single mineral species, and which is explicable on the above mentioned principle of isomorphism, is well illustrated by the results of analyses of specimens of *tetrahedrite* (or *grey copper ore*). The crystals of this mineral belong to the cubic system and are hemisymmetrically developed, being all closely related in figure to the regular tetrahedron, chemically they can be referred to the general formula $4R'SR''S_3$, where R' includes chiefly Cu, Fe, Zn, Ag, and Hg, and R'' includes Sb, As, and Bi. The composition shows the following variations: copper 15 to 42 p.c., iron 1 to 7, zinc 0 to 7, silver 0 to 31, mercury 0 to 17, antimony 12 to 30, arsenic 0 to 20, bismuth 0 to $1\frac{1}{2}$, a few of the varieties, those containing noteworthy proportions of silver or mercury, for instance, have been distinguished by special names. The difficulty of deciding whether a mineral product is to be regarded as a distinct species or as a variety of another species presents itself in the case of the specimens to which the name of *tennantite* has been given, crystallographically they show the same relationship to the regular tetrahedron which characterises the specimens of *tetrahedrite*, and chemically they are represented by the same general formula $4R'SR''S_3$, they are distinguished, however, by the absence of antimony (and bismuth) as in the chemical composition of the specimens belonging to the above crystallographic and chemical type there is thus a sudden transition from 12 p.c. of antimony (generally accompanied by more or less arsenic) to zero, the specimens of *tennantite* are generally regarded as entitled to rank as a distinct species. Similarly *aragonite*, *witherite*, *strontianite*, and *cerussite* belong to one chemical type $R''CO_3$, and are almost identical in their fundamental crystalline forms, for the several minerals R'' is essentially Ca, Ba, Sr, and Pb they are regarded as distinct species because in nature there is found no gradual transition of chemical composition. On the other hand, the isomorphous rhombohedral carbonates belonging to the above chemical type $R''CO_3$, where R'' is Ca, Mg, Fe, or Mn, present such a gradual shading into each other, both chemically and physically, that a perfect definition of species is impossible. Another instructive illustration of this difficulty is furnished by the group of minerals to which the name *garnet* has been given. The garnets crystallise in the cubic system, and in their forms the do-

decahedron is prominently developed, chemically they are represented by the general formula $3R''O R'''O_2 \cdot 3SiO_2$. They have in general very similar physical characters, though they differ considerably both in colour and specific gravity, they have in fact every claim to recognition as a natural family or group. But the differences of chemical composition are remarkable. R' and R'' are essentially, for *grossularite*, Ca and Al, for *pyrope*, Mg and Al, for *almandite*, Fe⁺ and Al, for *spessartite*, Mn and Al, for *andradite*, Ca and Fe⁺⁺⁺, for *brockbergite*, Ca, Mg, and Fe⁺⁺⁺, for *uvarovite*, Ca and Cr. As the above kinds of garnet are connected by various transitions, it may easily be imagined that the garnet group presents great difficulty as regards the definition of its species, and that it is possible to introduce a large number of unnecessary names into the science.

There are other natural groups or families, however, as for example, the *augite*, *hornblende*, *scapolite*, *felspar*, *mica*, *chlorite*, and *tourmaline* groups, in which the representation of the chemical composition of the whole group by a single chemical formula presents great difficulties. Indeed, the formulæ which are employed in mineral chemistry are at present for the most part empirical, constitutional formulæ, such as are employed in organic chemistry, are almost unknown. Much work must be done before we obtain a real insight into the structure of the more complex silicates. For attempts in this direction the student may refer to the following —

Augites and Hornblendes Tschermak's *Mineralogische Mittheilungen*, 1871 17, *Neues J M* 1, 43

Scapolites Sitz W 1863 [1st part] 1142

Felspars *ibid* 1864 [1st part] 566

Micas *ibid* 1878 [1st part] 5, *Am S* [3] 38, 384

Clintonites Sitz W 1878 [1st part] 555

Tourmalines P 139, 379, 547

Silicates in general Z K 17, 25

Minerals are of interest to the chemist as the source of the various chemical elements, the characters of which, and of their combinations, it is his province to study. A few of these elements are found in the uncombined state, notably carbon, sulphur, arsenic, bismuth, copper, silver, gold, and platinum. Some are present in large proportion in numerous mineral species, the latter themselves plentifully dispersed through the earth's crust, others are found in few mineral species, and these only in small quantities and in few localities.

It is the province of the mineralogical chemist to determine the chemical composition of each species and variety, and to represent it by means of an empirical or constitutional formula. A considerable number of minerals are pure or nearly pure chemical compounds of simple constitution, and these are specially interesting to the chemist, as they often furnish him with beautifully crystallised specimens of compounds which, as laboratory products, are either amorphous or are obtained as minute crystals only with much expenditure of time and trouble, among such compounds we may especially note the sulphides of lead, zinc, copper, iron, antimony, and arsenic, the various sulpharsenites

and sulphantimonites, the oxides of aluminium, iron, titanium, silicon, and tin, various silicates, molybdate, tungstate, and chromate, of lead, tungstate of calcium, the sulphates and carbonates of calcium, strontium, barium, and lead, various phosphates, arsenates, and vanadates, of copper, lead, and calcium

Time and energy are now rarely expended in the chemical analysis of a substance unless there is something extraordinary in its external characters, and it is by means of these characters that it is possible to describe for future recognition the substance of which an analysis has been made

Mineralogical chemistry deals, too, with the classification of all the chemical compounds met with in the mineral kingdom, in fact, the arrangement in large groups is now generally based on chemical composition, though the definition of the species is made to rest also on the crystalline form. The system of classification now generally adopted is as follows —

Division I Native elements, metallic and non metallic

Division II The compounds of metals with elements of the arsenic and sulphur groups, viz arsenides, sulphides, arseno sulphides, sulpho salts, and their analogues

Division III Chlorides, and their analogues

Division IV Compounds of oxygen (a) oxides, (b) oxy salts, namely, (1) carbonates, (2) silicates and titanates, (3) molybdates and tungstates, (4) chromates and sulphates, (5) borates, (6) nitrates, and (7) phosphates, arsenates, and vanadates

For details v Groth's *Tabellarische Uebersicht der einfachen Mineralien*, Braunschweig, 1889

The mineralogical chemist observes the reactions of minerals with various reagents, both in the wet and dry way, and on a large or a microscopic scale, and by a classification of reactions he provides means for the determination of the species by chemical methods

To the mineralogical chemist is further assigned the most important task of discovering the modes in which the various chemical compounds may be produced artificially, and the investigation of the processes and reactions by which these chemical compounds have been actually produced in nature. In this respect *pseudomorphs* (i.e. minerals presenting a form characteristic not of their own but of some other substance) are of great value. They are always results of chemical change, and are produced in various ways. Some are merely due to the investment of a substance afterwards removed from beneath the crust, these have been termed *epimorphs*. For example, hollow, well defined scalenohedra of *limonite* (hydrated ferric oxide) are met with, they owe their form to crystals of calcite upon which the limonite has been deposited. In other cases the original mineral is altered throughout its mass, and suffers a loss of a chemical constituent, as when *galena* (sulphide of lead) is found with the form of *anglesite* (sulphate of lead). Or, again, there has been an addition of a chemical constituent, as when *chessylite* (a hydrated carbonate of copper) is found with the form of *cuprite* (oxide of copper). Or, again, there has been an exchange of chemi-

cal constituents, as when *galena* is found with the form of *pyromorphite* (phosphate and chloride of lead), or *calcite* with the form of *gypsum*. Sometimes there is a complete exchange of material without loss of form, as when copper is found with the form of *aragonite*. Pseudomorphs illustrate the decomposing influences to which many minerals have been subjected, and throw valuable light on the order of succession in which, and the conditions under which, particular minerals have been formed and deposited and in furnishing sure proofs of conversions which we cannot hope to effect in the laboratory, they afford a knowledge of facts which can be arrived at in no other way

The following works, relative to the artificial production of substances met with as minerals, may be consulted —

Fuchs, *Die künstlich dargestellten Mineralien*, Haarlem, 1872

Daubrée, *Études synthétiques de géologie expérimentale*, Paris, 1879

Fouqué & Lévy, *Synthèse des minéraux et des roches*, Paris, 1882

Bourgeois, *Reproduction artificielle des Minéraux*, Paris, 1884

MINT The oil of spear mint (*Mentha viridis*) contains $C_{10}H_{16}O$ (225°), S G 952, and a terpene (Gladstone, J 1863, 548, cf Kane, A 32, 286)

Peppermint v MENTHOL

MOCHYLIC ALCOHOL $C_{15}H_{26}O$ [234°] Present in bird lime as mochyl palmitate $C_{15}H_{31}O_2$ (Divers a Kawakita, C J 53, 274) Slender lustrous prisms, insol water, v sl sol petroleum ether, v sol ether, m sol alcohol. Dissolves like birdlime in conc H_2SO_4 with a red colour. Heated with palmitic acid in sealed tubes to 160° a substance very similar to bird lime is obtained. Yields on distillation a hydrocarbon $C_{15}H_{32}$

MOLECULAR CONSTITUTION OF BODIES, THEORIES OF The theory of the molecular constitution of matter now universally accepted was held long before any crucial proof was given of its necessity. For though such phenomena as the enormous changes in volume which take place when a gas condenses to a liquid point most strongly to some such view, they cannot be held to be conclusive, unless it is considered axiomatic that a perfectly homogeneous structure is incapable of dilatation or contraction. The first attempt to give more elaborate reasoning in favour of the molecular theory of the constitution of matter seems to have been made by Cauchy, and was founded upon the dispersion which light experiences when it passes through transparent bodies. Since the velocity of light when passing through such bodies depends upon the wave length of the light, and also on the nature of the body, Cauchy argued, that since a velocity is not of the same dimensions as a length, the velocity cannot depend upon the wave length absolutely, but must depend merely upon its ratio to some other length, now the only length available is one derived from the body itself, and since the dispersion does not depend upon the dimensions of the transparent body, this length must be one intrinsic to the body, the body therefore cannot be homogeneous and without structure or

there would be no such length available; the body must therefore be coarse grained. This reasoning, however, is not quite conclusive, for light of different wave lengths has different times of vibration, and the phenomenon of dispersion might be expected if there were any time connected with the structure of the body which could come into comparison with the time of vibration of the light. Dispersion proves that the transparent body is coarse grained, either with respect to space or time, but does not prove that it is necessarily coarse grained with regard to space on a scale comparable with the wave length of light. In fact, in the theories which have been put forward to explain dispersion, this phenomenon is made to depend upon the ratio of the time of vibration of light to some time of vibration of the molecules of the system. A less ambiguous proof of the molecular structure of gases was given by Osborne Reynolds in his paper '*Some Dimensional Properties of Matter in the Gaseous State*' (*T* 1879). This proof was founded on the phenomenon called the 'thermal effusion' of gases. This phenomenon is of the following kind.—If we have a vessel divided into two portions by a porous diaphragm, say of stucco or meerschäum, and the gas on one side of the diaphragm is kept at a different temperature from that on the other side, it is found that, in order to prevent the gas flowing from the cold to the hot side of the diaphragm, the pressure of the gas on the hot side must be greater than that on the cold side, and that the difference of pressure required to prevent the flow obeys different laws according as the gas is dense or rare. If the gas is dense, this difference of pressure varies inversely as the density of the gas, while if the gas is rare, the difference of pressure varies directly as the density. Reynolds found that the density of the gas at which the law changed from one form to the other depended upon the fineness of the pores of the diaphragm, the finer the pores, the greater was the density at which the law changed. Thus, since the law connecting the difference of pressure with the density depends upon the diameter of the pores of the diaphragm, there must be some length in the gas with which this diameter of the pores can come into comparison, the gas must therefore have structure, and since the density of the gas when the law changes is greater for small pores than for large ones, the structure of the gas must be finer at great densities than at low ones. The investigations of Sir William Thomson (*Popular Lectures and Addresses*), Loschmidt, and others, have gone further than this, and have not merely furnished proofs that matter has structure, but have given limits below which the coarse grainedness of matter cannot lie. These investigations are founded on considerations about surface tension, the difference of potential which occurs when two metals are put in metallic connexion, the amount of polarisation at the surface of an electrode and an electrolyte, the viscosity, the diffusion, and the conductivity for heat, of gases. It will be sufficient for us to show how one of these leads to a limit for the dimensions of molecular structure, and we will take the one depending on the surface tension. The surface-

tension of a film of liquid is not likely to alter until the thickness of the film falls below the distance at which one molecule ceases to exert an appreciable influence on another, for it is only the molecules within a film of this thickness which can exert any influence on those at the surface, so that if we can find a limit to the thickness of a film which possesses an unaltered surface tension, we shall have a quantity comparable with the distance up to which one molecule exerts an appreciable effect on another. When a film is stretched, work is done on it against surface-tension, and this work is stored up in the film, so that if the surface-tension were the same for an infinitely thin film as for one of finite thickness, an infinite amount of work could be stored up in the film. But the greatest amount of energy which can be stored up in, say, a gram of water must be less than the amount required to separate the molecules to such a distance that they no longer exert any influence on each other, but this is exactly what is done when the water is vaporised, so that the greatest amount of energy which can be stored up in a gram of water is less than the amount required to convert it into steam. To convert one gram of water at 15° into steam requires the expenditure of $621 \times 4.2 \times 10^7$ ergs. If T be the surface tension of water, and if the gram of water is stretched into a thin film whose thickness is x , the work done in stretching it is $\frac{2T}{x}$, but this is not all the energy which

is communicated to the film, for, unless heat is supplied to the film as it stretches, it will cool, the amount of heat which must be supplied to the film, when measured in mechanical units, is about half the work done in stretching the film, so that the total energy communicated to the film is $3T$, this must be less than the work required

to vaporise the film, so that $\frac{3T}{x}$ must be less than $536 \times 4.2 \times 10^7$, or putting T at 15° equal to 74 , x cannot be less than 8.5×10^{-7} cm, that is, a thickness of 10^{-6} cm must be comparable with the range of molecular action of the water molecules.

Quincke (*P* 137, 402) investigated the range of molecular forces, by finding the thickness of a silver film which when deposited on glass just began to alter the capillary ascent or depression of water in contact with the glass. The results of this, and other investigations with a similar object, are summarised in the following extract from a paper by Rucker (*C J Trans* 1888 260) —

Table of properties of thin films and of molecular magnitude

Thickness of film = 118×10^{-7} cm

Superior limit to the radius of molecular action, deduced from Plateau's experiments (*Statique des Liquides*, 1873, i 210) on the pressure of a soap bubble, by using Maxwell's theory that the surface tension first diminishes when the thickness of the film equals the range of molecular action, $96 \times 10^{-7} - 45 \times 10^{-7}$

Between these limits the thickness of a film begins to be unstable, that is, the surface tension

begins to diminish Hence the radius of molecular action must be $< 96 \times 10^{-7}$ and $> 22 \times 10^{-7}$ cm

$$50 \times 10^{-7} \text{ cm}$$

Value of the range of molecular action deduced by Quincke (*P* 137, 402) from experiments on capillary elevation

$$12 \times 10^{-7} \text{ cm}$$

Average thickness of black soap films, measured by two independent methods

As the tension of a black film is equal to that of a thick film, the surface tension, which begins to diminish at 50×10^{-7} cm, must increase again, and reach its original value at 12×10^{-7} cm This is also about the thickness below which, according to O Wiener (*W* 31, 629), a thin silver plate will no longer produce the same effect on the phase of reflected light as a thick silver plate would do

$$10.5 \times 10^{-7} \text{ cm}$$

Thickness of the permanent water film observed by Bunsen (*W* 24, 322) on unwashed glass at a temperature (23°C) at which the vapour pressure of water is small

$$4 \times 10^{-7} \text{ to } 3 \times 10^{-7} \text{ cm}$$

Average distance from centre to nearest centre of molecules in gases under standard conditions, calculated by Meyer (*Die kinetische Theorie der Gase*)

$$3 \times 10^{-7} \text{ cm}$$

Thickness of metal films required to polarise platinum completely (Oberbeck, *W* 31, 331)

$$10^{-7} \text{ to } 2 \times 10^{-8} \text{ cm}$$

Thickness of electric double layer, according to Oberbeck and Falck (*W* 21, 157)

$$2 \times 10^{-8} \text{ cm}$$

Smallest thickness of silver which affects the phase of reflected light (Wiener, *l.c.*)

$$1.4 \times 10^{-8} \text{ to } 1.1 \times 10^{-8} \text{ cm}$$

Diameter of gaseous hydrogen molecule

$$7 \times 10^{-8} = 2 \times 10^{-8} \text{ cm}$$

This is given by combining (1) the specific inductive capacity and coefficient of viscosity, (2) the refractive index and coefficient of diffusion, (3) the law of expansion and the thermal conductivity

Average distance between centre of molecules supposed arranged uniformly in liquids and solids according to Thomson

$$2 \times 10^{-8} \text{ cm}$$

Inferior limit to the diameter of a gaseous molecule according to Thomson These results may be shortly summed up as follows —

10^{-7} cm		
118 × 1	Superior limit to range of molecular action	Plateau, Maxwell.
96—45	Range of unstable thickness begins.	Reinold and Rücker
59	Superior limit to range of molecular action	Plateau.
50	Magnitude of range of molecular action	Quincke
12	Range of unstable thickness ends.	Reinold and Rücker
12	Action of silver film on phase of reflected light alters.	Wiener
10.5	Thickness of permanent water film on glass at 23°	Bunsen.
4—3	Mean distance between centres of nearest molecules in gases at 760 mm and 0°C	O Meyer

10^{-7} cm.		
3—1	Thickness of metal films which polarise platinum	Oberbeck.
1—0.02	Thickness of electric double layer	Lippmann and Oberbeck.
2	Smallest appreciable thickness of silver film.	Wiener
14—11	Diameter of gaseous hydrogen molecule	Exner O Meyer Van der Waals.
07—02	Mean distance between centres of nearest liquid molecules	W Thomson.
02	Inferior limit to diameter of gaseous molecule.	W Thomson

Having obtained some idea of the coarseness of the structure of matter, we shall now consider various theories of that structure In order to see what has been explained by these theories, and what remains to be explained, let us enumerate the most important properties of matter in that state of aggregation when the properties are the most simple, *z.e.* the state of a so called 'perfect' gas

1 The relation between pressure, density, and temperature is expressed by the laws of Boyle and Charles, $p = \kappa \rho \theta$, when p is the pressure, ρ the density, and θ the absolute temperature of the gas, and κ is a quantity which remains constant for the same gas For different gases κ is inversely proportional to their combining weights In all such gases there is the same number of molecules in unit volume, provided the pressure and temperature are the same

2 The gases possess viscosity The coefficient of viscosity (unless the density is very greatly reduced) is independent of the density, but depends upon the temperature The most recent experiments show that the viscosity varies as the two third power of the absolute temperature (Barus, *Bulletin of the U S Geological Survey*, No 54, 1889)

3 The gases conduct heat with a facility depending on the temperature

4 They diffuse into each other with a rapidity depending upon the density and the temperature

5 They possess specific heats of various kinds, the ratio of the specific heat under constant pressure to that under constant volume being 1.4 for most gases

6 They exhibit phenomena of the type of those which occur in the radiometer

7 When they are raised to a high temperature they become luminous, and give out rays of definite periods, the periods being independent of the temperature They absorb light of the same periods as those they give out when hot

8 They possess very different electrical properties For low differences of potential they insulate almost perfectly, but when the electric intensity is raised sufficiently a spark passes through them, and the electricity is discharged

9 The various gases exhibit very different chemical properties

According to the kinetic theory of gases a gas consists of a great number of small particles called molecules, moving about with great velocity in all directions, but continually either striking against each other, or coming so near together that they exert forces on each other which produce effects somewhat similar to those produced by the collision of two elastic balls This theory has been shown by Maxwell and

Clausius (Maxwell's *Theory of Heat*, O E Meyer's *Die kinetische Theorie der Gase*) to be sufficient to explain all those properties of gases included under (1). These properties are independent of the nature of the molecule, and of the exact way in which two molecules act upon each other. They could be deduced equally well whether we supposed the molecules to be hard elastic spheres, or systems attracting or repelling each other when they come near to gether, or vortex rings. When we consider other properties besides those included in (1), we find the explanation less satisfactory. The theory gives an explanation of the viscosity, diffusion, and conduction of heat, but the exact way in which these properties vary with the temperature depends upon the nature of the action between the molecules. Two cases have been worked out by Maxwell. In the first case the molecules were supposed to be hard perfectly elastic spheres, and which only acted upon other molecules when in collision with them. According to this hypothesis, the coefficient of viscosity would be proportional to the square root of the absolute temperature. The other case which has been worked out by Maxwell is that in which the molecules are regarded as systems repelling each other with forces which vary inversely as the fifth power of the distance between them. According to this hypothesis, the coefficient of viscosity would be proportional to the absolute temperature. The experiments of Barus (*l c*) and others show that the coefficient of viscosity obeys neither of these laws, but varies more quickly with the temperature than the result obtained on the first hypothesis, and more slowly than that obtained on the second. Neither of these theories of the action of one molecule on another can be the true one. Sutherland (*P M* 24, 113, 168), by considering the results of Thomson and Joule's experiments on the cooling of gases passing through a porous plug, arrives at the conclusion that the force between two molecules must be inversely as the fourth power of the distance between them. The value of the temperature coefficient of the viscosity on this hypothesis has not, however, been worked out.

The fact that the ratio of the specific heat at constant pressure to the specific heat at constant volume is the same for the perfect gases with the same number of atoms in the molecule has not been explained by the kinetic theory, and in fact the results we should expect from the kinetic theory are so different from those actually observed that they constitute perhaps the gravest difficulty which the kinetic theory has yet encountered. The results to which the kinetic theory leads are easily found. Let T_m be the total kinetic energy of the molecules in unit mass, β the ratio of this to the energy due to the translatory motion of their centres of gravity, p the pressure, v the volume of unit mass, and θ the absolute temperature. Then

$T_m = \frac{3}{2} \beta p v$, so that k the specific heat at constant volume will equal $\frac{3 \beta p v}{2 \theta}$. If the pressure is kept constant, then, in addition to the energy spent in warming the gas, an amount

of work— $p \times$ (change in volume for one degree)—is done, but when the pressure is constant the volume is proportional to the absolute temperature, so that the change in volume for one degree = v/θ . Thus, if k_p be the specific heat at constant pressure, then

$$k_p = \frac{3 \beta p v}{2 \theta} + \frac{p v}{\theta},$$

$$\text{so that } \frac{k_p}{k_v} = \frac{3}{2} \beta + 1$$

or if the ratio of the specific heats be denoted by γ

$$\gamma = 1 + \frac{2}{3\beta}, \quad \beta - 1 = \frac{5 - 3\gamma}{3(\gamma - 1)}$$

Now $\beta - 1$ is the ratio of the internal energy of the molecules to that due to the motion of translation of their centres of gravity, and we see from the preceding equation that this depends only upon the ratio of the two specific heats, the constancy of this ratio for different gases shows that the proportion which the vibratory energy bears to the energy of translation must be the same for all such gases, so that at the same temperature the vibratory energy of all these gases must be the same. Now the number of lines visible in the spectrum of the various gases is very different, and we should therefore expect the gases to have very different capacities for vibratory energy. The fact that it is not so seems to indicate that the vibratory energy is not due, at any rate at low temperatures, to those quicker modes of vibration which manifest themselves as light, but must be due to some other modes common to all gases. This mode in a diatomic gas may, possibly, be the motion of the atoms relatively to each other, and this view is strengthened by the fact that the ratio of the vibratory to the translatory energy increases with the number of atoms in the molecule. Thus, if we take Dulong's values of the ratio of the specific heats for hydrogen, carbon dioxide, nitrous oxide, and ethylene, the values of the ratio of the vibratory energy to the energy of translation are respectively 75, 11, 11, and 2, thus, for these gases the ratios are very approximately proportional to the number of atoms in the molecule, and the result suggests that the vibratory energy at these low temperatures is that of the atoms in the molecule relatively to their common centre of gravity, and not of the quicker modes of vibration corresponding to the luminous rays. If experiments on the ratio of the specific heats could be made at temperatures high enough to produce luminosity of the gas, it is possible that the results might be different from those made at lower temperatures, and that they might be found to vary from gas to gas.

Theories of the Constitution of the Molecule
The kinetic theory of gases, as developed by Clausius and Maxwell, dealt chiefly with the behaviour of a large number of molecules, and said little about the constitution of individual molecules. Quite recently, however, Sir W Thomson (*Lectures on Molecular Dynamics and the Wave Theory of Light*) and Lindemann (*Ueber Molekularphysik, Physikalisch Gesellschaft, Königsberg, 39 [1888]*) have discussed the properties of a molecule consisting of a number of spherical

shells, one inside the other, each shell being connected to the one next it by an elastic spring. The external spherical shell is supposed to be acted on by the ether, or, what is the same thing, by a periodic force whose period is that of the vibrations which the ether is transmitting. We may say in passing that many of the results obtained do not depend upon this special view of the construction of the molecule, but would be true if we supposed the molecule to be a dynamical system whose configuration could be fixed by n co ordinates, that is, a system possessing n degrees of freedom. The behaviour of such systems when light falls upon them is investigated in the papers above mentioned, and expressions are obtained for the refractive index of a medium consisting of molecules of this kind for light of any period. These expressions explain the dispersion of light, and the results deduced from them agree with those found by experiment, they also explain the anomalous dispersion of the rays whose periods nearly coincide with those which are absorbed by the medium—a subject which was investigated experimentally for several substances, especially a solution of fuchsin in alcohol, by Kundt and Christoffel—the periods of vibration of the glowing gas being the periods of vibration of the spherical shells which constitute the molecule. The nature of the molecules assumed in this theory imposes a limit to the amount of energy due to the relative motions of the shells, for it is evident that the amplitude of vibration of any shell cannot be greater than the difference between its radius and that of a neighbouring shell. Thus, on Lindemann's theory, the internal kinetic energy reaches a maximum, and when it has reached this maximum any further exposure to light must lead to an increase in the translatory energy, and thus to an increase in the temperature of the system made up of such molecules.

Lindemann explains the development of heat which occurs on chemical combination by the transference of the internal kinetic energy into translatory energy, the substances after combination possessing less internal energy than before, the heat being produced by a loss of kinetic energy, and not, as in the ordinary explanation, by a loss of the potential energy of separation. If this view is correct, however, the internal kinetic energy must in certain gases be enormously greater than the translatory energy, thus, for example, in the combination of hydrogen and oxygen enough heat is produced to raise the temperature of the mixture nearly $20,000^{\circ}\text{C}$, and even assuming that after combination there is no internal kinetic energy, the internal kinetic energy before combination must be about 70 times the translatory energy. It is very difficult to see how this can be reconciled with the value found by experiment for the ratio of the specific heat at constant pressure to that at constant volume.

According to Lindemann, the electric properties of bodies are to be explained by internal kinetic energy due to vibrations which are too quick to be visible, in fact, roughly speaking, electricity is ultra violet light. This view is open to many difficulties, one of which is that it would not lead us to expect the great differ-

ence that exists between the electrical properties of the atom and the molecule. A molecule seems to be almost electrically neutral, thus it is impossible to communicate a charge of electricity to the molecules of a gas, though when the molecule is split up into atoms it exhibits most energetic electrical properties. Karl Pearson (*Proc London Math Soc*, 20, 38) has developed a theory in which the molecules are supposed to be made up of spheres vibrating in an uncompressible fluid, he obtains results similar to those of Sir W Thomson and Lindemann. In fact, as we said before, many of these results do not depend upon the nature of the molecule, but would be true if we supposed the molecule to be a dynamical system possessing n degrees of freedom.

Evidence as to molecular structure afforded by the spectra of bodies.—If we consider the oscillations of a dynamical system possessing n degrees of freedom, we find that there are n periods of vibration given by the roots of a determinantal equation, and that the relation between these roots depends on the nature of the system, a system could be constructed having n periods of any given value. If, however, the system possesses an infinite number of degrees of freedom, there will be an infinite number of periods, but the periods will be connected by more or less simple relations. Thus, if the vibrating system were like a stretched string, the periods would be proportional to the natural numbers, while if it were like a bar, the periods would be proportional to the natural numbers for the longitudinal and torsional vibrations, and to the roots of the equation $\frac{1}{x} \left\{ \epsilon^{\frac{1}{x}} + \epsilon^{-\frac{1}{x}} \right\} = \pm 2$ for the transver-

sal vibrations. If the system were a circular membrane, the frequencies would be proportional to the roots of an equation formed by equating a Bessels function to zero. If the system were a uniform elastic sphere, the frequencies would be the roots of a complicated equation given by Chree in the *Transactions of the Cambridge Philosophical Society* (14, 316, 317). Other periods which have been worked out are those of circular vortex rings. The frequencies of the higher vibrations about the circular form are proportional to $\sqrt{n(n-1)}$, where n is a large natural number, and the vibrations about the circular cross section are proportional to the natural numbers (J J Thomson, *On the Motion of Vortex Rings*, 35, 74). Many investigations have been made with the object of finding whether or not there are simple harmonics—that is, frequencies proportional to the natural numbers—in the spectra of bodies. In the case of the spectrum of hydrogen, Johnstone Stoney finds that the wave lengths 4102 37, 4862 11, 6563 93, which occur in this spectrum, are very accurately in the ratio $\frac{1}{32} : \frac{1}{7} : \frac{1}{5}$. Schuster, however, in his 'Report on the Genesis of Spectra' (*B A Reports*, 1882) says

'Other writers, as, for instance, Sorét (*P M* 1871 464), have from time to time drawn attention to harmonic ratios in various spectra, and the author of this report has during the last ten years collected a large quantity of material bearing on the question. The results have, on

the whole, not been favourable to the theory of harmonic ratios. In any spectrum containing a large number of lines it is clear that, owing to accidental coincidences, we shall always be able to find ratios which agree very closely with the ratios of small integer numbers. It is only by means of a systematic investigation that we can find out whether these coincidences are due to any real cause. We must, by means of the theory of probabilities, calculate the number of the coincidences, which we might expect to find on the supposition that the lines are distributed at random throughout the whole range of the visible spectrum. If on calculating out all fractions which can be formed in a spectrum by any pair of lines the number of ratios agreeing within certain limits with ratios of integer numbers greatly exceeds the most probable number, we should have reason to suppose that the lines are not distributed at random, but that the law suggested by Messrs. Lecoeq de Boisbaudran and Stoney is a true one.

The results of a long investigation conducted in this manner tend to show that the number of harmonic ratios is, if anything, smaller than was to be expected on the hypothesis of no connexion.

The simple harmonic ratio is not, perhaps, *a priori* the most probable relation between the periods. Balmer (*W* 25, 80) has shown that the wave lengths of a series of hydrogen lines are expressed by the formula $\frac{Cm^2}{m^2 - 4}$, where m

is an integer. Hagenbach (*Verhand d Naturforsch Ges zu Basel*, 1886) has compared the results of this formula with Cornu's measurements of the wave lengths of the hydrogen lines, the result of the comparison is given in the following table—

$$\lambda = 3645.42 \frac{m^2}{m^2 - 4}$$

Line	m	Calculated wave length	Observed wave-length	Difference
H α	3	6562.8	6563.1	+ 3
H β	4	4860.6	4860.7	+ 1
H γ	5	4339.8	4339.5	- 3
H δ	6	4101.1	4101.2	+ 1
H ϵ	7	3969.5	3969.2	- 3
H ζ	8	3888.4	3888.1	- 3
H η	9	3834.8	3834.9	+ 1
H θ	10	3797.3	3797.3	0
H i	11	3770.0	3769.9	- 1
H κ	12	3749.6	3750.2	+ 6
H λ	13	3733.8	3734.1	+ 3
H μ	14	3721.4	3721.1	- 3
H ν	15	3711.4	3711.2	- 2

These results seem to show that the hydrogen molecule is a system possessing an infinite number of degrees of freedom, and not a finite number of rigid particles mutually attracting each other.

It is worthy of notice that when m is large the formula previously quoted for the frequency of vibrations of a circular vortex ring becomes $\lambda = \frac{Cn}{n^4 - \frac{1}{2}}$ which is of the same type as Balmer's

A very striking feature in the spectra of some elements is the recurrence in the spectra of certain groups of lines—for example, triplets in the magnesium, doublets in the sodium, spectrum—and the most promising way of finding whether there is anything corresponding to overtones in the spectrum would be the investigation of the relation between the frequencies of the lines in these groups as they recur in the spectrum. It is stated by Schuster that no simple harmonic relations exist between these

groups. Deslandres (*C R* 104, 972) has shown that the periods of the recurring bands in the nitrogen spectrum are connected by a relation of the form $An + B$, where n is an integer.

The first explanation of the existence of these groups which suggests itself is that corresponding to a triplet we have three, to a doublet two, similar systems near together. Each of these systems, if free from the other's influence, would vibrate with the same period, but when placed so near together that they influence each other, the system of three will have three, and that of two will have two, nearly equal periods.

The theory of the oscillation of such systems shows that the gravest mode of the combined system will be lower, and the highest higher, than that of the original system. Thus in a triplet corresponding to each line of the original system, there will be three lines. If this view is correct, then any 'element' in whose spectrum doublets or triplets occur is capable of being split up into simpler systems, and the lines of the substance into which it is split up will be intermediate between those of the doublets or triplets. Thus, we should expect to find a tendency for these doublets to disappear as the temperature is raised. Though there does not seem much evidence to show that this tendency is widespread, it does appear to exist in the case of calcium, for in the drawing of the spectrum of this element given in Lockyer's *Studies in Spectrum Analysis* (191) there is in the violet end of the spectrum a doublet where the spark is taken without a jar in the circuit, but when a large jar is placed in the circuit the doublet is replaced by a single line intermediate to those of the doublet.

The widening of the lines of the spectrum of a gas when the pressure is increased might be explained on similar principles. A molecule when free from the influence of other molecules vibrates in certain definite periods, and shows sharp bright lines in its spectrum, when, however, it gets under the influence of another similar molecule its periods are slightly altered, and for each of the original periods we have two periods, the one graver and the other higher than the original period, the departure from the original period depending on the distance between the molecules. Thus, in the case of a gas so dense that the molecules influence each other, the molecules, instead of emitting light of a definite period, would emit light of different periods, some higher and some lower than the undisturbed one, thus, instead of a bright line in the spectrum, we should have a luminous band stretching across the original bright line.

Connexion between spectra of elements and those of their compounds—One of the most interesting subjects in connexion with molecular theories, and one from which we may hope to gain great insight into molecular structure, is the connexion between the spectrum of a compound and the spectra of its constituents. Mitscherlich showed that compounds have emission spectra of their own. A considerable amount of work bearing on the subject has been done by Gladstone and Dale, Abney and Festing, Kundt, and others, who have investigated the absorption spectra of compounds. But, however important these researches are from other points of view,

they cannot be said to have as yet thrown much light on the structure of matter

A relation between the lines in the spectrum of a compound and the lines in the spectra of its elements, based on a mathematical theory, which, however, does not seem yet to have been published, has been enunciated by Grunwald, who states (*P M* (6) 84, 354) —

'Let a be a primary chemical element, which is chemically combined with other elements in a gaseous substance A , and occupies the volume $[a]$ in the unit volume of A . Let the substance A combine chemically with another gaseous substance B , to form a third C . In this combination let the element a pass into a different chemical condition a' , giving up (or in exceptional circumstances taking up) a certain quantity of heat in order to permit the new compound to form, and in consequence chemically contracting (or exceptionally expanding). Let the volume which it occupies in the body C , after the new condition of chemical equilibrium has been established, be $[a']$, then the quotient $[a]/[a']$ is generally a very simple rational number in accordance with a known fundamental law of chemistry. If this is the case the wave-lengths λ of all the rays which belong to the element a in the line-spectrum of the free substance A , and are therefore radiated by it are related to the wave-lengths λ' of the corresponding rays which the same element emits in the new chemical condition a' , in which it exists in the more complex substance A within the newly formed compound C , as the corresponding volumes $[a]$ and $[a']$ '

This relation has been tested in the case of water vapour, as however both this substance and hydrogen have a good many lines in their spectra, it is not surprising that coincidences occur between the observed and calculated values of the wave lengths of the lines in the water vapour spectrum. We must, therefore, suspend our opinion as to the value of the relation given by Grunwald until the theoretical grounds on which it is based have been published. See also Ames (*N* 40, 19)

In a binary compound, AB , we may suppose that the atoms A and B are dynamical systems, which in the molecule of the compound are near together, and that the proximity of A causes the periods of B to be slightly different from the periods when B is vibrating by the influence of other systems, and *vice versa*. Then the theory of the vibrations of such a system shows that if p_1, p_2, \dots, p_n are the frequencies of A when free, q_1, \dots, q_n those of B , $\delta p_1, \delta p_n, \delta q_1, \delta q_n$, the increase in the frequencies p_i, \dots , respectively, due to the proximity of the two systems, then

$$p_1 \delta p_1 = \frac{f_{11}^2}{p_1^2 - q_1^2} + \frac{f_{12}^2}{p_1^2 - q_2^2} + \dots$$

With similar expression for δq_1 &c. The quantities f_{11} are quantities depending on the proximity of the systems

From this relation we see that the effect on the period, say p_1 , of the first system of the existence of a period, say q_2 , in the second, is to quicken the period of the first, if the first is quicker than the second, and to retard it, if it is slower than the second. If we observe the spectra, this result could be expressed by saying that the effect of the annexation was to make the line of B repel the lines of A . Thus on this theory the spectrum of the compound may be got by superposing the spectra of its constituents, A and B , and then supposing the lines of A to repel those of B , and the lines of B to repel those of A , the repulsion increasing with the proximity of the lines. Thus if we take two elements A and B , such that A and B have two lines nearly coinci-

dent, then in the compound AB these lines will be considerably displaced and the distance between them increased

Arrangement of the atoms in the molecule on the supposition that the atoms are vortex rings
There is one theory of the structure of the molecule which is worth mentioning, as it affords a possibility of the explanation of that remarkable alternation of properties with atomic weight which is expressed by the periodic law. If we assume that a molecule is built up of a number of vortex rings placed close together, then a section of the molecule, by a plane through the centre at right angles to the planes of the ring, will consist of two groups, each consisting of a number of small circles. The arrangement of the circles in either group will be very much the same as the arrangement, when in steady motion, of the cross sections of the same number of uniform straight parallel vortex columns with circular sections. These, when in steady motion, arrange themselves in a definite way, which may easily be discovered without calculation, as the arrangement is very nearly the same as that of the same number of equal uniform parallel magnets under the attraction of a magnetic pole, some distance away from the nearer poles of the parallel magnets, and of opposite sign to these poles. These magnets will take up definite positions of equilibrium, under the action of their mutual repulsion and the external attraction. The figures of equilibrium of the magnet are given by Mayer (*N* 18, 253) and Monckman (*Proc Camb Phil Soc* 6, 169). If we examine these figures, we see that as the number of magnets increase there is a tendency for certain peculiarities to recur, as, for example, the number of planes of symmetry, and the nature of the simpler groups of which we may imagine the more complex ones to be made up. Thus, if we imagine the molecules of all elements to be made up of the same primordial atom, and interpret increasing atomic weight to indicate an increase in the number of such atoms, then, on this view, as the number of atoms is continually increased, certain peculiarities in the structure will recur, which in all likelihood would be accompanied by a recurrence of some of the properties of the elements

Electrical theory of molecular structure — There is another view of molecular structure which is almost forced upon us by the laws of electrolysis, this is, that the forces between the atoms in the molecule are electrical in their origin. On this theory, the atoms in the molecule of a compound are supposed to be charged with definite quantities of electricity, the quantity of electricity on the atom being the same for all elements of the same valency, and being positive or negative, according as the element is electro positive or electro negative. The charge on an atom of a divalent element is assumed to be twice, and that on an atom of a trivalent element three times, the charge on the atom of a monovalent element (*v* Von Helmholtz [*Fara day Lecture*], *C J* 39, 277). This view of the structure of the molecule at once explains Faraday's law of electro chemical decomposition. It also explains the difference which exists between the electrical properties of the molecule and the atom, for in the molecule the positive and ne-

gative charges neutralise each other's effect at points outside the molecule, the free atom is, however, essentially charged and therefore capable of producing electrical effects. When we dissociate a gas into atoms, the dissociated gas, on this theory, consists of an equal number of electrically charged particles, some being charged with positive electricity, and an equal number (if the constituents of the molecule are of the same valency) charged with negative electricity. This collection of electrified particles would behave like a conductor of electricity, so that, if this theory of the structure of the molecule is correct, a gas whose molecules are dissociated by heat into atoms ought to be a conductor of electricity. J J Thomson (*P M* [6] 29, 358, 441) has recently made a series of experiments on the conduction of electricity through very hot gases, and has found that while some of these hot gases (hydriodic acid gas, for example) allow electricity to pass through them with ease, others (such as nitrogen) only allow it to do so with great difficulty, and it was found that whenever the electricity passed with ease through a hot gas, the dissociation of the gas could be detected by chemical means. These experiments are, therefore, in accordance with the result of this theory of molecular structure. On this view of molecular structure the 'bonds of affinity' of chemists have a distinct physical meaning, as they are the tubes of electrostatic force connecting the atoms.

A difficulty which arises on this theory, and one that seems to show that it requires modification, is the existence at low temperatures of what are called by chemists unsaturated compounds. For, according to this view of the structure of matter, an unsaturated compound is one in which there are not equal and opposite quantities of electricity in each molecule, so that the molecules of an unsaturated gas, being electrically equivalent to a number of positively and negatively charged particles, ought to be have like a conductor. But gases which are unsaturated at low temperatures, such as NO, behave at these temperatures with respect to electricity like saturated gases, they transmit electric induction. For example, a gold leaf electroscope will work perfectly well inside a glass vessel containing NO, and its leaves will be attracted by an electrified body outside the electroscope, and a current of electricity cannot be driven through a tube containing such gases by a battery containing only a small number of cells. We must therefore conclude that electrically such gases are saturated.

Maxwell, in the article on the 'Constitution of Bodies,' *Encyclopædia Britannica*, introduced the idea that in solids the molecules might arrange themselves in groups, some of which under the action of stresses might split up and form other stable groups in which the molecules are differently arranged, these new groups returning only slowly to their original configuration after the stresses are removed. This behaviour of the molecular groups shows itself in the 'elastic after effect' produced by torsion in metal wires and glass fibres. Ewing (*P M* [6] 30, 205) has lately applied the same idea to explain the behaviour of iron when magnetised, and has devised a model which illustrates very clearly

the breaking up of the old groups and the formation of new ones J J T

MOLECULAR WEIGHTS The article **ATOMIC AND MOLECULAR WEIGHTS**, in vol 1, describes the limits within which the term molecular weight may be applied with safety to solid and liquid bodies (*v* especially pp 347-350). Since that article was printed, an advance has been made in the methods by which the molecular weights of bodies which cannot be gasified without decomposition may be determined. This advance is based, for the most part, on the researches of Raoult. As the result of a long series of investigations into the lowering of the freezing point of water and various other solvents, produced by dissolving therein quantities of various compounds proportional to the formula-weights, or reacting weights, of these compounds, Raoult finds that such quantities of chemically similar compounds generally produce equal lowerings of the freezing points of water and some other solvents (Raoult's Memoirs will be found in *A Ch*, *v* especially [6] 8, 817).

Let P grams of a compound be dissolved in 100 g of water or other solvent, and let the observed lowering of freezing point of the solvent be C , then $\frac{C}{P}$ is called by Raoult the *coefficient of lowering of freezing point* for the compound in question. Putting M as the reacting-weight, or formula weight, of the compound, then $\frac{C}{P}M$ is called the *molecular lowering of freezing point* for this compound.

Raoult finds that $\frac{C}{P}M$ is generally constant for all the members of a series of chemically similar compounds. Thus, Raoult gives the following values for $\frac{C}{P}M$, water being the solvent —

- 19, for many organic compounds,
- 35, for salts of monovalent metals with monobasic acids,
- 40, for normal salts of monovalent metals with dibasic acids.

Raoult also gives the following values for $\frac{C}{P}M$, benzene being the solvent —

- 49, for many organic compounds,
- 25, for the lower members of homologous series of alcohols.

Other values for $\frac{C}{P}M$ were found when other solvents, *e g* acetic acid, were used.

Another form in which Raoult's results may be put is the following $-\frac{P}{C}$ = grams of dissolved body, in 100 g solvent, that lower freezing point of solvent 1°. Now $\frac{P}{C} \times c = M$, where c = constant determined experimentally for each series of chemically similar compounds, and for each solvent.

If the value of $\frac{C}{P}M$ is known for a group of compounds, or if the value of c is known in the

expression $\frac{P}{C} \times c = M$, it is possible to find the formula-weight of a member of this group. Thus, in the case of ether, the following data were obtained —

(i) 4.47 g ether were dissolved in 100 g water, the freezing point of the water was lowered by 1.05° , hence $\frac{C}{P} = \frac{1.05}{4.47} = 23^\circ$. But the

value of $\frac{C}{P}M$ for organic compounds generally dissolved in water is 19, hence, for ether, $M = \frac{19}{23} = 82$

(ii) 2.721 g ether were dissolved in 100 g benzene, the freezing point of the benzene was lowered by 1.826° , hence $\frac{P}{C} = 1.49$, $\therefore c = 1.49$ g ether in 100 g benzene lower the freezing point by 1° . But the constant for such organic compounds as the ethers dissolved in benzene is 49, hence, for ether, $M = 1.49 \times 49 = 73$

(iii) The value found for $\frac{C}{P}$ when ether was dissolved in acetic acid was 529° . But the value of $\frac{C}{P}M$ for organic compounds generally dissolved in acetic acid is 39, hence, for ether, $M = \frac{39}{529} = 74$

The mean of these three results gives 76.6 for the formula weight of ether, the molecular weight of ether gas, determined by applying Avogadro's law, is 74.

The empirical law of Raoult—quantities of chemically similar compounds proportional to the reacting weights, or formula weights, of these compounds produce equal lowerings of the freezing points of water and some other solvents—has been developed by van't Hoff, Arrhenius, and others (especially *Z P C* 1, 481 [translation in *P M*, August 1888], and *Z P C* 1, 631, 2, 284, 491). If an aqueous solution of a substance is contained in a vessel the walls of which are permeable by water molecules but not by the molecules of the dissolved substance, and the vessel is immersed in water, water will enter the vessel, and the pressure on the walls will increase until equilibrium results, after which no more water will enter. The pressure on the walls of the vessel is called *osmotic pressure*. If the vessel were furnished with a movable piston, the same condition of equilibrium might be attained, without the entry of water into the vessel, by compressing the solution with a pressure equal to the osmotic pressure. With such an arrangement the concentration of the liquid could be altered by increasing or decreasing pressure by means of the piston, as the process is reversible, the second law of thermodynamics may be applied.

The experiments of de Vries (*Z P C* 2, 415, 8, 103), Pfeffer (*Osmotische Untersuchungen* [Leipzig, 1887]), and others, show that the osmotic pressures of dilute aqueous solutions are proportional to the concentrations of these solutions. Now, to say that change of concentration of dilute solutions is proportional to the pressure exerted by the solutions, is equivalent

to saying that Boyle's law holds good for dilute solutions. Moreover, the proportionality of concentration to osmotic pressure may be deduced theoretically. If we assume, as seems justifiable, that osmotic pressure is due to the impact of the molecules of the dissolved substance, then the number of impacts in unit time must be proportional to the number of molecules in unit volume (on this point v. L. Meyer, *Z P C* 5, 23, and van't Hoff's reply, *Z P C* 5, 174, cf. Bredig, *Z P C* 4, 444). But this is the molecular conception of gaseous pressure. Hence, as in gases volume is inversely as pressure, the same proportionality should hold good in dilute aqueous solutions, in other words, Boyle's law should apply to these solutions. Van't Hoff then proceeds to deduce, by thermodynamical reasoning, that osmotic pressure is proportional to absolute temperature, concentration being constant. This conclusion is equivalent to saying that the law of Charles holds good for dilute aqueous solutions, inasmuch as concentration of solution corresponds with gaseous volume.

The experimental results of Pfeffer and of Soret (*A Ch* [5] 22, 293) are in keeping, on the whole, with the statement that the laws of Boyle and Charles hold good in dilute aqueous solutions.

Solutions which exert equal osmotic pressures are called *isotonic* solutions. Thermodynamical reasoning applied to these solutions leads to the conclusion that the osmotic pressure of a specified mass of a gasifiable substance in dilute solution is the same as the pressure exerted by the same mass of the same substance existing as a gas at the same temperature. If, then, osmotic pressure may be substituted for gaseous pressure, Avogadro's law may be extended to substances in dilute solution. This extension of the law of Avogadro is thus stated by van't Hoff: '*Equal volumes of different solutions, at the same temperature and osmotic pressure, contain equal numbers of molecules, which numbers are the same as would be contained in equal volumes of gases at the same temperature and pressure*'. This is van't Hoff's law of osmotic pressure. Various proofs of the accuracy of this law have been given.

To apply this law, it is necessary to find the mass of a substance present in a solution which is *isotonic* with another solution containing a known mass of a body of known molecular weight. The two solutions then contain equal numbers of molecules of the dissolved substances, and as the molecular weight of one substance is known, the molecular weight of the other can be found. There are many practical difficulties in determining whether or not two solutions are isotonic. Now van't Hoff has shown by thermodynamical reasoning (*Z P C* 1, 496) that solutions of different bodies in the same solvent, having equal freezing points, are isotonic at their freezing points. Put into other words, this conclusion asserts that solutions which have equal freezing points contain equal numbers of molecules in equal volumes, or, that solutions which contain equal numbers of molecules in equal volumes, and are therefore isotonic, have equal freezing points. But this is the law of lowering of freezing points am-

prically established by Raoult. In place of the somewhat vague term reacting weight, or formula weight, used in stating the law (*v supra*), we are now justified in employing the more definite term molecular weight. This law, as developed by van't Hoff, asserts that the product of the lowering of the freezing point of a 1 p.c. solution into the molecular weight of the dissolved body is a constant for different bodies dissolved in the same solvent. For solutions of most organic compounds in water, the constant is about 18.9.

Van't Hoff then proceeds to show that the molecular lowering of freezing point of a dilute solution bears a simple relation to the latent heat of fusion of the solvent (for proof *v Z P C* 1, 496-7). This relation is expressed, for very dilute solutions, by the equation

$$t = 0.1976 \frac{T^2}{W}$$

where t = molecular lowering of freezing point, already expressed as $\frac{C}{P}M$ (p. 417), T = freezing

point of solvent stated in absolute measure, and W = latent heat of fusion of solvent in gram-units (cf. Eykman, *Z P C* 3, 203, and especially *ib Z P C* 4, 512). If this conclusion is granted, it follows that equal numbers of molecules of all bodies dissolved in the same solvent must lower the freezing point to the same extent, provided the solutions are very dilute.

Let us take a case to exhibit the application of the law of molecular lowering of freezing point in the form given to it by van't Hoff. Thymol is dissolved in phenol, and the lowering of the freezing point of the phenol is observed. The constant for molecular lowering of freezing point of phenol is first calculated by van't Hoff's formula: the freezing point of the phenol used was 38° , in absolute measure this is $273^\circ + 38^\circ = 311^\circ$, the latent heat of fusion of the phenol was found to be 25, hence

$$t = 0.197 \frac{311^2}{25} = 76 \quad \text{The solution of thymol used}$$

contained 401 g thymol in 7559 g phenol, the freezing point was lowered by 2.49° , stating these results in parts of thymol per 100 of phenol, we find that 5.3 g thymol dissolved in 100 g phenol lowered the freezing point through 2.49° . Then $\frac{5.3}{2.49} = 2.12$, i.e. 2.12 g thymol in

100 g phenol lowered the freezing point through 1° . But this quantity, 2.12, is $\frac{1}{16}$ th of the molecular weight of thymol, therefore molecular weight of thymol = $2.12 \times 76 = 161$. The molecular weight calculated from the formula $C_{11}H_{19}O$ is 150.

In applying the law of molecular lowering of freezing point it is necessary to work with dilute solutions. The freezing point varies somewhat with concentration, in some cases this variation is very marked. Beckmann (*Z P C* 2, 742) recommends that a series of observations should be made, concentration varying so that the lowering of freezing point may range from $c. 2^\circ$ to $c. 2^\circ$. If possible, observations should be made with solutions in different solvents, care being taken to select solvents which do not react chemically, so far as is known, with the dissolved body, and the results should be checked by ob-

servations of the lowering of vapour pressure of some solvent produced by dissolving in it the substance whose molecular weight is being determined (*v infra*).

To sum up this part of the subject. Known weights of the substance, the molecular weight of which is to be determined, are dissolved in known weights of the solvent, so that the concentration of the solutions varies from, say, 1 to 5 or 6 p.c., the freezing point of each solution is determined. The freezing point of the solvent is determined. Two methods of calculation may then be adopted—

(i) The lowering of freezing point, brought about by 1 g of the substance dissolved in 100 g of solvent, is calculated from each observation made, let this = A . The value of the constant expressing the molecular lowering of freezing point of the solvent by the class of bodies to which the substance under examination belongs is known,

let this be C . Then $\frac{C}{A}$ gives approximately the

molecular weight of the substance.

(ii) The weight of the substance which would lower the freezing point of the solvent by 1° is calculated from each observation made, let this be B . Then $B \times C$ gives approximately the molecular weight (C has the same meaning as in (i)).

The values found for mol. w. from the different observations are compared, if the differences are small, the mean is taken, if there are marked differences, experiments are made with other solvents. It may be that the substance undergoes dissociation in all solvents, and that, therefore, the method is inapplicable (*v infra*).

The molecular lowering of freezing point of the solvent may be calculated by the use of van't Hoff's formula ($t = 0.1976 \frac{T^2}{W}$), provided the

latent heat of fusion of the solvent is known, the value thus found should agree with the constant determined by experiment.

The solvents commonly employed are water, benzene, phenol, and glacial acetic acid, naphthalene has also been used for some hydrocarbons, Eykman (*Z P C* 4, 512) recommends urethane, phenyl propionic acid, the higher acids of the acetic series, stearin, and *p*-toluidine. Various modifications of Raoult's original apparatus have been devised, references to the papers in which the most important are described will be found at the end of this article.

There are many apparent exceptions to Raoult's law, and therefore to van't Hoff's law of osmotic pressure. These exceptions are explained by the hypothesis, put into definite form by Arrhenius (*Z P C* 1, 631), that bodies whose behaviour is not directly expressed by these laws are partially dissociated in solution. Here again there is a marked analogy between gases and dilute solutions, as the pressure of the vapour obtained by heating ammonium chloride is greater than the pressure calculated by Avogadro's law, on the assumption that the vapour consists of molecules of NH_4Cl , but as the observed pressure agrees with the calculated pressure when it is assumed that the vapour consists of equal numbers of molecules of NH_3 and HCl , so the apparently abnormal osmotic pres-

tures of many solutions are brought into agreement with van't Hoff's law by assuming that the bodies in these solutions are more or less dissociated into simpler molecules. If the molecular weight of a substance, as determined by observations of freezing points, apparently decreases as dilution increases, it is likely that the substance is dissociated in the solution, and that the amount of dissociation increases as the quantity of solvent is increased. Substances may undergo dissociation in one solvent and not in another (for several carefully worked out examples, *v* Beckmann, *Z P C* 2, 715). Those substances, solutions of which show osmotic pressures agreeing with those calculated by van't Hoff's law from observations of the lowering of freezing-points of the solutions, are generally, if not always, non electrolytes. The apparent exceptions to the law of van't Hoff occur chiefly, if not wholly, among electrolytes. The dissociation-hypothesis of Arrhenius regards such electrolytes as more or less dissociated into their ions when they are dissolved in water. A great deal of work has been done in developing and applying the hypothesis of electrolytic dissociation, an account of this work, and of the chief results, will be found in the article *PHYSICAL METHODS*, section *Electrical methods*.

Raoult has found that in many cases the lowerings of vapour pressure produced by dissolving a comparatively non volatile substance in a considerably more volatile liquid is related to the number of molecules of the dissolved body in 100 molecules of the solvent, provided the solution be very dilute (*Z P C* 2, 353). Arrhenius has shown that this generalisation made by Raoult can be deduced from van't Hoff's law of osmotic pressures (*Z P C* 3, 115). The generalisation has been placed on a thermodynamical basis by van't Hoff (*Z P C* 1, 493) and Planck (*Z P C* 1, 577). The law may be put in the following form *At any specified temperature the ratio of lowering of vapour-pressure of a solvent, produced by dissolving a non-volatile body in it, to the vapour-pressure of the solvent, is equal to the ratio of the number of molecules of the dissolved body to the total number of molecules in the solution*. Let p = vapour pressure of solvent, $p-p'$ = observed lowering of vapour pressure, W = weight of dissolved body, W' = weight of solvent, m = molecular weight of dissolved body, and m' = molecular weight of solvent; then

$$\frac{p-p'}{p} = \frac{W}{m} \cdot \frac{W'}{m'}.$$

If W , W' , p , p' , and m' are known, m can be found. Suppose a grams of the substance are dissolved in 100 g solvent, then,

$$m = \frac{m' a}{100 p - p'}$$

Among the solvents which have been used in applying this law to the determination of molecular weights are ether, alcohol, CS_2 , and Hg. For descriptions of apparatus, and some of the data obtained, *v* Raoult (*Z P C* 2, 353), Walker (*Z P C* 2, 602), Beckmann (*Z P C* 4, 532), Ramsay (who determined mol w of several metals by using Hg as solvent) (*C J* 55, 521).

Raoult and others have found that the rise

in the boiling point of a liquid caused by dissolving a non-volatile substance therein bears a simple relation to the molecular weight of the dissolved substance. There is a constant which must be determined for each solvent which expresses the rise of b.p. caused by solution of 1 gram molecule of substance in 100 grams of solvent, for ether this constant is 21° (Beckmann, *Z P C* 3, 803). It is necessary then to find the weight of substance which, dissolved in 100 g ether, raises the b.p. 1° , the product of this weight multiplied by 21 gives, approximately, the molecular weight of the dissolved substance. Thus Beckmann (*l.c.*) found that 2.153 g aniline dissolved in 100 g ether raised the b.p. of the ether 484° , therefore $4.45 \text{ g aniline would raise the b.p. } 1^\circ$, but $4.45 \times 21 = 93.4$, the mol w of aniline is 93. Arrhenius contributes a note to Beckmann's paper (*Z P C* 4, 550) in which he shows, by thermodynamical reasoning, that the rise of boiling point (dT) of a solvent, caused by dissolving n gram molecules of another substance in 100 g of the solvent, is directly proportional to the quantity dissolved (n) and the square of the b.p., and is indirectly proportional to the heat of vaporisation of 1 g of the liquid, W . Put into an equation this state-

ment reads thus $-dT = \frac{92T^2}{W} n$. This equation

is precisely similar to that found by van't Hoff (*v* p 419) for the molecular lowering of the freezing point of a solvent, only in the present case the quantities T and W depend upon the boiling point of the solvent. A description of apparatus suited for applying the law of molecular raising of boiling point to molecular weight determinations will be found in Beckmann's paper, *Z P C* 4, 543.

References—Blagden, *T* 58, 277, de Coppet, *A Ch* [4] 23, 366, 25, 502, 26, 98, Rudorff, *P* 114, 63, 116, 55, 145, 599, Raoult, *A Ch* [5] 20, 217, 28, 133, [6] 2, 66, 93, 99, 115, 4, 401, 7, 289, 317, *Z P C* 2, 488, Eykman, *Z P C* 4, 497 (data bearing on question of series constants, influence of concentration, nature of solvent, &c).

The following memoirs bear chiefly on the similarities between the gaseous state and that of substances in dilute solutions—van't Hoff, *Z P C* 1, 481, 5, 175, 221, Arrhenius, *ibid* 1, 631, 2, 284, 491, Ostwald, *ibid* 2, 36, 270, Beckmann, *ibid* 2, 734, Nernst, *ibid* 2, 613, 4, 872, Planck, *ibid* 1, 577, 2, 343, Bredig, *ibid* 4, 444, Wiedemann, *ibid* 2, 241, and Ostwald's reply p 243, Pickering, *C J* 57, 331.

The following memoirs are on the application of van't Hoff's law and Raoult's method to special compounds—von Klobukow, *Z P C* 3, 351, 476 (shows that CHI_3 and morphine do not give the anomalous results which Raoult supposed they did), Eykman, *Z P C* 2, 966 (morphine), Paterno a Nasim, *B* 21, 2153 (I), Loeb, *Z P C* 2, 606 (I), Beckmann, *ibid* 5, 76 (I, P, S), Paterno, *ibid* 4, 457, Sabanejeff, *J R* 1889 [1] 515, abstract in *B* 23, [Ref] 87 (colloids); Brown a Morris, *C J* 53, 610, 55, 462 (carbohydrates), Löw, *B* 21, 271, 22, 470 (sugars), von Klobukow, *Z P C* 5, 28 (sugars), Heycock a Neville, *C J* 55, 666, 57, 376, Kuster, *Z P C* 5, 601 (isomorphous mixtures).

The following memoirs deal with relations between osmotic pressures, lowering of vapour-pressure and of freezing-point, and raising of boiling point, of a solvent by substances dissolved therein — Arrhenius, *Z P C* 8, 115, van't Hoff, *ibid* 1, 481, Planck, *ibid* 1, 577, van't Hoff a Reicher, *ibid* 3, 198, Guldberg, *C R* 70, 1349, Beckmann, *Z P C* 4, 532, Raoult, *C R* 110, 402, Raoult a Recoura, *C R* 110, 402

The following memoirs contain descriptions of apparatus — Auwers, *B* 21, 701, Hollemann, *B* 21, 860, Hentschel, *Z P C* 2, 306, Beckmann, *ibid* 2, 638, 4, 543 (raising b p), Eykman, *ibid* 2, 964, 3, 113, 4, 497, Fabinyi, *ibid* 3, 38, von Klobukow, *ibid* 4, 10, Raoult, *ibid* 2, 353 (lowering vapour pressures), Walker, *ibid* 2, 602 (lowering vapour pressures), Heycock a Neville, *C J* 55, 666 (using Hg as solvent), Ramsay, *C J* 55, 521 (lowering vapour-pressure of Hg) M M P M

MOLYBDATES Salts of molybdic acid v.

MOLYBDENUM, ACIDS OF, p 422

MOLYBDENUM Mo At. w 95.9 Mol w unknown SG 8.56 (Long, *Am. S* [2] 45, 131), 8.49 to 8.64 (Bucholz, *Nicholson's J* 20, 121) Does not melt in O-H flame at temp at which Pt melts (Debray, *A* 103, 250) SH 5°–15° 0659 (De la Rive a Marcet, *A Ch* [2] 75, 113) Emission spectrum (obtained by using electric spark) shows only a few lines, the most prominent in the blue and violet (v Thalen, *A Ch* [4] 18, 242) SVS c 113

Occurrence — Never uncombined The sulphide (molybdenite, or molybdenum glance), oxide (molybdenum ochre), molybdate of lead (wulfenite), and molybdate of cobalt (paterite), occur in small quantities in certain localities Many iron ores contain small quantities of Mo compounds (Braun, *Fr* 6, 86, Wohler, *Mineral analyse* [Göttingen, 1861]) The residues from the copper smelting ovens at the Mansfeld works sometimes contain as much as 28 p c Mo (Heine, *J pr* 9, 176, cf Stromeyer, *P* 28, 551, Steinberg, *J pr* 18, 379, Genth *J pr* 37, 193, Steinacker, *Ueber einige Molybdunverbindungen* [Göttingen, 1861] 22) An alloy of Mo and Pb, in plates 30 mm long, was found in Utah by Silliman (*Am S* [3] 6, 128) According to Lookyer (*Pr* 27, 279), Mo occurs in the sun

Scheele in 1778 (*Opusc* 1, 200) distinguished native Mo sulphide from galena In 1782, Hjehn obtained the metal from the sulphide (v *Crell Ann* for 1790, 1791, 1792, and 1794) For an historical account of this metal v Svanberg a Struve, *J pr* 44, 257 The name molybdenum was given from *μολύβδαινα*, or *molybdæna*, used by Dioscorides and Pliny to designate galena and other lead compounds

Formation — 1 By reducing MoO₃ by H, K or Na, C, or KCN — 2 By reducing Mo chlorides by H at a high temperature — 3 By reducing acid K molybdate by C — 4 By electrolysing molten MoO₃, or (NH₄)₂MoO₄ in solution

Preparation — The starting-point is pure MoO₃ (v *infra*) — 1 MoO₃ is heated in a crucible in a stream of pure H (which must be passed over a long layer of red-hot Cu), the mass is transferred to a tube of unglazed porcelain, and very strongly heated in the H stream, finally the last traces of oxide are removed by gently

warming in a stream of dry HCl as long as a white wool like sublimate (MoO₃, 2HCl) is formed (Liechti a Kempe, *A* 169, 344) The metal in contact with the porcelain tube is not pure (Debray, *C R* 56, 732, cf Wohler a von Uslar, *A* 94, 256, Rammelsberg, *P* 127, 284) — 2 MoO₃ is mixed with C and heated in a graphite-crucible, in the O-H flame, the product always contains a few per cents of C (Debray, *C R* 46, 1098) — 3 An intimate mixture of 1 pt MoO₃ and 1½ pts KCN is placed in a crucible, the cover is luted on, the crucible is placed in another which is filled with C, and the whole is kept at white heat for 12 hours By this method Loughlin (*Am S* [2] 45, 131) obtained metal with 98.7 p c Mo

There are many methods for preparing MoO₃ from Mo ores Finely powdered molybdenite (MoS₂) is roasted in an open porcelain vessel, with frequent stirring, until SO₂ ceases to be evolved, Brunner (*D P J* 150, 672) recommends to mix the ore with its own weight of fine quartz-sand, previously washed with HClAq, and to heat in a flat Pt dish, to incipient redness, till the residue is citron yellow when hot, and white when cold The roasted ore is treated with NH₄Aq, to the solution are added a few drops of NH₄ sulphide, the ppd CuS is removed, the filtrate is evaporated to dryness, the residue is dissolved in NH₄Aq, and the solution is evaporated to the crystallisation point (Wohler) Svanberg a Struve (*J pr* 44, 264) add excess of K₂CO₃ to the ammoniacal solution of the roasted ore, separate ppd Al₂O₃, evaporate to dryness, and strongly heat the residue in a Pt crucible, they treat with water, filter from CuO and Al₂O₃, evaporate to dryness, add S equal to double the weight of the residue, heat in a glass vessel on a sand bath to full redness till excess of S is burnt off, wash with warm water (the last washings containing a little K₂CO₃) as long as the washings are coloured, and until the MoS₂ is perfectly black, and then roast the pure MoS₂ thus obtained, or oxidise it to MoO₃ by HNO₃Aq Delifs (*A* 106, 376) decomposes molybdenite by HNO₃Aq Wittstein forms soluble sulpho-molybdates by roasting with S, and decomposes by H SO₄Aq, finally roasting the sulphide to MoO₃ (*R P* [2] 73, 155, cf Wicke, *A* 45, 373, Wohler, *ibid* p 374) Molybdenite may also be decomposed by calcining with alkali carbonates (v Christl, *D P J* 124, 398, Elbers, *A* 83, 219), or by the use of H₂SO₄ (Elbers, *lc*)

Properties — As prepared by reducing the chloride in H, Mo appears as dull silver-coloured, somewhat malleable, plates (Wohler a von Uslar, *A* 94, 256) The metal obtained by reducing MoO₃ by H at very high temperatures is lustrous, if the reduction is effected at lower temperatures, an ashen grey powder is obtained which becomes metal-like when rubbed The metal obtained by Debray, which contained 4–5 p c C, was silver white, and harder than topaz The SG of Mo is given by Loughlin (*Am S* [2] 45, 131) as 8.56, this metal contained 98.7 p c Mo Debray (*C R* 46, 1098) gives SG 8.6 for the metal obtained by reducing MoO₃ by C, and containing 4–5 p c C

Mo, whether in powder or in pieces, is unchanged in ordinary air, when heated it becomes brown, then blue, then white, on the surface,

and at high temperatures it is burnt to MoO_3 . It is oxidised by fusion with KNO_3 , or slowly by KOH , also by HNO_3 , or Cl water, it burns when heated in steam, evolving H_2 . Mo is insol in HClAq , dilute $\text{H}_2\text{SO}_4\text{Aq}$, and HFAq , it is not acted on by KOHAq . Mo combines directly with O , Cl , and Br , but not with I . The metal is infusible at white heat (Debray, *A* 108, 250).

The atomic weight of Mo has been determined (1) by analysing, and determining $V D$ of MoCl_3 (Liechti & Kempe, *A* 169, 344, Debray, *C R* 66, 732), (2) by determining $S H$ of Mo (De la Rive & Marcet, *A Ch* [2] 75, 113), (3) by determining Cl in MoO_3Cl_2 (H Rose, *P* 40, 400, Svanberg & Struve, *A* 68, 209), (4) by synthesis of Ag_2MoO_4 (Debray, *C R* 66, 732), (5) by analyses of MoCl_3 and MoCl_4 (L & K, *A* 169, 344, cf L Meyer, *ibid* p 360), (6) by reducing MoO_3 in H (*S* & *S*, *J pr* 44, 301, Dumas, *A Ch* [3] 55, 143, Debray, *C R* 66, 732, Rammelsberg, *B* 10, 1776), (7) by oxidising MoS_2 to MoO_3 (*S* & *S*, *A* 68, 209), (8) by conversion of MoCl_3 and MoCl_4 to MoS_2 (L & K, *A* 169, 344, cf Meyer, *ibid* p 360). The atom of Mo is pentavalent in the gaseous molecule MoCl_5 .

Mo is probably both metallic and non-metallic in its chemical relations. Few, if any, definite salts are known obtained by replacing the H of acids by Mo , but Mo_2O_3 and MoO_2 seem to dissolve in acids without evolution of O . The oxide MoO_3 is an anhydride, molybdic acid, H_2MoO_4 , and molybdates, besides tri tetra n molybdates, are known. MoO_3 also combines with some other anhydrides to form complex bodies which react as acids, *e g* P_2O_5 , 20MoO_3 , $38\text{H}_2\text{O}$, SiO_2 , 12MoO_3 , $26\text{H}_2\text{O}$. MoS_2 reacts as an acidic sulphide, forming sulphosalts M^+MoS_4 . Several haloid compounds, and many oxyhaloid compounds, of Mo are known. Mo is closely related to Cr , W , and U , and is less closely related to S , Se , and Te . Mo occurs in Group VI series 6, the following table exhibits the position of Mo relatively to the other members of the group

Even series—				
2	4	6	8	10
$\text{O} = 16$	$\text{Cr} = 52$	$\text{Mo} = 96$	—	$\text{W} = 184$

Odd series—				
3	5	7	9	11
$\text{S} = 32$	$\text{Se} = 79$	$\text{Te} = 125$	—	—

(*v* CHROMIUM GROUP OF ELEMENTS, vol 11 p 168, cf CLASSIFICATION, vol 11 p 207)

Reactions—1 Heated in air or oxygen, Mo is burnt to oxide, if the temperature is sufficiently high, MoO_3 is formed—2 Oxidised, to MoO_3 , by molten nitre, and slowly by molten potash—3 Heated in steam, blue oxide ($?\text{Mo}_2\text{O}_3$), and then MoO_3 , is formed—4 Pulverulent Mo is oxidised by rubbing with silver oxide, or mercuric oxide—5 Oxidised by nitric acid, conc hot sulphuric acid, or chlorine water—6 Mo does not dissolve in hydrochloric, hydrofluoric, nor dilute sulphuric acid, nor does it react with potash solution.

Combinations—1 Mo combines with chlorine to form MoCl_3 , and with bromine to form MoBr_3 , it does not combine directly with I —2 Heated in oxygen, Mo forms oxides, MoO_3 being the final product of the combination.

Detection and Estimation—The greater

number of the Mo compounds, including most of the molybdates of the alkaline earths and heavy metals, are insol in water. Lower oxides, and also the sulphides, produce MoO_3 when heated in air, MoO_3 readily dissolves in alkali solutions. Insol molybdates are brought into soluble forms by fusion with alkali carbonates. All Mo compounds may be changed to soluble alkali sulphomolybdates by fusion with Na_2CO_3 , and S . Phosphomolybdates are readily soluble in NH_4Aq . Mo compounds impart a yellow colour to the borax bead when heated in the oxidising flame, the colour disappears on cooling, in the reducing flame, they give a dark brown colour, the bead is opaque if excess of Mo compound is present. They give a clear green microcosmic salt bead in the reducing flame, and in the oxidising flame a bead which is greenish when hot and colourless when cold. In solution, Mo compounds give a brown black pp with H_2S , soluble in NH_4 sulphide, solutions of molybdates give blue coloured compounds by the action of reducing agents. When MoO_3 or a molybdate is evaporated with conc H_2SO_4 , and then allowed to cool, the acid becomes deep blue in colour, this reaction distinguishes molybdates from tungstates, the blue colour does not appear if antimonious or stannous salts are present, unless the substance has been wetted with a few drops of phosphoric acid, and evaporated to dryness, before addition of H_2SO_4 (Schönn, *Fr* 8, 379, Maschke, *Fr* 12, 383). If Zn is put into solution of a molybdate, a few drops of conc KCNSAq are added, and then enough HClAq or $\text{H}_2\text{SO}_4\text{Aq}$ to cause slow evolution of H_2 , a carmine red colour is produced, even with $\frac{1}{50000}$ part MoO_3 , ether removes the coloured compound (Braun, *Fr* 2, 36).

Liquids containing molybdates yield deep red solutions (sulphomolybdates) by boiling with yellow NH_4 sulphide (Braun, *Fr* 6, 86).

Mo is estimated (1) as metal by reducing in H (Rammelsberg, *P* 127, 281), (2) as MoO_3 by oxidising in air or by HNO_3Aq (v d Pfordten, *B* 15, 1927), (3) as PbMoO_4 (Chatard, *B* 4, 280, v also Ullik, *A* 144, 217), (4) by reduction with Zn and titration by means of standard KMnO_4Aq (Wernecke, *Fr* 14, 1, v d Pfordten, *B* 15, 1927), (5) by the reaction of MoO_3 with an acidified solution of KI , whereby I is set free and is determined by standard $\text{Na}_2\text{S}_2\text{O}_4\text{Aq}$ (Mauro & Danesi, *Fr* 20, 507).

Molybdenum, Acids of, and their salts. The oxide MoO_3 is slightly sol in water, c 500 parts cold water dissolve 1 part MoO_3 . The solution reddens litmus. Several hydrates of MoO_3 have been isolated. $\text{MoO}_3 \cdot \text{H}_2\text{O} = \text{H}_2\text{MoO}_4$, was obtained by Ullik, and also by Vivier (*v infra*), the hydrates $\text{MoO}_3 \cdot 2\text{H}_2\text{O}$, $2\text{MoO}_3 \cdot \text{H}_2\text{O}$, $4\text{MoO}_3 \cdot \text{H}_2\text{O}$, $5\text{MoO}_3 \cdot \text{H}_2\text{O}$, and $8\text{MoO}_3 \cdot \text{H}_2\text{O}$, have also been prepared, some of these are soluble, and some are insoluble, in water. Several series of salts are known, these salts may be represented as derived from hydrates of MoO_3 , in some cases the hypothetical hydrates have been isolated, and in other cases they have not been isolated. Besides reacting as the anhydride of more than one acid, MoO_3 reacts with some oxides more acidic than itself to form compounds, *e g* $\text{MoO}_3 \cdot \text{SO}_3$, it also combines with

certain acids—*e.g.* with HCl , H_3PO_4 , H_2AsO_4 —the compounds thus formed have acidic reactions and yield salts (*v* ANTIMONO MOLYBDATES, ARSENO MOLYBDATES, PHOSPHO MOLYBDATES, p 425, *v* also MOLYBDENUM TRIOXIDE, p 432)

MOLYBDIC ACIDS 1 H_2MoO_4 . The normal hydrate $\text{MoO}_3 \cdot \text{H}_2\text{O}$, or hydroxide $\text{MoO}_3(\text{OH})_2$, was obtained by Ullik, but only once, by mixing MgMoO_4 with an equivalent of HNO_3 aq and allowing to stand (*A* 153, 374) Vivier (*C R* 106, 601) obtained this compound by allowing a solution of NH_4 molybdate in HNO_3 aq to stand for some time, also by heating ordinary NH_4 molybdate solution with its own volume of water to 50° – 60° for some days H_2MoO_4 forms a mass of minute needles, it is quite insol water

2 $\text{H}_2\text{Mo}_2\text{O}_{11}$. The hydrate $2\text{MoO}_3 \cdot \text{H}_2\text{O}$ ($=\text{MoO}_3(\text{OH})_2 \cdot \text{MoO}_3$) was obtained by Ullik (*A* 144, 329) by the action of excess of H_2SO_4 aq on the Ba salt formed by adding BaCl_2 aq to ordinary NH_4 molybdate solution, filtering, placing over H_2SO_4 and drying the gum like residue over H_2SO_4 for some months, or at 100° (Ullik, *A* 153, 373) $\text{H}_2\text{Mo}_2\text{O}_{11}$ is a gum like amorphous solid, *e sol* water

3 $\text{H}_4\text{Mo}_3\text{O}_{15}$, and $\text{H}_4\text{Mo}_4\text{O}_{16}$. The hydrates $4\text{MoO}_3 \cdot \text{H}_2\text{O}$ ($=\text{MoO}_3(\text{OH})_2 \cdot 3\text{MoO}_3$), and $8\text{MoO}_3 \cdot \text{H}_2\text{O}$ ($=\text{MoO}_3(\text{OH})_2 \cdot 7\text{MoO}_3$) are formed similarly to $2\text{MoO}_3 \cdot \text{H}_2\text{O}$, the drying being conducted at 120° , and 160° – 170° , respectively (*U*, *l c*) These compounds are amorphous solids, *e sol* water Sabanejeff (*J R* 1889 [1] 515, abstract in *B* 23 (*Ref*) 87) obtained a colloidal form of $\text{H}_4\text{Mo}_3\text{O}_{15}$, *sl sol* water, by drying for some weeks over H_2SO_4 , the formula was determined by applying Raoult's law (*cf* MOLECULAR WEIGHTS, p 417) For other hydrates of MoO_3 , viz $\text{MoO}_3 \cdot 2\text{H}_2\text{O}$ and $5\text{MoO}_3 \cdot \text{H}_2\text{O}$, *v* HYDRATES OF MOLYBDENUM TRIOXIDE, p 432

Graham (*C J* 1864) obtained a soluble colloidal form of molybdic acid (he does not give the composition) by dialysing an aqueous solution of Na molybdate, to which excess of HCl aq had been added, until the liquid in the dialyser was free from NaCl and HCl Graham describes the solution of molybdic acid as 'yellow, astringent to the taste, acid to test paper, and possessed of much stability', it decomposed Na_2CO_3 with evolution of CO_2 , and became in *sol* when heated for some time with a strong acid

Chloromolybdic acid $\text{H}_2\text{MoO}_4\text{Cl}_2$ ($=\text{MoO}_3(\text{OH})_2\text{Cl}_2$) (*Molybdenum hydroxychloride*) This compound is obtained by passing HCl over MoO_3 at 150° – 200° (Debray, *C R* 46, 1101) It forms a loose, white, crystalline, mass, *e sol* water, sublimed unchanged in HCl

Thiomolybdic acid and thiomolybdates, *v* MOLYBDENUM, THIOACIDS OR, AND THEIR SALTS, p 434

MOLYBDATES A great many molybdates are known, they are more or less closely related to the chromates The following classification is that given by Ullik (*W A B* 60 [2] 295), and now generally adopted $\text{R} = \text{divalent metal}$

Monomolybdates, RMO_4 , or RO MoO_4 ,
Dimolybdates, $\text{R}_2\text{Mo}_2\text{O}_{11}$, or $\text{R}_2\text{O } 2\text{MoO}_3$,
Trimolybdates, $\text{R}_3\text{Mo}_3\text{O}_{15}$, or $\text{R}_3\text{O } 3\text{MoO}_3$,
Tetramolybdates, $\text{R}_4\text{Mo}_4\text{O}_{19}$, or $\text{R}_4\text{O } 4\text{MoO}_3$,
Octomolybdates, $\text{R}_8\text{Mo}_8\text{O}_{27}$, or $\text{R}_8\text{O } 8\text{MoO}_3$,
Decamolybdates, $\text{R}_{10}\text{Mo}_{10}\text{O}_{31}$, or $\text{R}_{10}\text{O } 10\text{MoO}_3$,
Tribasic heptamolybdates, $\text{R}_3\text{Mo}_7\text{O}_{24}$, or $3\text{R}_2\text{O } 7\text{MoO}_3$,

There are a few molybdates which do not belong to any of these classes, *e.g.* $5\text{Al}_2\text{O}_3 \cdot 2\text{MoO}_3$, and $2\text{BaO } 5\text{MoO}_3$

The alkali *monomolybdates* are produced by dissolving MoO_3 , xH_2O , or MoO_3 , in equivalent quantities of alkali, in solution or molten, they are easily *sol* in water The other salts of this series are generally *insol* in water, they are produced by *ppn* from the alkali salts, or, in crystals by fusing Na_2MoO_4 with NaCl and metallic chlorides (Schultze, *A* 126, 55) The alkali *dimolybdates* are formed by fusing MoO_3 and alkali carbonates in the proper proportion, or by the action of MoO_3 on molten alkali nitrates (Ullik, *A* 144, 214) The *trimolybdates* are formed by boiling metallic carbonates with considerable excess of MoO_3 and water, filtering, and allowing to evaporate in the air These salts are generally easily *sol* in hot water, but only slightly *sol* in cold water, many of them exist both in crystalline and amorphous forms, the amorphous varieties are readily *sol* in cold water (Ullik, *A* 144, 227, 153, 376) The *tetramolybdates* may be obtained by rapidly evaporating very small quantities of the solutions which yield *trimolybdates* when allowed to evaporate slowly They form amorphous, brittle, masses, which slowly decompose with formation of *trimolybdates* (Ullik, *A* 144, 321) Crystallisable acid *tetramolybdates* are sometimes obtained by adding a fair excess of acid to solutions of mono or *dimolybdates*, addition of a little acid to such solutions generally throws down *trimolybdates* Very few *octo-* or *decamolybdates* have been prepared

The following table presents the composition of the chief molybdates The small figures after the formulæ refer to the list of memoirs given after this table —

MOLYBDATES

<i>Monomolybdates</i> , RMO_4 , or RO MoO_4	
$(\text{NH}_4)_2\text{MoO}_4$ (also <i>infra</i> , Mg salts) (23, 27)	
$5(\text{NH}_4)_2\text{MoO}_4 \cdot \text{Mn}_2\text{Mo}_2\text{O}_{21}$ 12aq (22)	
BaMoO_4 (1, 23)	
$\text{BeMoO}_4 \cdot \text{BeO} \cdot \text{H}_2$ 6aq (24)	
CaMoO_4 (7)	
CoMoO_4 (1, 89)	
$\text{CoMoO}_4 \cdot 2\text{NH}_3$ aq (33)	
$3\text{CuMoO}_4 \cdot \text{CuO} \cdot \text{H}_2$ 4aq (11)	
$\text{Fe}_2(\text{MoO}_4)_3 \cdot \text{MoO}_3$ 7aq (34, 1)	
$\text{Fe}_2(\text{MoO}_4)_3 \cdot 2\text{MoO}_3$ 16aq (20)	
$5\text{Li}_2\text{MoO}_4$ 2aq (17, 26)	
MgMoO_4 5aq (12, 21, 26)	
$\text{MgMoO}_4 \cdot (\text{NH}_4)_2\text{MoO}_4$ 2aq (12)	
$\text{MgMoO}_4 \cdot \text{K}_2\text{MoO}_4$ 2aq (12)	
MnMoO_4 (<i>v supra</i> , NH_4 salts, and <i>infra</i> , K salts) (1, 22, 39)	
Hg_2MoO_4 (29)	
$\text{NiMoO}_4 \cdot 2\text{NH}_3$ aq (1, 33)	
K_2MoO_4 (also <i>supra</i> , Mg salts) (23, 26, 34)	
$2\text{K}_2\text{MoO}_4 \cdot \text{Mn}_2\text{Mo}_2\text{O}_{21}$ 12aq (22)	
$\text{K}_2\text{MoO}_4 \cdot 2\text{Na}_2\text{MoO}_4$ 14aq (16, 26)	
$3\text{Rb}_2\text{MoO}_4 \cdot 4\text{MoO}_3$ 4aq (26, 31)	
Ag_2MoO_4 (33)	

derived from H_2MoO_4 , or $\text{H}_2\text{O MoO}_3$,
" " H_2MoO_4 , or $\text{H}_2\text{O } 2\text{MoO}_3$,
" " *hypothetical* $\text{H}_2\text{Mo}_2\text{O}_{11}$, or $\text{H}_2\text{O } 3\text{MoO}_3$,
" " $\text{H}_2\text{Mo}_3\text{O}_{15}$, or $\text{H}_2\text{O } 4\text{MoO}_3$,
" " $\text{H}_2\text{Mo}_4\text{O}_{19}$, or $\text{H}_2\text{O } 5\text{MoO}_3$,
" " *hypothetical* $\text{H}_2\text{Mo}_8\text{O}_{27}$, or $\text{H}_2\text{O } 10\text{MoO}_3$,
" " *hypothetical* $\text{H}_2\text{Mo}_7\text{O}_{24}$, or $3\text{H}_2\text{O } 7\text{MoO}_3$,

$\text{Ag}_2\text{MoO}_4 \cdot 4\text{NH}_3$ (35)
 $2\text{Ag}_2\text{MoO}_4 \cdot 3\text{MoO}_3$ (28)
 $\text{Na}_2\text{MoO}_4 \cdot 2\text{aq}$ (also *supra*, K salts). (28, 28, 36)
 SrMoO_4 (1)
 Ti_2MoO_7 (28, 37)
 $8\text{Ti}_2\text{MoO}_7 \cdot 5\text{MoO}_3$ (37)
 $8\text{Ti}_2\text{MoO}_7 \cdot 3\text{MoO}_3$ (37)
 ZnMoO_4 (1, 39)

Dimolybdates, $\text{R}_2\text{Mo}_2\text{O}_7$, or $\text{RO } 2\text{MoO}_3$, or $\text{R}_2\text{MoO}_4 \cdot \text{MoO}_3$

$\text{Al}_2(\text{MoO}_4)_3 \cdot 3(\text{NH}_4)_2\text{Mo}_2\text{O}_7 \cdot 20\text{aq}$ (18)
 $\text{Al}_2(\text{MoO}_4)_3 \cdot 3\text{K}_2\text{Mo}_2\text{O}_7 \cdot 20\text{aq}$ (28)
 $\text{Al}_2(\text{MoO}_4)_3 \cdot 3\text{Na}_2\text{Mo}_2\text{O}_7 \cdot 22\text{aq}$ (28)
 $(\text{NH}_4)_2\text{Mo}_2\text{O}_7$ (also *supra*, Al salts, and *infra* Cr salts) (28)
 $\text{Cr}_2(\text{MoO}_4)_3 \cdot 3(\text{NH}_4)_2\text{Mo}_2\text{O}_7 \cdot 20\text{aq}$ (19)
 $\text{Cr}_2(\text{MoO}_4)_3 \cdot 3\text{K}_2\text{Mo}_2\text{O}_7 \cdot 20\text{aq}$ (19)
 $\text{Cr}_2(\text{MoO}_4)_3 \cdot 3\text{Na}_2\text{Mo}_2\text{O}_7 \cdot 21\text{aq}$ (19)
 $\text{Hg}_2\text{Mo}_2\text{O}_7$ (29)
 $\text{Na}_2\text{Mo}_2\text{O}_7$ (2)

Trimolybdates, $\text{R}_3\text{Mo}_3\text{O}_{10}$, or $\text{RO } 3\text{MoO}_3$, or $\text{R}_3\text{MoO}_4 \cdot 2\text{MoO}_3$

$(\text{NH}_4)_3\text{Mo}_3\text{O}_{10} \cdot \text{aq}$ (25, 32)
 $(\text{NH}_4)_3\text{Na}_3\text{Mo}_3\text{O}_{10} \cdot \text{aq}$ (38)
 $\text{BaMo}_3\text{O}_{10} \cdot 8\text{aq}$ (23)
 $\text{CaMo}_3\text{O}_{10} \cdot 6\text{aq}$ (7)
 $\text{CoMo}_3\text{O}_{10} \cdot 10\text{aq}$ (11)
 $2\text{CuMo}_3\text{O}_{10} \cdot 9\text{aq}$ (11)
 $\text{MgMo}_3\text{O}_{10} \cdot 10\text{aq}$ (12)
 $\text{K}_3\text{Mo}_3\text{O}_{10} \cdot 8\text{aq}$ (8, 23)
 $\text{Na}_3\text{Mo}_3\text{O}_{10} \cdot 4\text{aq}$, and 7aq (5, 28)
 $(\text{NaNH}_4)_3\text{Mo}_3\text{O}_{10} \cdot \text{aq}$ (38)
 $\text{ZnMo}_3\text{O}_{10} \cdot 10\text{aq}$ (11)

Tetramolybdates, $\text{R}_4\text{Mo}_4\text{O}_{13}$, or $\text{RO } 4\text{MoO}_3$, or $\text{R}_4\text{MoO}_4 \cdot 3\text{MoO}_3$

$(\text{NH}_4)_4\text{Mo}_4\text{O}_{13} \cdot \text{aq}$ (25)
 $\text{BaH}_4(\text{Mo}_4\text{O}_{13}) \cdot 17\text{aq}$ (6)
 $\text{CaMo}_4\text{O}_{13} \cdot 9\text{aq}$ (7)
 $\text{CaH}_4(\text{Mo}_4\text{O}_{13}) \cdot 17\text{aq}$ (7)
 $\text{MgH}_4(\text{Mo}_4\text{O}_{13}) \cdot 19\text{aq}$ (12)
 $\text{KHM}_4\text{O}_{13} \cdot 6\text{aq}$ (10)
 $\text{NaHM}_4\text{O}_{13} \cdot 8\text{aq}$ (14)
 $\text{Na}_4\text{Mo}_4\text{O}_{13} \cdot 6\text{aq}$ (3)
 $\text{ZnMo}_4\text{O}_{13} \cdot 8\text{aq}$ (11)

Octomolybdates, $\text{R}_8\text{Mo}_8\text{O}_{23}$, or $\text{RO } 8\text{MoO}_3$, or $\text{R}_8\text{MoO}_4 \cdot 7\text{MoO}_3$

$\text{Na}_8\text{Mo}_8\text{O}_{23} \cdot 4\text{aq}$ (1*, 6)
 $\text{NaHM}_8\text{O}_{23} \cdot 4\text{aq}$ (1*, 6)

Decamolybdates, $\text{R}_{10}\text{Mo}_{10}\text{O}_{29}$, or $\text{RO } 10\text{MoO}_3$, or $\text{R}_{10}\text{MoO}_4 \cdot 9\text{MoO}_3$

$\text{Na}_{10}\text{Mo}_{10}\text{O}_{29} \cdot 12\text{aq}$ (1*, 6)

Tribasic heptamolybdates (sometimes called *seven thirds molybdates*), $\text{R}_3\text{Mo}_3\text{O}_{24}$, or $3\text{RO } 7\text{MoO}_3$, or $\text{R}_3\text{MoO}_4 \cdot 2\text{RO } 6\text{MoO}_3$

$(\text{NH}_4)_3\text{Mo}_3\text{O}_{24} \cdot 4\text{aq}$ (ordinary ammonium molybdate) (16, 28, 26, 27)
 $\text{Ba}_3\text{Mo}_3\text{O}_{24} \cdot 9\text{aq}$ (28, 31)
 $\text{Mg}_3\text{Mo}_3\text{O}_{24} \cdot 20\text{aq}$ (12)
 $\text{K}_3\text{Mo}_3\text{O}_{24} \cdot 4\text{aq}$ (28, 26)
 $\text{Na}_3\text{Mo}_3\text{O}_{24} \cdot 22\text{aq}$ (18, 26, 36)

Molybdates not included in foregoing classes

$5\text{Al}_2\text{O}_3 \cdot 2\text{MoO}_3$ (18, 28)
 $4(\text{NH}_4)_3\text{O} \cdot 5\text{MoO}_3 \cdot \text{aq}$ (30)

$2\text{BaO } 5\text{MoO}_3 \cdot 6\text{aq}$ (28)
 $\text{CuO } 2(\text{NH}_4)_2\text{O } 5\text{MoO}_3 \cdot 9\text{aq}$ (21)
 $7(\text{NH}_4)_2\text{O } 2\text{Na}_2\text{O } 21\text{MoO}_3 \cdot 15\text{aq}$ (26)
 $7(\text{NH}_4)_2\text{O } 3\text{Na}_2\text{O } 25\text{MoO}_3 \cdot 30\text{aq}$ (26)

For some other complex molybdates, v Bärwald, *C C* 1885 424 (Abstract in *C J* 50, 17)

Molybdates of Cd (1), Di (Frerichs & Smith, *A* 191, 355), La (F & F, *lc* p 365), Pb (28), and Manross, *A* 82, 358), and Sm (Clève, *Bl* [2] 43, 170), have also been prepared

Memoirs where accounts of the molybdates will be found — (1) Schultze, *A* 126, 55 (1*) Ullik, *W A B* 60 [2], 295 (2) Ullik, *A* 144, 214 (3) *Ibid id* 144, 321 (4) *Ibid id* 144, 227, 153, 376 (5) *Ibid id* 144, 223 (6) *Ibid id* 144, 336 (7) *Ibid id* 144, 231, 327, 334 (8) *Ibid id* 144, 208 (9) *Ibid id* 144, 230 (10) *Ibid id* 144, 334 (11) *Ibid id* 144, 232, 233 (12) *Ibid id* 144, 345, 153, 368 (13) *Ibid id* 144, 219 (14) *Ibid id* 144, 233 (15) *Ibid id* 144, 339 (16) Rammeisberg, *P* 127, 298 (17) *Ibid id* 128, 311 (18) Struve, *J pr* 61, 449 (19) *Ibid id* 61, 457 (20) *Ibid id* 61, 459 (21) *Ibid N Petersb Acad Bull* 12, 142 (22) *Ibid J pr* 61, 460 to 466 (23) Svanberg & Struve, *J pr* 44, 257 (24) Atterberg, *J* 1873 258 (25) Berlin, *J pr* 49, 445 (26) Delafontaine, *J pr* 95, 136 (27) Flückiger, *P* 86, 594, cf Werneke, *Fr* 14, 14 (28) Gentile, *J pr* 81, 414 (29) Hirzel, *J* 1852 419 (30) Jean, *C R* 78, 1436 (31) Jorgensen, *Gm K* 2 [2] 217 (32) Kammerer, *J pr* [2] 6, 358 (33) Sonnenschein, *J pr* 53, 340 (34) Steinacker, *J* 1861 238 (35) Widmann, *Bl* [2] 20, 64 (36) Zenker, *J pr* 58, 486 (37) Flemming, *J* 1868 250 (38) Mauro, *B* 14, 1379 (39) Coloriano, *Bl* [2] 50, 451 For an account of the crystalline forms of many molybdates v Zepharovich, *W A B* 58 [2], 111

To indicate the methods of preparation and properties of the molybdates, short descriptions are given of the NH_4 and Na salts, for the others, reference must be made to the original memoirs

Ammonium molybdates (1) *Monomolybdate*, $(\text{NH}_4)_2\text{MoO}_4$. Prepared by dissolving MoO_3 in excess of very conc NH_4Aq , and ppg by alcohol, obtained in small monoclinic crystals by spontaneous evaporation of the trimolybdate in conc NH_4Aq . Effloresces in air, with loss of NH_3 , with water, forms an acid salt. Forms double salts with MgMoO_4 and MnMoO_4 . (2) *Dimolybdate*, $(\text{NH}_4)_4\text{Mo}_2\text{O}_7$. Prepared by evaporating the mother liquor from which the monomolybdate has separated, a white crystalline powder. Forms double salts with $\text{Al}_2(\text{MoO}_4)_3$ and $\text{Cr}_2(\text{MoO}_4)_3$. (3) *Trimolybdate*, $(\text{NH}_4)_6\text{Mo}_3\text{O}_{10}$. Prepared by slow decomposition of ordinary NH_4 molybdate solutions, at temperatures below 10° . Lustrous needles, easily sol hot water, sl sol cold water. Forms a double salt with $\text{Na}_2\text{Mo}_4\text{O}_{10}$. (4) *Tetramolybdate*, $(\text{NH}_4)_8\text{Mo}_4\text{O}_{13}$. Prepared by slowly decomposing solution of ordinary NH_4 molybdate by HClAq or HNO_3Aq , the solution eventually becomes filled with crystals of the tetra-salt, and the liquid is free from Mo. (5) *Tribasic heptamolybdate*, $(\text{NH}_4)_3\text{Mo}_3\text{O}_{24} \cdot 4\text{aq}$ (= $3(\text{NH}_4)_3\text{O} \cdot 7\text{MoO}_3 \cdot 4\text{aq}$). This is the *ordinary*

ammonium molybdate used as a reagent for phosphoric acid, &c. Crystallises from solution of MoO_3 in NH_4Aq in large monoclinic prisms, unchanged in air. The same salt crystallises with 12aq from the mother liquor.

Treatment of ammonium molybdate residues. Venator (*Ar Ph* [3] 23, 713) recommends to add enough FeCl_3Aq to give a brownish colour to the solution, to ppt phosphoric acid by NH_4Aq , to filter, and add BaCl_2Aq whereby Ba molybdate and BaSO_4 are ppt. The pp is thoroughly washed with hot water, and boiled for a long time with $(\text{NH}_4)_2\text{SO}_4\text{Aq}$, $\text{NH}_4\text{molybdate}$ separates from the filtrate on evaporation.

Sodium molybdates. (1) **Monomolybdate**, $\text{Na}_2\text{MoO}_4\cdot 2\text{aq}$. Prepared by fusing together equivalent quantities MoO_3 and Na_2CO_3 , dissolving in water, and evaporating, also by dissolving MoO_3 in $\text{Na}_2\text{CO}_3\text{Aq}$, also by neutralising the solution of any of the other Na salts by Na_2CO_3 . Small lustrous tablets, sol water, solution has an alkaline reaction. Solutions evaporated under 6° give a salt with 10aq, these crystals effloresce to the salt with 2aq. Forms a double salt with K MoO_4 . (2) **Dimolybdate**, $\text{Na}_2\text{Mo}_2\text{O}_7$. Prepared by fusing together MoO_3 and Na_2CO_3 in the proper proportion, and treating with a little cold water, also by adding MoO_3 to the equivalent quantity of molten NaNO_3 (only half of this is decomposed), and treating with cold water. Small lustrous needles, sol with difficulty in water, hot or cold. Melts at incipient redness, and crystallises on cooling. An aqueous solution of this salt evaporated to a syrup yields crystals of $\text{Na MoO}_4\text{aq}$, easily sol water. (3) **Trimolybdate**, $\text{Na}_2\text{Mo}_3\text{O}_{10}\cdot 7\text{aq}$. Prepared by saturating warm $\text{Na}_2\text{CO}_3\text{Aq}$ with MoO_3 , filtering, and allowing to evaporate spontaneously, also by adding excess of HNO_3Aq to a conc solution of MoO_3 in $\text{Na}_2\text{CO}_3\text{Aq}$, also by adding acetic acid to solution of $\text{Na}_2\text{MoO}_4\cdot 2\text{aq}$, also by spontaneous evaporation of solution of $\text{NaHMoO}_4\cdot 8\text{aq}$ after addition of 1 or 2 formula weights of Na_2CO_3 . A voluminous pp consisting of fine needles. Small quantities of a hot conc solution of this salt give $\text{Na}_2\text{Mo}_3\text{O}_{10}\cdot 4\text{aq}$ on rapid evaporation, amorphous mass, easily sol water. Forms a double salt with $(\text{NH}_4)_2\text{Mo}_3\text{O}_{10}$. (4) **Tetramolybdates**, (a) **Normal salt**, $\text{Na}_2\text{Mo}_4\text{O}_{23}$. Prepared by adding the proper quantity of HClAq to $\text{Na}_2\text{MoO}_4\text{Aq}$, and evaporating, also by spontaneous evaporation of solution of $\text{NaHMoO}_4\cdot 8\text{aq}$, to which a little NaCl has been added. Easily sol hot water, sl sol cold water. (b) **Acid salt**, $\text{NaHMo}_4\text{O}_{23}\cdot 8\text{aq}$. Prepared by adding the proper quantity of HClAq to $\text{Na MoO}_4\text{Aq}$. Large monoclinic crystals, sol water, hot or cold. Melts below redness. (5) **Octomolybdates**, (a) **Normal salt**, $\text{Na}_2\text{Mo}_8\text{O}_{52}\cdot 4\text{aq}$. Prepared by digesting the acid salt with 1 formula weight Na_2CO_3 in water. A white insol powder. (b) **Acid salt**, $\text{NaHMo}_8\text{O}_{52}\cdot 4\text{aq}$. Prepared by adding excess of conc HNO_3Aq to boiling $\text{Na}_2\text{MoO}_4\text{Aq}$. A white pp. (6) **Decamolybdate**, $\text{Na}_2\text{Mo}_{10}\text{O}_{60}\cdot 12\text{aq}$. Prepared by adding 2 HClAq to $\text{Na MoO}_4\text{Aq}$, and evaporating on the steam bath. Separates as a white, almost insol, crystalline powder. (7) **Tri-basic heptamolybdate**, $\text{Na}_3\text{Mo}_7\text{O}_{42}\cdot 22\text{aq}$ ($= 3\text{Na}_2\text{O}\cdot 7\text{MoO}_3\cdot 22\text{aq}$). Prepared by dissolving MoO_3 in the calculated quantity of Na_2CO_3 in water, also by mixing equivalent quantities of

$\text{Na}_2\text{Mo}_2\text{O}_7$ and Na_2CO_3 in solution, and evaporating, also from a solution of MoO_3 in NaOH Aq or $\text{Na}_2\text{CO}_3\text{Aq}$, by adding HNO_3Aq as long as the pp of trimolybdate redissolves and until the liquid acquires an acid reaction. Large, lustrous, monoclinic prisms, easily sol water, solution reacts acid. Melts when warmed, and crystallises on cooling, the molten salt is insol water. There are also several complex double sodium molybdates.

ANTIMONO-, ARSENO-, FLUO-, PHOSPHO-, SILICO-, VANADO-, MOLYBDATE.

Antimonoso-molybdates. These salts are most simply regarded as compounds of Sb_2O_3 with MoO_3 , and basic oxides, they are obtained by boiling SbOCl with an acid molybdate solution (Gibbs, *P Am A* 21, 93).

Antimono molybdates. Some of these salts, compounds of Sb_2O_3 with MoO_3 , and basic oxides, are described by Gibbs (*P Am A* 21, 105).

Arsenoso molybdates. Described by Gibbs (*lc* 21, 81). The former salts are obtained by boiling As_2O_3 with conc solutions of acid molybdates, the latter by oxidising the former in alkaline solution. (For composition of foregoing complex salts v abstracts of Gibbs' papers in *C J Abstracts*, 1886 426, 511, 1887 113, v also *Pufahl, B* 17, 217, *Debray, C R* 78, 1408, *Struve, J pr* 58, 493.)

Arseno molybdic acids. Compounds of H_2AsO_4 with MoO_3 , v *Severth, B* 6, 391, *Debray, C R* 78, 1408, *Struve, J pr* 58, 493.

Regarding the relations of the antimono and arseno molybdates to the phospho molybdates, v *Phosphomolybdates*, p 426.

Fluo molybdates. A number of salts are obtained by dissolving molybdates in HFAq and evaporating, others are formed by dissolving metallic oxides along with MoO_3 in HFAq and evaporating, other salts are produced by dissolving hydrated MoO_3 along with metallic oxides in HFAq and evaporating, others by dissolving MoCl_3 in alkali fluorides. These salts may be regarded as *fluomolybdates*, most of them belong to one of the series $\text{M}^i\text{Mo}_i\text{O}_i\text{F}_i$, $\text{M}^i\text{Mo}_i\text{O}_i\text{F}_i$, or $\text{M}^i\text{Mo}_i\text{OF}_i$, the first may be looked on as salts of the hypothetical acid $\text{H}_i\text{MoO}_i\text{F}_i$, derived from H_iMoO_i , the second may be looked on as salts of the hypothetical acid HMoO_iF_i , derived from H_iMoO_i , and the third as salts of the hypothetical acid H_iMoOF_i , derived from the hypothetical H_iMoO_i . The foregoing, and other, fluomolybdates may also be regarded as compounds of metallic fluorides with MoO_3F_2 and MoOF_3 , respectively.

Fluomonomolybdates, $\text{M}^i\text{Mo}_i\text{O}_i\text{F}_i$. These salts, which may also be represented as $2\text{MF MoO}_3\text{F}_i$, are generally obtained by dissolving monomolybdates in HFAq , most of them are sol water, and crystallise well. When carefully heated in air, many form molybdates, others give residues of metallic oxides (Delafontaine, *J* 1867 233). The chief fluomonomolybdates are the following $(\text{NH}_4)_2\text{MoO}_4\text{F}_i\cdot \text{H}_2\text{O}$, trichlorides, by adding slight excess of HFAq to a strongly ammoniacal solution of ordinary $\text{NH}_4\text{molybdate}$, and evaporating, when heated, H_2O and HF are evolved and blue oxide of Mo remains $(\text{NH}_4)_2\text{MoO}_4\text{F}_i$, rhombic plates (Mauro, *Mem R Acad des Lincei* [4] 4, 481). $\text{CoMoO}_3\text{F}_i\cdot 6\text{H}_2\text{O}$, dark red efflorescent crystals,

obtained by dissolving equivalent weights of CoO and MoO₃ in HFAq, and evaporating K₂MoO₄F₂H₂O, lustrous, triclinic crystals, obtained by dissolving K₂MoO₄ in HFAq, also by adding KF to MoO₃ dissolved in HFAq. Other

ever, exceed 6, in one case $n=14$ when $R=Ag$, (Gibbs). R = basic metal usually Na, K, or NH₄, x is generally large, varying from 8 to 46. The chief sub classes of these phospho molybdates are the following (Gibbs) —

10MoO ₃ ·2P ₂ O ₅ ·5R ₂ O xaq,	$R=NH_4$, and K , $x=7$ and 20
10MoO ₃ ·2P ₂ O ₅ ·6R ₂ O xaq,	$R=NH_4$, K , Ag , $x=14$, $R=Na$, $x=23$.
32MoO ₃ ·2P ₂ O ₅ ·6R ₂ O xaq,	$R=NH_4$, $x=28$
36MoO ₃ ·2P ₂ O ₅ ·2R ₂ O xaq,	$R=Na$, x undetermined, >10
36MoO ₃ ·2P ₂ O ₅ ·4R ₂ O xaq,	$R=Na$, x >8
44MoO ₃ ·2P ₂ O ₅ ·5R ₂ O xaq,	$R=K$, $x=22$ "
44MoO ₃ ·2P ₂ O ₅ ·6R ₂ O	$R=NH_4$
44MoO ₃ ·2P ₂ O ₅ ·6R ₂ O xaq,	$R=NH_4$, $x=18$ and 24
44MoO ₃ ·2P ₂ O ₅ ·6R ₂ O xaq,	$R=K$, $x=24$
44MoO ₃ ·2P ₂ O ₅ ·14R ₂ O xaq,	$R=Ag$, $x=23$
48MoO ₃ ·2P ₂ O ₅ ·2R ₂ O xaq,	$R=\frac{1}{2}Co(NH_3)_4(NO_3)_3O$, $x=46$
48MoO ₃ ·2P ₂ O ₅ ·4R ₂ O xaq,	$R=K$, $x=8$

salts of this series are CdMoO₄F₂, NiMoO₄F₂·6H₂O, Rb₂MoO₄F₂·H₂O, 2Na₂MoO₄F₂·H₂O, Ti₂MoO₄F₂·H₂O, ZnMoO₄F₂·6H₂O.

Fluomolybdates, M₁Mo₂O₇F₂ or M₂MoO₄F₂ (Delafontaine, *J* 1867 233). These salts may also be represented as MF MoO₄F₂, they are obtained by dissolving dimolybdates in HFAq, or by treating the salts M₂MoO₄F₂ with HFAq. The chief salts of this series are NH₄MoO₄F₂·H₂O, and KMoO₄F₂·H₂O.

Fluomolybdates other than the foregoing. The salts (NH₄)₂MoO₇F₂ (or MoO₄F₂·3NH₄F) and (NH₄)₂MoO₇F₂ (NH₄)₂MoO₄ (or MoO₄F₂·4NH₄F) (NH₄)₂MoO₄ are described by Mauro (*Mem R Acad der Lincei* [4] 4, 481). The same chemist (*G* 19, 179, *v* also *B* 15, 2509) describes the following salts — K₂MoO₇F₂·H₂O (or MoO₄F₂·2KF·H₂O), obtained by dissolving MoCl₅ or MoO₃·xH₂O in conc hot KFAq, and crystallising K₂Mo₂O₇F₂·H₂O (or 3MoO₄F₂·5KF·H₂O), by dissolving the preceding salt in HFAq and evaporating (NH₄)₂MoO₇F₂ (or MoO₄F₂·2NH₄F), by dissolving MoO₃·xH₂O in HFAq, adding NH₄Aq till the green liquid becomes reddish, then adding HFAq till a green colour is produced again, and evaporating considerably at a moderate temperature (NH₄)₂MoO₇F₂·H₂O (or 3MoO₄F₂·5NH₄F·H₂O), by dissolving the preceding salt in warm HFAq and evaporating.

Phospho-molybdates. The phospho molybdates have been examined by Berzelius, Svanberg a Struve (*J* pr 44, 299), Debray (*B* [2] 5, 404), Rammelsberg (*B* 10, 1776), Finkener (*B* 11, 1638), and especially by Wolcott Gibbs (*P Am A* 17, 62, 18, 232, 21, 50). These salts are produced when phosphoric acid or a soluble phosphate is added to solution of a molybdate, with or without the presence of a free acid, also by fusing together phosphates and molybdates, by dissolving molybdates in soluble in water in phosphoric acid, by digesting MoO₃ with an alkaline phosphate, and by treating mixtures of insoluble phosphates and molybdates with a dilute acid. The phospho-molybdates of the fixed alkaline bases may be dehydrated by careful heating, but MoO₃ is generally partially volatilised. Solutions of these salts in NH₄Aq are decomposed by H₂S with formation of thio-molybdates.

The phospho molybdates belong to the general form $mMoO_3 \cdot 2P_2O_5 \cdot nR_2O$ xaq, m is always an even number varying from 10 to 48, n varies from 2 to 6, and is generally = 6, n may, how-

The only *phospho molybdic acid* which has been isolated is 48MoO₃·2P₂O₅·6H₂O xaq, three hydrates are known in which $x=54$, 92, and 100 (Gibbs).

The salts with 6R₂O are regarded by Gibbs as normal salts, those with less than 6R₂O generally have acid reactions. To the *ordinary ammonium phospho-molybdate* — obtained by adding excess of a mineral acid to a solution containing MoO₃, P₂O₅, and an NH₄ salt — Gibbs assigns the composition 48MoO₃·2P₂O₅·5(NH₄)₂O·16aq (cf Svanberg a Struve, *J* pr 44, 291, Nutting, *Pharmaceut Vierteljahresschrift*, 4, 549, Sonnenschein, *J* pr 53, 342, Lupowitz, *P* 109, 135, Seligsohn, *J* pr 67, 470, Hundeshagen, *Fr* 23, 141, 172, who says that when dried above 130° the salt is 12MoO₃·(NH₄)₂PO₄).

Gibbs (*lc*) describes four other series of phospho molybdates —

Metaphospho molybdates,
 $mMoO_3 \cdot nH_2PO_4 \cdot pR_2O$ xaq

Hypophospho molybdates,
 $mMoO_3 \cdot nH_2PO_2 \cdot pR_2O$ xaq

Phosphoroso molybdates,
 $mMoO_3 \cdot nH_2PO_3 \cdot pR_2O$ xaq

Phosphoroso phospho molybdates,
 $mMoO_3 \cdot nP_2O_5 \cdot nH_2PO_3 \cdot pR_2O$ xaq

Metaphospho molybdates, represented by 10MoO₃·4NH₄PO₃·3(NH₄)₂O 9aq and

14MoO₃·3Ba(PO₃)₂·BaO 55aq are converted by acids into oithophospho molybdates (Gibbs, *P Am A* 21, 116).

Hypophospho molybdates are represented by 8MoO₃·2H₂P₂O₄·2(NH₄)₂O 2aq, this salt was obtained by mixing NaH₂PO₄Aq with solution of (NH₄)₂Mo₂O₇, and adding HClAq (*G*, *P Am A* 18, 232).

Phosphoroso-molybdates are represented by 24MoO₃·4H₂PO₃·4(NH₄)₂O xaq, in which $x=17$ and 25, obtained by adding H₂PO₃Aq (produced by adding water to PCl₅) to solution of (NH₄)₂Mo₂O₇ (*G*, *P Am A* 18, 237, 21, 89).

Phosphoroso phospho molybdates are represented by 72MoO₃·3P₂O₅·2H₂PO₃·9(NH₄)₂O 38aq, obtained by adding the product of the decomposition of PCl₅ by H₂O to solution of 10MoO₃·2P₂O₅·6(NH₄)₂O 14aq (*G*, *lc* 21, 96).

Note — *Pyrophospho-tungstates* have been isolated, of the form 22WO₃·9P₂O₅·pR₂O, $p=13$, 18, and 20, an *ortho-metaphospho tungstate* has also been prepared,

22WO₃·3(P₂O₅·3K₂O) 6NaPO₃·2K₂O 2Na₂O 42aq.

Corresponding molybdates have not yet been isolated

Besides the foregoing phospho molybdates, representatives of phosphovanado- and phosphostanno- molybdates have been prepared by Wolcott Gibbs

Phosphovanado molybdates are represented by $28\text{MoO}_3 \cdot 2\text{P}_2\text{O}_5 \cdot 16\text{V}_2\text{O}_5 \cdot 16(\text{NH}_4)_2\text{O}$ 100aq, and $48\text{MoO}_3 \cdot 2\text{P}_2\text{O}_5 \cdot \text{V}_2\text{O}_5 \cdot 7(\text{NH}_4)_2\text{O}$ 30aq. These salts are obtained by digesting MoO_3 with solution of phosphovanadates, $\text{mP}_2\text{O}_5 \cdot \text{nV}_2\text{O}_5 \cdot \text{pR}_2\text{O}_5$, by heating solutions of vanado molybdates, $\text{mMoO}_3 \cdot \text{nV}_2\text{O}_5 \cdot \text{pR}_2\text{O}_5$, with alkaline phosphates in the presence of acid, and by heating V_2O_5 with solution of an alkaline phospho molybdate (Gibbs, *l.c.* 18, 253)

Phosphostanno molybdates, the salt $16\text{MoO}_3 \cdot 3\text{P}_2\text{O}_5 \cdot 4\text{SnO}_2 \cdot 3(\text{NH}_4)_2\text{O}$ 28aq is obtained by pouring solution of $\text{SnCl}_2 \cdot 2\text{NH}_4\text{Cl}$ into a hot solution of the acid NH_4 phospho molybdate $10\text{MoO}_3 \cdot 2\text{P}_2\text{O}_5 \cdot 5(\text{NH}_4)_2\text{O}$ 7aq (Gibbs, *l.c.* 21, 120)

The arseno- molybdates already referred to (p 425) belong to the series $\text{mMoO}_3 \cdot \text{nAs}_2\text{O}_5 \cdot \text{pR}_2\text{O}_5$ (arsenoso- compounds) where $m=6, 8$, and 12 , $n=3, 2$, and 5 , $p=2$ and 3 ($\text{R}=\text{NH}_4, \frac{1}{2}\text{Mn}, \frac{1}{2}\text{Ba}$), and $x=6, 13$, and 24 (Gibbs, *P Am A* 21, 81). The antimo- molybdates (*v* p 425) belong to the series $17\text{MoO}_3 \cdot 3\text{Sb}_2\text{O}_5 \cdot 6(\text{NH}_4)_2\text{O}$ 21aq (antimonoso- compounds), and the series $7\text{MoO}_3 \cdot 4\text{Sb}_2\text{O}_5 \cdot 5(\text{NH}_4)_2\text{O}$ 12aq (antimono compounds) (Gibbs, *P Am A* 21, 93 a 105)

Silico-molybdates Alkali molybdate solutions produce yellow pps when added to solutions of alkali silicates in presence of HNO_3 (Knop, *C C* 1857 691, 861, Richter, *D P J* 199, 183). According to Paumentier (*C R* 92 1231) these pps have the composition $13\text{MoO}_3 \cdot \text{M}_2\text{SiO}_5 \cdot x\text{H}_2\text{O}$. P obtained the acid $12\text{MoO}_3 \cdot \text{H}_2\text{SiO}_3 \cdot 24\text{H}_2\text{O}$ from the mercurous salt (*C R* 94, 213). This acid forms large yellow regular octahedra, melting at 45° and decomposing below 100° , easily soluble in water and dilute acids, decomposed by excess of NH_4Aq or alkali carbonates with separation of silica

Vanado molybdates These salts belong to two series (Gibbs, *P Am A* 18, 240 a 264)

Vanadio molybdates These salts are represented by $6\text{MoO}_3 \cdot \text{V}_2\text{O}_5 \cdot 2(\text{NH}_4)_2\text{O}$ 5aq, $16\text{MoO}_3 \cdot 2\text{V}_2\text{O}_5 \cdot 5\text{BaO}$ 29aq, and $18\text{MoO}_3 \cdot \text{V}_2\text{O}_5 \cdot 8(\text{NH}_4)_2\text{O}$ 15aq, they are obtained by digesting V_2O_5 (free from VO) with solutions of alkaline molybdates, by heating together solutions of alkaline vanadates and molybdates especially in presence of acid, by boiling solutions of alkaline vanadates or meta vanadates (*e.g.* NH_4VO_3) with MoO_3 , and by the decomposition of phosphovanadomolybdates (*v supra*)

Vanadio-vanadico molybdates The representatives of this series which have been isolated are $28\text{MoO}_3 \cdot \text{VO}_2 \cdot 4\text{V}_2\text{O}_5 \cdot 11(\text{NH}_4)_2\text{O}$ 20aq and $80\text{MoO}_3 \cdot 3\text{VO}_2 \cdot 2\text{V}_2\text{O}_5 \cdot 14\text{BaO}$ 48aq. They are obtained by boiling VO_2 and V_2O_5 with an acid molybdate, by the partial reduction of vanadio molybdates, and by digesting solutions of acid molybdates with solutions containing VO_2 and V_2O_5

Alumino-, chromico-, ferrico-, and manganomolybdates have been obtained by Struve (*Petersburg Acad Bull* 12, 142) and Paumentier (*C. R.* 98, 389). These salts are

classified by Gibbs (*P Am A* 21, 121) as belonging to the forms $10\text{MoO}_3 \cdot \text{M}_2\text{O}_3 \cdot 2\text{K}_2\text{O}$ 15aq, where $\text{M}=\text{Al}, \text{Cr}, \text{Fe}$, $12\text{MoO}_3 \cdot \text{M}_2\text{O}_3 \cdot 6\text{R}_2\text{O} \cdot x\text{H}_2\text{O}$, where $\text{M}=\text{Al}, \text{Cr}, \text{Fe}$, $\text{R}=\text{NH}_4, \text{K}$, or Na , and x is generally $=20$ to 22 , and $16\text{MoO}_3 \cdot \text{Mn}_2\text{O}_5 \cdot 5\text{R}_2\text{O}$ 12aq where $\text{R}=\text{NH}_4$, and K

Molybdenum, alloys of Mo alloys with several metals. With aluminum, a crystalline powder consisting of microscopic rhombic prisms, approximating in composition to MoAl_3 , is obtained by melting together 1 pt MoO_3 dissolved in HFAq and evaporated to dryness, 2 pts cryolite, 2 pts KCl NaCl , and 1 pt Al , and dissolving excess of Al from the regulus by NaOHAq (Wohler a Michel, *A* 115, 102). With iron, grey, hard, brittle, alloys are obtained by directly fusing the metals together, by reducing $\text{Fe}_2\text{O}_3 \cdot 4\text{MoO}_3$ in H at full red heat, an alloy of the composition Mo_3Fe is obtained (Stenacker, *Ueber einige Molybdänverbindungen* [Göttingen, 1861]). Alloys with Cu, Au, Pt, and Ag have been described. An alloy of Mo with lead was found in plates in Utah (Silliman, *Am S* [3] 6, 128)

Molybdenum, bromides of Mo combines directly with Br to form three bromides MoBr_3 , MoBr_2 , and MoBr , no MoBr_4 corresponding with MoCl_4 has been isolated. When Br is passed over strongly heated Mo, an oxybromide is formed, then a greenish grey sublimate of MoBr , near the heated part of the tube, the metal becomes yellow red and is eventually converted into MoBr_2 , between the MoBr_2 and MoBr_3 , isolated lustrous black needles of MoBr are formed. The only halogen compound of Mo which has been gasified, and the mol w of which is known in the gaseous state, is MoCl_4

MOLYBDENUM DIBROMIDE, MoBr_2 ($\text{Mo}_2\text{Br}_4, \text{Br}$) (*Molybdous bromide*)

Preparation —1. By passing Br vapour, much diluted by CO_2 , over strongly heated Mo (Atterberg, *J* 1872 260) —2. By passing Br vapour over Mo at a moderate temperature, or over a heated mixture of MoO_3 and C, strongly heating the MoBr_2 thus produced, and treating the residue with water whereby unchanged Mo may be washed away (Blomstrand, *J pr* 82, 433)

Properties and Reactions —A golden yellow powder, unchanged by heat, insoluble in water and in all acids. Concentrated alkali solutions produce alkali bromide and $\text{MoO}_3 \cdot x\text{H}_2\text{O}$ (Blomstrand, *J pr* 77, 91). Dilute alkali solutions produce alkali bromide, and yellow solutions from which CO_2 ppts $\text{Mo}_2\text{Br}_4(\text{OH})_2 \cdot 8\text{H}_2\text{O}$. This compound, known as *molybdenum bromohydroxide*, is prepared by slowly adding CO_2 to the solution of MoBr in KOHaq , or by adding acetic acid and then passing in CO , or by decomposing the solution when hot by NH_4Cl (*v Molybdenum bromohydroxide*). The reaction of MoBr_2 with KOHaq points to the formula $\text{Mo}_2\text{Br}_4 \cdot \text{Br}$, for the dibromide, the radicle Mo_2Br_4 combines with acid radicles to form salts, *v Molybdenum bromohydroxide*, p 428

MOLYBDENUM TRIBROMIDE, MoBr_3 (*Molybdic bromide*) Obtained by passing Br vapour over gently heated Mo, or by heating a mixture of MoO_3 and C in Br vapour, any MoBr_2 and $\text{MoO}_3 \cdot \text{Br}$ formed are removed by heating in the stream of Br, as these compounds are more volatile than MoBr_3 (Blomstrand, *J pr* 82, 433).

Forms a blackish green mass of small interlaced needles. Sublimes with difficulty, at bright red heat gives MoBr_2 and Br. Unchanged by water, insoluble in conc HClAq and cold dilute HNO_3Aq . Slowly acted on by dilute alkali solutions, decomposed by boiling alkali solution with ppn of black $\text{Mo}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$ (Blomstrand, *l c*).

MOLYBDENUM TETRABROMIDE, MoBr_4 (Molybdic bromide) Formed in small quantity by heating Mo in Br, appears as single, black, lustrous, needles. Easily decomposed by heat to MoBr_2 and Br, the temperatures at which it is formed and decomposed being not very different. Fusible and volatile, forming brown vapours. Deliquescent, soluble in water, forming a brown-yellow solution from which alkalis ppt rusty brown $\text{MoO}_3 \cdot x\text{H}_2\text{O}$ (Blomstrand, *J pr* 82, 433).

Molybdenum, bromochlorides of

$\text{Mo}_2\text{Br}_2\text{Cl}_2 \cdot 3\text{H}_2\text{O}$ and $\text{Mo}_2\text{Cl}_2\text{Br}_2 \cdot 3\text{H}_2\text{O}$ Produced by adding HClAq to a solution of MoBr_2 in KOHAc , and by adding HBrAq to MoCl_2 in KOHAc respectively, cf *Molybdenum bromohydroxide, infra*, and *Molybdenum chlorohydroxide, p 430*.

Molybdenum, bromochlorohydroxide of
 $\text{Mo}_2\text{Cl}_2\text{Br}_2\text{OH} \cdot x\text{H}_2\text{O}$, obtained by adding water to an alcoholic solution of $\text{Mo}_2\text{Cl}_2\text{Br}_2 \cdot 3\text{H}_2\text{O}$, cf *Molybdenum chlorohydroxide de*.

Molybdenum, bromofluoride of

$\text{Mo}_2\text{Br}_2\text{F}_2 \cdot 3\text{H}_2\text{O}$ Produced by adding HFAq to solution of MoBr_2 in KOHAc , cf *Molybdenum bromohydroxide*.

Molybdenum, bromohydroxide of

$\text{Mo}_2\text{Br}_2(\text{OH})_2$ Prepared by dissolving MoBr_2 in KOHAc , and passing in CO_2 or adding acetic acid, or by adding NH_4Cl to a hot solution of MoBr_2 in KOHAc (Blomstrand, *J pr* 82, 433, cf Atterberg, *J* 1872 260). The yellow crystalline pp, consisting of $\text{Mo}_2\text{Br}_2(\text{OH})_2 \cdot 8\text{H}_2\text{O}$, is dried at 100° or *in vacuo* over H_2SO_4 , by placing the pp. over H_2SO_4 , the dihydrate $\text{Mo}_2\text{Br}_2(\text{OH})_2 \cdot 2\text{H}_2\text{O}$ is produced.

$\text{Mo}_2\text{Br}_2(\text{OH})_2$ is a red powder, sol in KOHAc , this solution probably contains $\text{Mo}_2\text{Br}_2\text{OK}$. Addition of acids to this solution generally ppts a compound of the acid radicle with the group Mo_2Br_2 , thus HClAq ppts $\text{Mo}_2\text{Br}_2\text{Cl}_2 \cdot 3\text{H}_2\text{O}$, HBrAq ppts $\text{Mo}_2\text{Br}_2\text{Br}_2 \cdot 3\text{H}_2\text{O}$, HFAq ppts $\text{Mo}_2\text{Br}_2\text{F}_2 \cdot 3\text{H}_2\text{O}$, and HIAq ppts the double compound $(\text{Mo}_2\text{Br}_2\text{I}_2) \cdot \text{Mo}_2\text{Br}_2(\text{OH})_2 \cdot 8\text{H}_2\text{O}$. $\text{H}_2\text{SO}_4\text{Aq}$ ppts $\text{Mo}_2\text{Br}_2\text{SO}_4 \cdot 3\text{H}_2\text{O}$. $\text{K}_2\text{Cr}_2\text{O}_7\text{Aq}$ ppts black $\text{Mo}_2\text{Br}_2\text{CrO}_4 \cdot 2\text{H}_2\text{O}$. Addition of NH_3 , molybdate and acetic acid ppts the molybdate $\text{Mo}_2\text{Br}_2\text{MoO}_4 \cdot 2\text{H}_2\text{O}$, $\text{H}_3\text{PO}_4\text{Aq}$, $\text{H}_2\text{C}_2\text{O}_4\text{Aq}$, and HNO_3Aq also yield pps, that by HNO_3Aq dissolves in excess of the acid. Acetic acid and CO_2 reppt the hydroxide from its solutions in alkalis. The haloid compounds of the radicle Mo_2Br_2 are yellow solids insol in excess of the haloid acids, they are decomposed by boiling water giving $\text{Mo}_2\text{Br}_2(\text{OH})_2 \cdot x\text{H}_2\text{O}$. The sulphate is yellow, and is sol in excess of $\text{H}_2\text{SO}_4\text{Aq}$, this solution gives no pp with AgNO_3Aq , addition of HClAq ppts $\text{Mo}_2\text{Br}_2\text{Cl}_2 \cdot 3\text{H}_2\text{O}$.

The compound $\text{Mo}_2\text{Br}_2(\text{OH})_2 \cdot 8\text{H}_2\text{O}$ might be represented as an oxybromide of Mo, viz as $\text{Mo}_2\text{OBr}_2 \cdot 9\text{H}_2\text{O}$, but the production of $\text{Mo}_2\text{Br}_2(\text{OH})_2$, by drying the hydrate, and the reactions of this compound with acids, are better expressed by representing the compound as a compound of the radicle Mo_2Br_2 , than as an

oxybromide. A corresponding chlorohydroxide, $\text{Mo}_2\text{Cl}_2(\text{OH})_2$, exists (*v p 430*).

Molybdenum, bromo-iodide of, $\text{Mo}_2\text{Br}_2\text{I}_2$ This compound is only known in combination with $\text{Mo}_2\text{Br}_2(\text{OH})_2$, the compound has the composition $(\text{Mo}_2\text{Br}_2\text{I}_2) \cdot \text{Mo}_2\text{Br}_2(\text{OH})_2 \cdot 8\text{H}_2\text{O}$, and is produced by adding HIAq to a solution of $\text{Mo}_2\text{Br}_2(\text{OH})_2$ in KOHAc , cf *Molybdenum bromohydroxide, supra*.

Molybdenum, chlorides of MoCl_3 is formed when Mo is heated in Cl free from O , by heating MoCl_3 in H_2 , MoCl_3 is produced, and MoCl_3 produces MoCl_2 , and MoCl_3 when heated in CO_2 . The formula MoCl_3 represents the molecular composition of the gaseous pentachloride, the mol wt of none of the other chlorides in the gaseous state has been determined.

MOLYBDENUM DICHLORIDE MoCl_2 (? $\text{Mo}_2\text{Cl}_2\text{Cl}_2$) (Molybdous chloride)

Formation—1 By heating MoCl_3 in an in different gas (Blomstrand, *J pr* 77, 95, Liechti a Kempe, *A* 169, 344)—2 By heating Mo with HgCl_2 —3 By carefully heating Mo in Cl largely diluted with CO_2 .

Preparation— MoCl_3 , as pure as possible, is placed in several porcelain boats, which are heated to dull redness in a glass tube, while a slow stream of perfectly dry CO_2 , free from an, is passed through the tube, the contents of the first boat are not quite pure MoCl_2 , but may be purified by gently warming with very dilute HNO_3Aq (Liechti a Kempf, *A* 169, 344).

Properties and Reactions—An amorphous dull yellow powder, unchanged in air, but when heated forms Mo_2O_3 , and then MoO_3 , volatilised with difficulty, insoluble in alcohol, dissolves in alcohol and ether (*L a K, l c*). Dissolves in HClAq and crystallises from this solution as $\text{MoCl}_2 \cdot \text{H}_2\text{O}$, $\text{MoCl}_2 \cdot 2\text{H}_2\text{O}$, or $2\text{MoCl}_2 \cdot 3\text{H}_2\text{O}$ (Blomstrand, *J pr* 77, 95, *v infra*, *Combinations*, No 1). Very slightly sol in HNO_3Aq , dissolved by $\text{H}_2\text{SO}_4\text{Aq}$, soluble in NH_4Aq , on boiling a brown powder containing N is ppd. Decomposed by hot conc alkali solutions with ppn of $\text{MoO}_3 \cdot x\text{H}_2\text{O}$. Dissolved by dilute alkali solutions to form a yellow liquid from which CO_2 or dilute acetic acid ppts $\text{Mo}_2\text{Cl}_2(\text{OH})_2 \cdot x\text{H}_2\text{O}$, *v infra* *Molybdenum chlorohydroxide*. This reaction suggests the formula $\text{Mo}_2\text{Cl}_2\text{Cl}_2$ for the dichloride (cf *Combinations*, No 2), the radicle Mo_2Cl_2 combines with acid radicles to form salts, *v Molybdenum chlorohydroxide*.

Combinations—1 With water to form the hydrates $\text{MoCl}_2 \cdot \text{H}_2\text{O}$, $\text{MoCl}_2 \cdot 2\text{H}_2\text{O}$, and $2\text{MoCl}_2 \cdot 3\text{H}_2\text{O}$ (or $\text{Mo}_2\text{Cl}_2\text{Cl}_2 \cdot 3\text{H}_2\text{O}$, $\text{Mo}_2\text{Cl}_2\text{Cl}_2 \cdot 6\text{H}_2\text{O}$, and $(\text{Mo}_2\text{Cl}_2\text{Cl}_2) \cdot 9\text{H}_2\text{O}$). The first of these hydrates is formed by dissolving MoCl_2 in rather dilute HClAq and allowing to stand, it forms thin yellow plates, insol water. The second hydrate is formed by warming a solution of MoCl_2 in HClAq on a water bath, or by slowly diluting a conc solution, it forms long thin prisms, sol water, alcohol, and ether. Dilute solutions of this hydrate deposit $\text{Mo}_2\text{Cl}_2(\text{OH})_2 \cdot 2\text{H}_2\text{O}$. The third hydrate is formed by dissolving MoCl_2 in hot HClAq and allowing to cool, it forms lustrous needles, insol water, decomposed by hot water, decomposed by heat with loss of H_2O , Cl, and HCl (Blomstrand, *J pr* 77, 95)—2 With the alkali haloid salts to form compounds of the type $2\text{MX} \cdot \text{Mo}_2\text{Cl}_2 \cdot x\text{H}_2\text{O}$

where $M = NH_4$ or K , and $X = Cl, Br$, or I (Blomstrand, *lc*) Obtained by adding excess of haloid acid, HX , to solutions of $MoCl_3$ in KOH or NH_4OH , and evaporating These compounds are decomposed by water, with solution of KX , or NH_4X , and ppt of $Mo_2Cl_4 \cdot x \cdot 3H_2O$ From solutions of these compounds $AgNO_3$ Ag ppts only half of the halogen present, thus 4Br is pptd from $Mo_2Cl_4 \cdot Br_2 \cdot 2KBr$, and 4Cl from $Mo_2Cl_4 \cdot Cl_2 \cdot 2KCl$ These compounds are not decomposed by H_2S in presence of acids, nor by K_2FeCy_4

MOLYBDENUM TRICHLORIDE, $MoCl_3$ (Molybdo-molybdic chloride)

Formation—1 By passing vapour of $MoCl_3$ over heated Mo (Berzelius)—2 By heating $MoCl_3$ by means of an ordinary spirit lamp, in a stream of H (Blomstrand, *J pr* 71, 449)—3 By passing CO_2 charged with $MoCl_3$ through a tube heated in one spot, $MoCl_3$ remains behind the heated place as a thick crystalline crust (Blomstrand, *lc*)

Preparation— $MoCl_3$ is prepared in a hard glass tube, narrowed at intervals so as to form 3 or 4 divisions (*v* *Molybdenum pentachloride, Preparation*), the greater part of the $MoCl_3$ is in the second division, while a little is in the first division Pure dry H is passed through the tube for some time, the second division of the tube is then heated until the $MoCl_3$ begins to sublime into the third division, the temperature of the third division is kept at 250° , at which temperature reduction to $MoCl_2$ proceeds When reduction is complete (known by the change of colour of black $MoCl_3$ into red $MoCl_2$), the H is replaced by a stream of dry CO_2 , and the small quantity of $MoCl_3$ in the first division of the tube is sublimed over the $MoCl_2$, which is not yet quite pure (Liechti & Kempe, *A* 169, 344)

Properties and Reactions—A dark brownish red solid, resembling amorphous P Unchanged in air Heated in a tube drawn to a fine opening, $MoCl_3$ forms as a red crystalline sublimate, heated in air, gives a white woolly sublimate, then brownish red, and finally dark blue, vapours, while impure $MoCl_3$ remains (*L a K, lc*) Decomposes at red heat into $MoCl_2$ and $MoCl$, Insol water and HCl Aq, sol in hot HNO_3 Aq, also in conc H_2SO_4 , forming a deep blue liquid (*L a K, lc*) Decomposed by boiling water, also by KOH Aq or $NaOH$ Aq, with pptn of $Mo_2O_3 \cdot 3H_2O$, warm NH_3 Aq produces a brown powder which contains N A compound of KCl with $MoCl_3$ was obtained by Berzelius by the action of K -amalgam on MoO_3 dissolved in HCl Aq

MOLYBDENUM TETRACHLORIDE, $MoCl_4$ (Molybdo-chloride)

Preparation— $MoCl_4$ is placed in a porcelain boat and heated to dull redness, in a long tube of hard glass, in a very slow stream of pure, air-free, CO_2 , $MoCl_4$ remains in the boat, and $MoCl_3$ sublimes, and is carried forward some distance by the CO_2 (Liechti & Kempe, *A* 169, 344)

Properties—A brown semi-crystalline solid, the vapour is intensely yellow Deliquescent, dissolves in water with hissing sound, the solution shows the reactions of salts of MoO_3 . Somewhat sol in alcohol and ether, dissolved by H_2SO_4 Aq with evolution of HCl , sol in conc HNO_3 Aq

Reactions—1 Kept in contact with air, oxy chlorides are produced—2 Heated in air, $MoO_3 \cdot Cl_2$ and $MoO_3 \cdot 2HCl$ are formed—3 Heated in carbon dioxide, forms $MoCl_3$ and $MoCl_2$, at a higher temperature the $MoCl_3$ is decomposed to $MoCl_2$ and $MoCl$

Combinations—1 With phosphorus pentachloride to form $MoCl_2 \cdot 2PCl_5$, a bluish amorphous solid, obtained by the reaction of PCl_5 with $MoCl_3$ or $MoO_3 \cdot Cl_2$ When this compound is heated, $MoCl_3$, PCl_5 is produced, a metal like lustrous solid, soluble in water to form a black liquid (Cronander, *Bl* [2] 19, 500)—2 With ammonium chloride to form $3MoCl_3 \cdot 2NH_4Cl \cdot 6H_2O$, obtained by saturating NH_4Cl Aq with $MoCl_3$, filtering from NH_4Cl which separates, and allowing to crystallise forms green, deliquescent octahedra (Blomstrand, *J pr* 71, 449)

MOLYBDENUM PENTACHLORIDE, $MoCl_5$ Mol wt 272.75 V.D. at $350^\circ = 137$ This chloride was supposed by Berzelius, also by Blomstrand, to be tetrachloride, its composition was determined by Debray (*C R* 66, 732)

Formation—1 By gently heating Mo or MoS_2 in Cl —2 By strongly heating in Cl a mixture of MoO_3 and C

Preparation—A tube of hard glass is narrowed at intervals so that 4 or 5 divisions are formed, a porcelain boat containing Mo is placed in the tube, and the Mo is heated in a stream of dry HCl so long as any white woolly sublimate of $MoO_3 \cdot 2HCl$ is formed, this sublimate is driven out of the tube by warming in the stream of HCl After cooling, dry air free Cl is passed through the tube for at least an hour, to ensure the removal of every trace of air, and the Mo is then gently heated in the stream of Cl $MoCl_5$ is formed and deposited immediately in front of the porcelain boat, by careful heating while the stream of Cl passes, the $MoCl_5$ can be sublimed into the different divisions of the tube, each of which is then sealed off (Liechti & Kempe, *A* 169, 344)

Properties—A black, crystalline, lustrous mass, a shade of greenish colour indicates presence of oxychloride (*L a K, lc*) Melts at 194° , and boils at 268° (Debray, *C R* 66, 732) The vapour is dark red V.D. at $350^\circ = 137$ (Debray, *lc*, Rieth, *B* 3, 668) May be sublimed unchanged in Cl or CO_2 Decomposed by heating in air with production of $MoO_3 \cdot Cl_2$ Fumes in air, and deliquesces to a brown liquid Dissolves in water with decomposition, solution in a little water is brown, but becomes colourless on dilution Sol in alcohol and ether, forming green liquids Sol in H_2SO_4 Aq and HNO_3 Aq

Reactions—1 Heated in air forms $MoO_3 \cdot Cl_2$, same change results by exposure to air—2 Reacts towards some organic compounds as a carrier of chlorine, e.g. CCl_4 and S_2Cl_2 are produced by passing Cl into CS_2 mixed with $MoCl_3$ (Aronheim, *B* 9, 1788)—3 Heated with ammonia or ammonium chloride produces compounds of Mo with N and H (*v* *Molybdenum, nitride of*, p 430)

Combinations—With phosphoryl chloride, to form $MoCl_3 \cdot POCl_3$, large dark green crystals, melting at 125° – 127° , and boiling at 170° with separation into its constituents (Pirutti, *B* 12, 1326) This compound is formed by heating PCl_5 with MoO_3 , in the ratio $3PCl_5 : MoO_3$, to 170° in a sealed tube, pouring off the brown liquid

from the crystals, washing the latter with CS_2 and drying them in a stream of CO_2 .

Molybdenum, chlorobromides of
 $\text{Mo}_2\text{Br}_2\text{Cl}_2 \cdot 3\text{H}_2\text{O}$ and $\text{Mo}_2\text{Cl}_2\text{Br}_2 \cdot 3\text{H}_2\text{O}$ Produced by adding HClAq to a solution of MoBr_2 in KOHAg , and by adding HBrAq to a solution of MoCl_2 in KOHAg respectively* (cf. *Molybdenum bromohydroxide*, p 428, and *Molybdenum chlorohydroxide*, *infra*)

Molybdenum, chlorobromohydroxide of
 $\text{Mo}_2\text{Cl}_2\text{Br}_2\text{OH} \cdot 2\text{H}_2\text{O}$, obtained by adding water to an alcoholic solution of $\text{Mo}_2\text{Cl}_2\text{Br}_2 \cdot 2\text{H}_2\text{O}$ (v next article)

Molybdenum, chlorohydroxide of
 $\text{Mo}_2\text{Cl}_2(\text{OH})_2 \cdot 2\text{H}_2\text{O}$ Prepared by dissolving MoCl_2 in KOHAg , and neutralising by acetic acid, the octohydrate $\text{Mo}_2\text{Cl}_2(\text{OH})_2 \cdot 8\text{H}_2\text{O}$ is obtained by adding CO_2 or NH_4Cl in place of acetic acid. The dihydrate a light yellow amorphous powder, insol water and alcohol, when freshly pptd and washed with cold water it dissolves readily in acids, but on boiling the solution in HNO_3Aq or $\text{H}_2\text{SO}_4\text{Aq}$ the hydrate is reprecipitated, and is now quite insoluble in acids. The octohydrate forms yellow lustrous crystals (Blomstrand, *J pr* 77, 95)

By adding HBrAq or HIAq to MoCl_2 , or $\text{Mo}_2\text{Cl}_2(\text{OH})_2 \cdot 2\text{H}_2\text{O}$, warming, and allowing to cool, the compounds $\text{Mo}_2\text{Cl}_2\text{Br}_2 \cdot 3\text{H}_2\text{O}$ and $\text{Mo}_2\text{Cl}_2\text{I}_2 \cdot 3\text{H}_2\text{O}$ are obtained, and by evaporating the mother liquor from the second of these, on the water bath, the compound $\text{Mo}_2\text{Cl}_2\text{I}_2 \cdot 6\text{H}_2\text{O}$ is produced. These haloid compounds of the radicle Mo_2Cl_2 are yellowish red crystals, soluble in alcohol, crystallisable from dilute HBrAq and HIAq respectively. Addition of water to the alcoholic solution of $\text{Mo}_2\text{Cl}_2\text{Br}_2 \cdot 3\text{H}_2\text{O}$ ppts $\text{Mo}_2\text{Cl}_2\text{Br}_2\text{OH} \cdot 2\text{H}_2\text{O}$ (Blomstrand, *lc*)

The compounds $\text{Mo}_2\text{Cl}_2(\text{OH})_2 \cdot 2\text{H}_2\text{O}$ and $\text{Mo}_2\text{Cl}_2(\text{OH})_2 \cdot 8\text{H}_2\text{O}$ might be represented as oxybromides of Mo, viz as $\text{Mo}_2\text{OCl}_2 \cdot 3\text{H}_2\text{O}$ and $\text{Mo}_2\text{OCl}_2 \cdot 9\text{H}_2\text{O}$, but the reactions of these compounds with HBrAq and HIAq , and the composition and properties of the salts $2\text{KX Mo}_2\text{Cl}_2\text{X}_2 \cdot x\text{H}_2\text{O}$ (v *Molybdenum dichloride*), are better expressed by representing them as hydrated hydroxides of the radicle Mo_2Cl_2 , than as hydrated oxybromides. A corresponding bromohydroxide, $\text{Mo}_2\text{Br}_2(\text{OH})_2$, exists (v p 428)

Molybdenum, chloro-iodides of
 $\text{Mo}_2\text{Cl}_2\text{I}_2 \cdot x\text{H}_2\text{O}$, $x=3$ and 6. Obtained by adding HIAq to MoCl_2 or to $\text{Mo}_2\text{Cl}_2(\text{OH})_2 \cdot 2\text{H}_2\text{O}$ (cf preceding article)

Molybdenum, fluorides of No fluoride of Mo has been isolated. Solutions of $\text{Mo}_2\text{O}_3 \cdot x\text{H}_2\text{O}$, $\text{MoO}_3 \cdot x\text{H}_2\text{O}$, and $\text{MoO}_2 \cdot x\text{H}_2\text{O}$, in HFAq may contain the corresponding fluorides (Berzelius). The first of these solutions is purple, on evaporation a purple coloured varnish is obtained, addition of KF , NH_4F , or NaF , to this solution, and evaporation, ppts rose coloured powders which may be double compounds of alkali fluoride and MoF_5 . Solution of $\text{MoO}_2 \cdot x\text{H}_2\text{O}$ in HFAq is rose red, but goes colourless on addition of much HFAq , on evaporation it yields a crystalline solid which dissolves in water, KF ppts a reddish-brown solid. Solution of MoO_3 in HFAq is colourless, on evaporation it yields a yellow syrup, addition of KF produces

$2\text{KF MoO}_2\text{F}_2 \cdot \text{H}_2\text{O}$ ($=\text{K}_2\text{MoO}_2\text{F}_2 \cdot \text{H}_2\text{O}$), (cf. *Fluomolybdates* under *Molybdates*, p 425)

Molybdenum, fluobromide of

$\text{Mo}_2\text{Br}_2\text{F}_2 \cdot 3\text{H}_2\text{O}$, prepared by adding HFAq to solution of $\text{Mo}_2\text{Br}_2(\text{OH})_2$ or MoBr_2 in KOHAg cf *Molybdenum, bromohydroxide of*, p 428

Molybdenum, haloid compounds of The haloid compounds of Mo which have been isolated are MoBr_2 , MoBr_3 , MoBr_4 , MoCl_2 , MoCl_3 , MoCl_4 , and MoCl_5 , the three bromides are reformed by the direct union of Mo and Br, MoCl_2 is produced by heating Mo in Cl, MoCl_3 is obtained by partial reduction (by H) of MoCl_5 , and MoCl_2 and MoCl_3 are the products of the decomposition by heat of MoCl_5 . The only one of these compounds whose molecular weight is known in the gaseous state is MoCl_5 . The reactions of the dibromide and dichloride leave little doubt that the molecular formulae of these compounds are not less than Mo_2X_4 , these compounds react as $\text{Mo}_2\text{X}_4\text{X}_2$, the radicles Mo_2X_4 combine with acid radicles to form salts such as $\text{Mo}_2\text{Br}_2\text{SO}_4$ and $\text{Mo}_2\text{Cl}_2\text{Br}_2$. These radicles Mo_2X_4 also combine with alkali haloid compounds to form $2\text{MY Mo}_2\text{X}_4\text{Y}_2 \cdot x\text{H}_2\text{O}$ where M = alkali metal and Y = halogen

No fluoride or iodide of Mo has been isolated with certainty

Molybdenum, hydroxides or hydrated oxides of, v *Molybdenum, oxides of*, p 431, also *Molybdic acids*, p 423

Molybdenum, hydroxychlorides of, $\text{Mo}(\text{OH})_2\text{Cl}$, v *Chloromolybdic acid*, p 423, $\text{Mo}_2\text{Cl}_2(\text{OH})_2$, v *Molybdenum, chlorohydroxide of*, *supra*

Molybdenum, hydroxybromide of, $\text{Mo}_2\text{Br}_2(\text{OH})_2$, v *Molybdenum, bromohydroxide of*, p 428

Molybdenum, iodides of None has been isolated with certainty. A solution of $\text{MoO}_2 \cdot x\text{H}_2\text{O}$ in HIAq gives a crystalline solid on evaporation, this solid is sol water, on heating it gives MoO_2 and HI (Berzelius)

Molybdenum, iodobromide of $\text{Mo}_2\text{Br}_2\text{I}_2$ Known only in combination with $\text{Mo}_2\text{Br}_2(\text{OH})_2$, as $(\text{Mo}_2\text{Br}_2\text{I}_2)_2 \cdot \text{Mo}_2\text{Br}_2(\text{OH})_2 \cdot 8\text{H}_2\text{O}$, which is produced by adding HIAq to a solution of $\text{Mo}_2\text{Br}_2(\text{OH})_2$ in KOHAg , cf *Molybdenum, bromohydroxide of*, p 428

Molybdenum, iodochloride of
 $\text{Mo}_2\text{Cl}_2\text{I}_2 \cdot x\text{H}_2\text{O}$, $x=3$ and 6. Obtained by adding HIAq to MoCl_2 or $\text{Mo}_2\text{Cl}_2(\text{OH})_2 \cdot 2\text{H}_2\text{O}$, cf *Molybdenum, chlorohydroxide of*, *supra*

Molybdenum, nitride of Mo_2N_2 , or (?) Mo_2N , A greyish black powder, obtained by passing dry NH_3 over MoCl_5 heated to full redness (Uhrlaub, P 101, 605). Heated to whiteness in NH_3 , Mo is produced

COMPOUNDS OF MOLYBDENUM WITH NITROGEN AND HYDROGEN These compounds are produced by heating MoCl_5 or MoO_3 in NH_3 , also by heating NH_4Cl with MoCl_5 . MoCl_5 melts when heated in dry NH_3 , NH_4Cl volatilises, and on continued warming a blackish mass remains, which when powdered, quickly washed with water, and dried over H_2SO_4 , has the composition $\text{Mo}_2\text{N}_2\text{H}_4$ (Uhrlaub, P 101, 605). Wöhler (A 108, 258) regarded this compound as 4MoN , $\text{Mo}(\text{NH}_2)_2$. By heating to incipient redness, a body of the composition $\text{Mo}_2\text{N}_2\text{H}_4$ is produced (Uhrlaub, *lc*). Both these compounds are black powders, heated in air they burn to MoO_3 , and give off NH_3 , with molten KOH they evolve NH_3 , and with NaClOAg

they evolve N (Uhrlaub, *lc*) By heating to a temperature slightly above that at which NH_4Cl is sublimed, Tuttle obtained Mo_3N_4 (*A* 101, 285) These compounds are all reduced to Mo by very strongly heating in NH_3 According to Tuttle (*lc*), purple coloured to black crystals are obtained by heating MoO_3 to redness in NH_3 , and these crystals consist of MoO_3 , Mo nitride, and Mo amide

Molybdenum, oxides and hydrated oxides of The three oxides MoO , Mo_2O_3 , and MoO_3 have been isolated, there also exist one or more oxides with more O than Mo O, and less than MoO_3 , a hydrate of the monoxide (MoO) was obtained by Blomstrand, but little is known concerning it The oxides Mo_2O_3 and MoO_3 are basic, they dissolve in acids to form salts, but little is known with certainty regarding these salts MoO_3 is acidic, it is the anhydride of molybdic acid HMoO_4 , and several other acidic hydrates of MoO_3 are known, *v* **Molybdenum, acids of**, p 423 MoO_3 also combines with some acids, *eg* HCl , and acidic radicles, *eg* SO_3 The mol wt of none of the oxides is known with certainty, as none has been gassed

HYDRATED MOLYBDOS OXIDE, $\text{MoO} \cdot x\text{H}_2\text{O}$ By treating MoBr_3 with conc KOH aq, Blomstrand obtained a black hydrate of MoO (*J pr* 77, 91) This compound has been examined only slightly

MOLYBDENUM SESQUIOXIDE, Mo_2O_3 (*Molybdomolybdic oxide*) This oxide was regarded by Berzelius as the protoxide, its composition was determined correctly by Blomstrand (*J pr* 71, 454, *cf* Svanberg a Struve, *J pr* 44, 257)

Formation—1 By heating MoO_3 to redness for some time in a stream of H (*S a S, lc*) — 2 By long continued digestion of MoO_3 with Zn and HCl aq, the product oxidises readily in air (Berzelius) — 3 By decomposing MoCl_5 by KOH aq, washing the pp of $\text{Mo}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$ with cold water, pressing between paper, drying over H_2SO_4 and heating *in vacuo* (Blomstrand, *J pr* 71, 455)

Preparation— $\text{MoO}_3 \cdot 3\text{H}_2\text{O}$ is obtained by one of the following methods it is dried over H_2SO_4 , and heated to a moderate temperature, *in vacuo*, or out of contact with air (if heated in air it is oxidised to MoO) To obtain $\text{Mo}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$, HCl aq is added to a conc solution of a molybdate till the pp which forms is redissolved, this solution is digested with Zn until the colour changes from blue to red brown and finally to black (Blomstrand, *J pr* 71, 455, Rammelsberg, *P* 127, 284, Weincke, *Fr* 14, 1, Pisani, *Fr* 4, 420, Macagno, *B* 8, 258, *cf* O von der Ploiden, *B* 15, 1925) Addition of NH_4 aq to this black solution ppts $\text{Mn}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$, care should be taken to ppt as little $\text{ZnO} \cdot \text{H}_2\text{O}$ as possible along with the $\text{Mo}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$ (the $\text{ZnO} \cdot \text{H}_2\text{O}$ begins to come down when most of the $\text{Mo}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$ is pptd) The pp is washed with water containing a little HCl , to remove $\text{ZnO} \cdot \text{H}_2\text{O}$, then with pure cold water (Blomstrand, *lc*), and is then pressed between paper $\text{Mo}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$ is also obtained by reducing the HCl solution of a molybdate by K amalgam containing very little K, and then ppg by NH_4 aq (Berzelius), also by adding NH_4 aq to the solution obtained by boiling HCl aq and Cu with PbMoO_4 (von Kobell, *J pr* 41, 158, *cf* Hirzel, *J* 1850 309), or with MoO_3 aq (Rammelsberg, *P* 127, 281)

Properties and Reactions—A black solid. As obtained by reducing MoO_3 with Zn and HCl aq (*v* **Formation** No 2), Mo_2O_3 is brass-yellow and shows the same crystalline form as the MoO_3 from which it is prepared, this variety of Mo_2O_3 oxidises in air more rapidly than the black variety obtained by heating the hydrate Heated in air, burns to MoO_3 Insol acids

HYDRATE OF MOLYBDENUM SESQUIOXIDE, $\text{Mo}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$ A black solid obtained as described under **Molybdenum sesquioxide**, **Preparation** The moist hydrate oxidises by exposure to air Dissolves slowly in acids to form purple solutions, which are opaque but become transparent and greyish brown when much diluted By evaporation, dark grey or black, non crystallisable, salts are obtained, these salts have not been examined at all satisfactorily, a phosphate, nitrate, borate, and several sulphates were obtained by Berzelius (*P* 6, 331, 369)

MOLYBDENUM DIOXIDE, MoO_2 (*Molybdic oxide*) By strongly heating NH_4 molybdate, in a closed crucible, Bucholz obtained a dark metallic mass which he regarded as Mo dioxide (*Scher J* 9, 485), but Uhrlaub (*P* 101, 605) and Tuttle (*A* 101, 285) showed that this substance contains N and H

Formation—1 By reducing MoO_3 or $\text{Hg} \cdot \text{MoO}_3$ (*H Rose, P* 75, 319) by heating in H (Svanberg a Struve, *J pr* 44, 257, Rammelsberg, *P* 127, 281) — 2 By melting $\text{Na}_2\text{Mo}_2\text{O}_7$ with $\frac{1}{3}$ its weight of Zn, added in small pieces, and repeatedly washing the cold mass first with warm conc KOH aq and then with warm HCl aq, the product usually contains a little MoO_3 (Ullrich, *A* 144, 227) — 3 By strongly heating $\text{Na}_2\text{Mo}_2\text{O}_7$ in H and removing Na MoO_4 produced by very dilute KOH aq (*S a S, lc*) ($? \text{Na}_2\text{Mo}_2\text{O}_7 + 2\text{H}_2 = \text{Na MoO}_4 + 2\text{H}_2\text{O} + 2\text{MoO}_2$)

Preparation—A mixture of 1 part MoO_3 , 1 part K CO_3 , and $\frac{1}{2}$ part H_2BO_3 , is very strongly heated in a Pt crucible, the crystals of MoO found in the mass on cooling are washed with H_2O , then with dilute NH_4 aq, and then with dilute HCl aq (Mauro a Panebianco, *G* 11, 501, *cf* Muthmann, *A* 238, 108)

Properties and Reactions—A dark brown powder, appearing somewhat purple in direct sunlight Obtained by fusing MoO_3 with K_2CO_3 and H_2BO_3 (*v* **Preparation**), MoO_2 forms copper red to greyish, metallic, very lustrous prisms, *SG* 6.44 at 10° (*M a P, lc*) Unacted on by alkali solution, HCl aq or HFA aq, oxidised to MoO_3 by HNO_3 aq, or by heating in air, heated in Cl_2 forms MoO_2Cl_2 Slightly soluble in conc. H_2SO_4 , and cream of tartar solution

HYDRATE OF MOLYBDENUM DIOXIDE, $\text{MoO}_2 \cdot x\text{H}_2\text{O}$ Obtained by adding NH_4 aq to solution of MoCl_5 , washing the pp with NH_4Cl aq, then with alcohol, pressing between paper, and drying *in vacuo* over H_2SO_4 The solution of MoCl_5 may be prepared by digesting a conc solution of MoO_3 in HCl aq with powdered Mo or with Cu (in absence of air) till all is dissolved (Rammelsberg, *P* 127, 281, *cf* Hirzel, *J* 1850 309)

$\text{MoO}_2 \cdot x\text{H}_2\text{O}$ is reddish-brown to nearly black, it closely resembles $\text{Fe}_2\text{O}_3 \cdot 8\text{H}_2\text{O}$ Slightly soluble in water, forming a reddish yellow liquid, which reddens litmus, and from which salts (*eg* NH_4Cl) reppt the hydrate This solution gela-

latimises after long standing, and dries to a brown black, insoluble, hydrate (Berzelius, *Lehrbuch*) Moist $\text{MoO}_3 \cdot x\text{H}_2\text{O}$ oxidises superficially in air, forming the blue oxide Heated *in vacuo* MoO_3 is produced $\text{MoO}_3 \cdot x\text{H}_2\text{O}$ is insoluble in caustic alkali solution, but dissolves in alkaline carbonates and bicarbonates, these solutions reprecipitate the hydrate on boiling, by standing in air molybdates are formed (Berzelius)

$\text{MoO}_3 \cdot x\text{H}_2\text{O}$ dissolves in acids to form salts These salts are generally purple red to black, only a few are crystallisable, their solutions oxidise in air, alkalis precipitate $\text{MoO}_3 \cdot x\text{H}_2\text{O}$ from these solutions The same salts may be prepared by digesting MoO_3 and Mo with acids till the colour is reddish-brown Berzelius (*P* 4, 153, 6, 331, 369, 7, 261) prepared some of the salts of MoO_3 , but they have not been examined satisfactorily By mixing MoO_3 in HClAq with NH_3 , molybdate, Rammelsberg obtained a crystalline salt $(\text{NH}_4)_2\text{O} \cdot 2\text{MoO}_3 \cdot 4\text{MoO}_3 \cdot 9\text{H}_2\text{O}$ (*P* 127, 281)

MOLYBDENUM TRIOXIDE, MoO_3 (Molybdic anhydride) This oxide was obtained by Scheele in 1778 soon after he had distinguished native Mo sulphide from galena

Occurrence—As molybdenum ochre in small quantities in a few localities

Formation—1 By heating Mo, or any of the lower oxides, for some time in air—2 By prolonged heating of the lower oxides in steam (Regnault, *A Ch* 62, 356)—3 By roasting MoS_2 in air, or oxidising it by HNO_3Aq (*v* Molybdenum, Preparation, p 421)—4 From NH_3 , molybdate, by evaporating with excess of HNO_3 and washing the residue with water (Brunner, *D P J* 150, 372), also by heating in thin layers spread out in a flat porcelain dish (Wilke, *A* 144, 205)

Preparation—Impure MoO_3 is dissolved in NH_3Aq , H_3PO_4 is removed by ppt with MgCl_2Aq , the filtrate is evaporated, the crystals of NH_4 molybdate which separate are heated with excess of HNO_3Aq and the residual MoO_3 is washed with water (For details as to preparation of MoO_3 from Mo ores *v* MOLYBDENUM, Preparation, p 421)

Properties—A white, light, porous, solid, when thrown into water it separates into small, thin, lustrous, scales S G 4.39 at 21° (Schafarik, *J pr* 90, 12) Strongly heated in an open vessel, sublimes in colourless, transparent, rhombic needles (Nordenskjöld, *P* 112, 160) Melts at red heat Soluble in 500 parts cold water (Bucholz, *Scher J* 9, 485), in 960 parts hot water (Hatchett, *T* 1795 323) The solution reddens litmus, tastes metallic (Wittstein, *J* 1860 159, Muller, *J pr* 80, 119) Soluble in acids, but scarcely soluble after being strongly heated Dissolves in alkali, alkaline carbonate, and cream of tartar solutions

Reactions—1 By heating in hydrogen MoO_3 is reduced to MoO_2 , Mo_2O_3 , or Mo, according to the temperature employed—2 Reduced to Mo by treating with potassium or sodium—3 Reduced to Mo_2O_3 by digesting with hydrochloric acid and zinc or copper, &c., also by potassium-amalgam—4 Reduced to the blue oxide (*v* p 433) by a little hydrogen sulphide (excess of H_2S produces black Mo sulphide), also by solution of sulphur dioxide, hydrogen iodide solution, boiling hydrochloric acid, nitric oxide, or stannous

chloride—5 Heated with ammonia, compounds of Mo with N, and with N and H, are produced (*v* Molybdenum, nitride of, p 430)—6 Heated with sulphur, MoS_2 is produced—7 Oxy chlorides, along with molybdates, oxybromides, or oxyfluorides are formed by heating MoO_3 with metallic chlorides, bromides, or fluorides (Schulze, *J pr* [2] 21, 440)—8 Various reduction products result by heating with potassium iodide (Schulze, *l c*)—9 Hydrogen chloride produces $\text{MoO}_3 \cdot 2\text{HCl}$ ($=\text{MoO}(\text{OH})\text{Cl}_2$, *v* Chloromolybdic acid, p 423)—10 Hydrofluoric acid probably produces MoO_2F_2 (*v* Molybdenum, oxyfluoride of, p 423)—11 Dissolves in fairly conc sulphuric acid, on evaporation, crystals of $\text{MoO}_3 \cdot \text{SO}_3$ separate (*v* infra, Combinations, No 3) Hot dilute $\text{H}_2\text{SO}_4\text{Aq}$, saturated with MoO_3 , gelatinises on cooling—12 Reacts with alkalis to form molybdates, $\text{R} \cdot \text{MoO}_4$, &c. (*v* p 423)—13 Oxy chloride, MoO_2Cl_2 , is formed by heating in chlorine (H Rose, *P* 75, 319)—14 Oxychlorides are produced by heating with phosphorus tri- or penta chloride (Michaelis, *J* 1871 249, Schiff, *A* 102, 116)

Combinations—1 With water to form various acidic hydrates (*v* Molybdic acids, p 423, also infra, Hydrates of molybdenum trioxide)—2 With hydrogen chloride to form $\text{MoO}_3 \cdot 2\text{HCl}$ This compound is produced as a white, loose, sublimate, by heating MoO_3 in HCl to 150° – 200° , it is easily soluble in water, and can be sublimed unchanged in HCl (Debray, *C R* 46, 1101) This compound may be regarded as an acid and called chloromolybdic acid, $\text{MoO}(\text{OH})\text{Cl}$ (*v* p 423)—3 With sulphur trioxide to form $\text{MoO}_3 \cdot \text{SO}_3$, and with SO_2 and H_2O to form $\text{MoO}_3 \cdot \text{SO}_3 \cdot 2\text{H}_2\text{O}$ $\text{MoO}_3 \cdot \text{SO}_3$ was obtained by Schultz Sellac (*B* 4, 14) by evaporating a solution of MoO_3 in fairly conc H_2SO_4 , it forms lustrous, transparent, deliquescent crystals, which lose SO_3 when heated $\text{MoO}_3 \cdot \text{SO}_3 \cdot 2\text{H}_2\text{O}$ was obtained by Anderson (*J B* 22, 161) by adding excess of $\text{H}_2\text{SO}_4\text{Aq}$ to BaMoO_4 , filtering, and evaporating—4 Combines with the oxides of phosphorus, arsenic, antimony, vanadium, and silicon, in presence of basic oxides, to form salts of complex acids, *v* Anti-mono, &c., molybdates, pp 425 et seq—5 With ammonium fluoride to form $\text{MoO}_3 \cdot 2\text{NH}_4\text{F}$ (? NH_4 salt of $\text{MoO}(\text{OH})\text{F}_2$), obtained in lustrous, light yellow, octahedral crystals, by adding NH_3Aq to an aqueous solution of $\text{MoO}_3 \cdot \text{F}_2 \cdot 3\text{NH}_4\text{F}$ (this salt is obtained by evaporating a solution of NH_4 molybdate in excess of NH_4FAq acidified by HFAq) (Mauro, *Mem R Acad dei Lincei*, [4] 4, 481)

HYDRATES OF MOLYBDENUM TRIOXIDE Six compounds $x\text{MoO}_3 \cdot y\text{H}_2\text{O}$ have been isolated Four of these have been described under Molybdic acids (p 423), the remaining two are described here

Monomolybdic dihydrate, $\text{MoO}_3 \cdot 2\text{H}_2\text{O}$ A yellow, crystalline, solid Slowly separates from a solution of 15 grs ordinary NH_3 molybdate in 1000 cc water mixed with 1000 cc HNO_3Aq S G 1.16, may be washed with cold water, sol in water and acids, solution reddens litmus and turns turmeric brown, conc HNO_3Aq separates MoO_3 (Millingk, *Gm -K* 2 [2], 170)

Ootomolybdic monohydrate, $8\text{MoO}_3 \cdot \text{H}_2\text{O}$ Obtained by ppg solution of or

dinary NH_4 molybdate by BaCl_2 Aq. decomposing the Ba salt by exactly the equivalent quantity of dilute boiling H_2SO_4 Aq. filtering, evaporating over H_2SO_4 , and drying at $160^\circ\text{--}170^\circ$ (Ullik, *A* 144, 329, 153, 373) Soluble in water

OXIDES OF MOLYBDENUM OTHER THAN THE DI-, SESQUI- AND TRI- OXIDE Several oxides appear to exist intermediate between MoO_3 and MoO_2 , their compositions are not yet settled Some of them form hydrates of which some are soluble and some insoluble in water The substance known as 'blue oxide of molybdenum,' formed by the action of reducing agents on acid solutions of MoO_3 , is probably Mo_2O_5

References— Mo_2O_5 , by heating lower oxides in air (Berzelius, *P* 6, 331, 369) Mo_2O_5 , by heating $(\text{NH}_4)_2\text{O} \cdot 2\text{MoO}_3 \cdot 4\text{MoO}_3 \cdot 9\text{H}_2\text{O}$ (Rammelsberg, *P* 127, 281) Mo_2O_5 , by heating NH_4 molybdate with MoO_3 (Berlin, *J pr* 49, 447, Uhrlaub, *P* 101, 605), also by electrolysis of molten MoO_3 (Wöhler a Bañ, *A* 110, 275) Mo_2O_5 , by heating MoO_3 with KI (Schulze, *J pr* [2] 21, 440) $\text{Mo}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$, by adding NH_4 molybdate or MoO_3 to MoO in HCl Aq (Berzelius, Rammelsberg, *lc*) Regarding properties of the blue products obtained by reducing acid solutions of MoO_3 , v Schiff, *A* 120, 47, Maschke, *Fr* 12, 384, Berzelius, *Gm K* 2 [2], 164, O von der Pfordten, *B* 15, 1925 Fairley (*C J* 31, 142) obtained indications of an oxide of Mo with more O than MoO_3 , by adding H_2O Aq to MoO_3 dissolved in excess of alkali solution

Molybdenum, oxybromide of $\text{MoO} \cdot \text{Br}_2$ Obtained by passing Br vapour over heated MoO_3 , by melting MoO_3 with H_3BO_3 , powdering, and then heating with KBr, by the reaction of MoO_3 on many metallic bromides (Schulze, *J pr* [2] 21, 442) Yellow crystals, may be sublimed, deliquescent and soluble in water The compound $\text{Mo}_2\text{Br}(\text{OH}) \cdot 8\text{H}_2\text{O}$ may be represented as an oxybromide $\text{Mo}_2\text{OBr} \cdot 9\text{H}_2\text{O}$, but is better regarded as hydrated bromohydroxide (*q v*, p 428)

Molybdenum, oxychlorides of Many oxychlorides of Mo exist They are produced by the reaction of Mo oxides with Cl, Mo chlorides with O, and PCl_5 and metallic chlorides with MoO_3

I Green oxychloride $\text{Mo}_2\text{O}_5\text{Cl}_2$, more probably MoOCl_2 (Blomstrand, *J pr* 71, 459, Putzbach, *A* 201, 123) By heating a mixture of Mo and MoO_3 in air free Cl, and in many other ways Melts and sublims below 100° Soluble in water Reduced by H at high temperatures, Putzbach (*lc*) describes $\text{Mo}_2\text{O}_5\text{Cl}_2$ as a product of this reduction

II Yellow white oxychloride MoO_2Cl_2 By passing Cl over heated MoO_3 or MoO (H Rose, *P* 75, 319), by heating Mo chlorides in presence of O (Blomstrand, *lc*), and in many other ways (*v* Schulze, *J pr* [2] 21, 441, Michaelis, *J* 1871 249, Schiff, *A* 102 216, Putzbach, *lc*) Melts at a high temperature when heated under pressure, at ordinary pressure sublims without melting, soluble in water and alcohol

III Violet oxychloride $\text{Mo}_2\text{O}_5\text{Cl}_4$, perhaps MoOCl_3 (Blomstrand, *lc*) Produced along with the green oxychloride, than which it is much less easily volatilised Dark violet-red crystals, deliquescent

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IV Brown oxychloride, $\text{Mo}_2\text{O}_5\text{Cl}_3$ (Blomstrand, *lc*) Produced, along with $\text{Mo}_2\text{O}_5\text{Cl}_2$, by heating Mo_2O_5 in dry Cl, purified by sublimation in H Forms large dark brown crystals, melts easily, deliquescent and soluble in water Putzbach (*lc*) describes a brown oxychloride, $\text{Mo}_2\text{O}_5\text{Cl}_4$, obtained in the preparation of MoOCl_3 , also a red oxychloride, $\text{Mo}_2\text{O}_5\text{Cl}_3$, obtained by heating the violet compound in CO_2

The compound $\text{Mo}_2\text{Cl}_4(\text{OH}) \cdot 2\text{H}_2\text{O}$ may be represented as an oxychloride, $\text{Mo}_2\text{OCl}_3 \cdot 3\text{H}_2\text{O}$, but its reactions are more those of chlorohydroxide (*q v*) $\text{MoO} \cdot 2\text{HCl}$, which might be represented as $\text{MoO}_2\text{Cl}_2 \cdot \text{H}_2\text{O}$, is described as chloromolybdic acid (*q v*)

Molybdenum, oxyfluoride of MoO_2F_2 Produced by heating MoO_3 with metallic fluorides, as a blue-white amorphous sublimate (Schulze, *J pr* [2] 21, 442) A number of compounds are known which may be regarded as double compounds of MoO_2F_2 with alkali fluorides, others may be regarded as double compounds of MoOF_3 with alkali fluorides, these are described as fluo molybdates (*p* 425)

Molybdenum, phosphide of Mo_2P_2 A grey crystalline powder, SG 6 17, oxidised by heating in air, with conc HNO_3 Aq forms H_3PO_4 and MoO_3 , heated in Cl forms MoCl_3 and PCl_5 , reduces Cu solutions with ppn of Cu Produced by heating a mixture of 1 pt MoO_3 with 2 pts glacial phosphoric acid to a high temperature for some time, and washing with HCl Aq and then with NaOH Aq (Wöhler a Rautenberg, *A* 109 374)

Molybdenum, salts of Little is known with certainty of the salts obtained by replacing H of oxyacids by Mo MoO_2 and Mo_2O_3 dissolve in acids to form salts, but accurate analyses and description of those salts are lacking (*v* Molybdenum dioxide and sesquioxide, p 431) MoO_3 combines with SO_2 to form MoSO_3 , and with HCl to form $\text{MoO} \cdot 2\text{HCl}$, but those compounds are probably rather acids than salts (*v* Molybdenum trioxide, p 432) The radicles Mo_2Br , and Mo_2Cl , combine with certain acidic radicles to form salts (*v* Molybdenum bromohydroxide, and chlorohydroxide, pp 428, 430)

Molybdenum, selenide of MoSe_2 Obtained, but not free from a greyish blue substance which is produced along with it, by saturating acid NH_4 molybdate with H Se gas (Uelsmann, *A* 116, 125)

Molybdenum, sulphides of Three are known, MoS_2 , MoS , and MoS_3 As none has been gasified the molecular weight of none is known with certainty MoS and MoS_3 , and probably also MoS_2 , are all acidic towards the alkali sulphides

MOLYBDENUM DISULPHIDE, MoS_2 Occurs native, as molybdenite, SG 4 138 to 4 569 Prepared by heating to redness a mixture of 1 pt MoO_3 and 2 pts S, in a crucible, till excess of S is burnt off, and washing the residue with warm water as long as the washings are coloured (Svanberg a Struve, *J pr* 44, 264) According to Carnot (*Bl* [2] 32, 164) compounds of Mo generally yield MoS , when heated in a stream of dry H_2S A black, lustrous powder, insol water, sol conc H_2SO_4 , with evolution of SO_2 , and production of a blue liquid Forms MoO_3 , when roasted in air, or heated with conc.

F F

HNO_3Aq , readily attacked by Cl , more slowly by Br , not acted on by aqueous alkali but after fusion with alkali dissolves in water (? with formation of a thio-salt) Slowly reacts with steam at high temperatures (Regnault, *A. Ch.* 62, 385) Reduced to Mo by strongly heating in H (O von der Pfordten, *B* 17, 731)

MOLYBDENUM TRISULPHIDE, MoS_3 (Thiomolybdic anhydride) Obtained by passing H_2S into solution of a molybdate until decomposition is complete, i.e. till all molybdate is changed to a thiomolybdate, adding dilute HClAq in slight excess, warming gently (Atterberg, *J* 1873 258), washing, and drying at a moderate temperature A black powder, red-brown when moist Goes to MoS_2 when heated E sol alkali sulphide solutions, slowly dissolved by alkali or alkali hydrosulphide solution, the solutions thus obtained contain *thiomolybdates*, M_2MoS_3 (*v* THIOMOLYBDIC ACIDS AND THEIR SALTS, *infra*) MoS_3 is reduced to Mo by heating in H (O von der Pfordten, *B* 17, 731)

MOLYBDENUM TETRASULPHIDE, MoS_4 Prepared by fusing a mixture of 2 pts MoO_3 and 1 pt K_2CO_3 , powdering, boiling with water, filtering, diluting, saturating with H_2S , washing the pp with cold and then with hot water, then treating with HClAq , and heating the chocolate brown powder thus obtained to 140° in a stream of H_2S (Kruss, *B* 16, 2044) Also obtained by heating MoS_3 to 100° in H_2S (K, *lc*) Also by boiling MoS_3 with $\text{K}_2\text{MoS}_4\text{Aq}$, washing the pp of MoS_3 and K_2MoS_4 with cold water, dissolving in boiling water, and adding acid to this solution (Berzelius) MoS_4 is a cinnamon-brown powder, partially oxidised in moist air With basic sulphides forms *perthiomolybdates* M_2MoS_5 (*v* THIOMOLYBDIC ACIDS AND THEIR SALTS, *infra*)

Molybdenum, sulpho-acids of, *v* Molybdenum, thio acids of, and their salts

Molybdenum, thio-acids of, and their salts A number of molybdates are known in which O is partly or wholly replaced by S These salts may be divided into two classes, those containing both O and S , these may be called *oxy-thiomolybdates*, and those containing only S , these may be called *thiomolybdates* The latter salts are classed as thiomolybdates, derived from MoS_3 , and perthiomolybdates, derived from MoS_4 The only thio acid of Mo which has been isolated with certainty is H_2MoS_3

PERTHIOMOLYBDIC ACID, H_2MoS_4 A red pp obtained by adding a weak acid to K_2MoS_4 (which separates during the preparation of MoS_3), washing with cold water, and drying over H_2SO_4 (Kruss, *A* 225, 1) Insol water, alcohol, and ether Sol warm NaOHAq , not acted on by NH_4Aq , evolves H_2S slowly when boiled with conc HClAq

OXY-MONOTHIOMOLYBDATES, $\text{M}_2\text{MoO}_3\text{S}$ The Na salt is prepared by melting pure MoO_3 with dry Na_2CO_3 , till CO_2 is all expelled, powdering the $\text{Na}_2\text{Mo}_2\text{O}_7$, thus obtained, dissolving in freshly-prepared NaHSaQ , withdrawing the reddish heavy oil which separates, washing the crystals which are formed by allowing this oil to stand, with alcohol, ether, CS_2 , and then again with alcohol and ether, and drying over H_2SO_4

(Krüss, *A* 225, 1). A hygroscopic salt, sol water

OXY-DITHIOMOLYBDATES, $\text{M}_2\text{MoO}_2\text{S}_2$ The NH_4 salt is prepared, as golden-yellow needles, by passing H_2S into NH_4 molybdate dissolved in NH_4Aq , at c 6° (Krüss, *lc*) The K salt is also described by Krüss (*lc*)

OXY-PYRODITHIOMOLYBDATES, $\text{M}_2\text{Mo}_2\text{O}_2\text{S}_2$ ($= 2\text{M}_2\text{MoO}_2\text{S}_2 - \text{M}_2\text{S}$) The salts of this series at present known are acid salts The NH_4 salt, $\text{NH}_4\text{HMo}_2\text{O}_2\text{S}_2$, is prepared, as a reddish-yellow pp, by adding NH_4HS drop by drop to NH_4 molybdate at c 90° , filtering after addition of NH_4Cl , washing with alcohol, and drying over H_2SO_4 Corresponding K and Na salts are described by Krüss (*lc*)

TETRA-BASIC DIOXY DITHIOMOLYBDATES The salt $\text{K}_2\text{O} \cdot 3\text{K}_2\text{S} \cdot 2\text{MoO}_3 \cdot 2\text{MoS}_3$ ($= \text{K}_4\text{Mo}_2\text{S}_8\text{O}_{10}$) was obtained by Krüss (*lc*) by adding alcohol to a solution of $\text{K}_2\text{Mo}_2\text{O}_7$, repeatedly shaking the oil which forms with alcohol, separating the black flaky crystals from the yellow crystals and the brownish red powder, all of which are produced on standing, washing and drying

THIOMOLYBDATES, M_2MoS_3 These salts are derived from the hypothetical thiomolybdic acid H MoS_3 , of which acid MoS_3 is the anhydride The soluble thiomolybdates are obtained by directly combining MoS_3 with basic sulphides, also by passing H_2S into conc solutions of the corresponding molybdates, the insoluble thio molybdates are obtained by ppg salt solutions by soluble thiomolybdates The soluble thiomolybdates may generally be crystallised, the crystals show fluorescence, green in reflected, red in transmitted, light Solutions of these salts are red The thiomolybdates as a class are decomposed by heat, giving a mixture of sulphide of the basic metal and MoS_2 , the K salt is only partly decomposed at white heat Conc solutions are stable in presence of excess of MoS_3 , when diluted, and especially in presence of free alkali, the solutions become turbid with formation of thiosulphates and thiomolybdates richer in S than the original salts Boiled in absence of air, the solutions evolve H_2S and form MoS_2 and perthiomolybdates, M_2MoS_5 (Berzelius, *P* 7, 261, Krüss, *A* 225, 1, cf Bodenstein, *J* pr 78, 186)

Potassium thiomolybdate, K_2MoS_3 Prepared by gradually heating, finally to a high temperature, a mixture of equal parts K_2CO_3 and S , with a little C , and a large excess of MoS_3 , dissolving in water, filtering, and crystallising (Hirzel, *J* 1850 319) Krüss (*lc*) obtained this salt by dissolving 5 g K_2MoO_4 in 16 cc water, adding 50 cc KOHAc , S 1 46, and saturating with H_2S Red, rhombic, crystals, with greenish fluorescence Sol water, acetic acid ppts $\text{K}_2\text{Mo}_2\text{S}_5$

The other thiomolybdates which have been examined, more or less fully, are the salts of NH_4 , Ba , Cd , Ca , Ce , Cr , Co , Cu , Au , Fe , Li , Mg , Mn , Hg , Ag , Na , Sr , Sn , U , and Zn

PERTHIOMOLYBDATES, M_2MoS_5 The soluble salts of this class are formed by boiling thiomolybdates M_2MoS_3 with MoS_3 , the insoluble salts by ppg salt solutions by soluble perthiomolybdates The perthiomolybdates are generally dark red, solutions of the alkali salts give

a pp of MoS_3 on addition of acids (Berzelius, *P* 7, 261)

Potassium perthiomolybdate, K_2MoS_6 . Prepared by saturating an acid K molybdate solution with H_2S (a mixture of K_2MoS_4 and MoS_3 is thus produced), boiling for some hours in a retort, cooling, collecting the pp of K_2MoS_6 mixed with MoS_3 , washing with water till the washings give a flocculent dark red pp on addition of HClAq , extracting with boiling water, and evaporating the solution. Also by heating dilute $\text{K}_2\text{MoS}_4\text{aq}$ with excess of MoS_3 , nearly to 100° till the liquid is nearly dry. Also by reacting on MoS_3 with KHSAq . A red mass, resembling an extract, on one occasion Berzelius obtained ruby-red crystalline granules. Insol cold water, sol hot water. Scarcely acted on by cold HClAq or KOHaq .

The other perthiomolybdates examined, more or less fully, are the salts of NH_4 , Ba, Bi, Ca, Au, Fe, Hg, Ag, Na, Sr, and Sn.

TRIANGULAR DITHIOMOLYBDATES Krüss (*A* 225, 1) obtained the salt $\text{K}_2\text{Mo}_2\text{S}_6$ ($=3\text{K}_2\text{S} \cdot 2\text{MoS}_3$) = $\text{K}_2\text{MoS}_3 \cdot \text{MoS}_3 \cdot 2\text{K}_2\text{S}$ by passing H_2S into $\text{K}_2\text{MoO}_4\text{aq}$ containing KOHaq , $\text{SG} > 1.46$ (*cf* Potassium thiomolybdate, *supra*). Small redish orange rhombic crystals, sol water, insol alcohol and ether. Decomposes slowly in air, giving K_2MoS_3 . M M P M.

MONARDA OIL The essential oil of *Monarda punctata*, which smells like thyme, contains thymol $\text{C}_{10}\text{H}_{14}\text{O}$ [48°] and a terpene (Arrpe, *A* 58, 41).

MONO This prefix is very seldom used in this dictionary. Compounds whose names begin with mono are described under the name to which mono has been prefixed.

MORIN or MORIC ACID $\text{C}_{15}\text{H}_8\text{O}_4\text{aq}$ (B a H), $\text{C}_{15}\text{H}_{10}\text{O}_4\text{aq}$ (from alcohol), $\text{C}_{15}\text{H}_{10}\text{O}_4 \cdot 2\text{aq}$ (from water) (L). S 0.25 at 20° , 0.94 at 100° . Occurs in old fustic (*Morus tinctoria*) to which it imparts its dyeing properties (Wagner, *J pr* 51, 82, Hlasiwetz a Pfandler, *J pr* 90, 445, 94, 65, *A* 127, 352, Löwe, *Z* 1875, 117, *Fr* 14, 119, Benedikt a Hazura, *M* 5, 165, 667).

Preparation—1 Deposited as calcium salt from an infusion of the wood on cooling. This deposit is extracted with alcohol, water is added to the alcoholic extract when calcium morate is ppd while maclurin remains in solution. The calcium salt is then decomposed by oxalic acid (Wagner)—2 The deposit is washed, heated with very dilute HClAq , freed from CaCl_2 by washing, and repeatedly dissolved in alcohol and ppd by water (H a P, B a H).

Properties—Needles, v sl sol water, v sol alcohol, m sol ether, insol CS_2 . The solutions are deep yellow. Blackens at 300° . Conc H_2SO_4 dissolves it, without change, forming a brownish yellow solution. Its alkaline solutions are deep yellow. Its solution does not ppt gelatin. It stains the skin yellow. FeCl_3 colours the alcoholic solution deep olive green. It reduces Fehling's solution on warming and ammoniacal AgNO_3 in the cold. Its alcoholic solution exhibits green fluorescence on the addition of an aluminium salt (Goppelsröder, *J pr* 101, 406, 104, 10, *Z* [2] 4, 154, 607). Morin yields resorcin on dry distillation.

Reactions—1 Sodium-amalgam turns the alkaline solution indigo blue and finally yellow,

the solution then containing phloroglucin—2. Potash-fuson yields phloroglucin and resorcin. 3 Nitric acid in HOAc oxidises it to (4, 2, 1)-di-oxy-benzoic acid.—4 Bromine added to its alcoholic solution forms ethyl-tetra-bromomorn $\text{C}_{15}\text{H}_2\text{EtBr}_4\text{O}$, 4aq which is ppd on adding water and melts at 135° . Tin and HCl convert this ether into tetra-bromo-morn $\text{C}_{15}\text{H}_2\text{Br}_4\text{O}$, 2aq which becomes anhydrous at 110° and then melts at 258° . It dyes silk and wool yellow without a mordant.

Salts— $\text{KC}_2\text{H}_3\text{O}_4$, yellow needles (from $\text{K}_2\text{CO}_3\text{aq}$)— $\text{Ca}(\text{C}_2\text{H}_3\text{O}_4)_2$, yellow precipitate $\text{Ba}(\text{C}_2\text{H}_3\text{O}_4)_2$, red-brown powder— $\text{Zn}(\text{C}_2\text{H}_3\text{O}_4)_2$, lemon-yellow needles, insol water, sol alkalis— $\text{PbC}_2\text{H}_3\text{O}_4$, egg yellow pp— $\text{Pb}_2\text{C}_2\text{H}_3\text{O}_4$ (L).

Morin sulphonic acid $\text{C}_{15}\text{H}_8\text{O}_4 \cdot \text{SO}_3\text{H}$, 2aq. Formed by heating morin with conc H_2SO_4 . Golden brown powder, sl sol. cold, v sol hot, water— $\text{K}_2\text{C}_2\text{H}_3\text{SO}_4$, 1aq golden needles— $\text{BaC}_2\text{H}_3\text{SO}_4$, Golden flocculent pp, converted by nitric acid into tri-nitro phloroglucin.

Isomorn Formed by adding sodium-amalgam to an alcoholic solution of morin containing HCl until the solution is deep-purple, and then evaporating (H a P). Purple-red prisms. When heated alone or in alcoholic solution, or more quickly by treatment with alkalis, it is reconverted into ordinary morin. Its solution mixed with dilute alum is dichroic.

Paramorn $\text{C}_{15}\text{H}_8\text{O}_4$. Obtained, together with a larger quantity of resorcin, by distilling morin mixed with sand (Benedikt, *B* 8, 606). Yellowish, woolly needles (from ether). Tasteless. May be sublimed. Reduces Fehling's solution. V sol hot water and ether (unlike morin). It dissolves unchanged in conc H_2SO_4 . Alkalis form a yellow solution. Unlike morin, its alcoholic solution is not ppd by $\text{Pb}(\text{OAc})_2$.

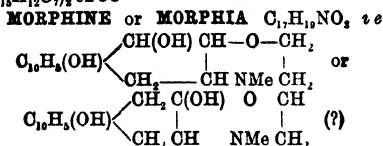
MORINDIN $\text{C}_{22}\text{H}_{16}\text{O}_{15}$ (A) [245°] (Stein). Extracted by alcohol from the root of various species of *morinda* used as a dye ('Suranyi') in India (Anderson, *Tr E* 16 [6], 435, *A* 71, 216). Slender lustrous orange needles, when heated it gives off orange vapours which condense to red needles of morindone. It is insol ether, v sl sol cold water and alcohol, sol aqueous KOH and conc H_2SO_4 , forming reddish violet solutions. Gives a red lake with alum, and a cobalt blue pp with baryta water. Morindin resembles ruberythric acid $\text{C}_{22}\text{H}_{16}\text{O}_{14}$ (Roehlede, *Sitz B* 7, 806, Stenhouse, *C* 17, 333) but differs from it in being insoluble in ether and in the behaviour of the red alkaline solution on boiling, for this becomes deep purple in the case of ruberythric acid, but does not change in that of morindin (Stein, *J pr* 97, 234, Thorpe a. Greenall, *C* 51, 52, 53, 171).

Morindone $\text{C}_{15}\text{H}_{10}\text{O}_4$. Tri-oxy methyl-anthraquinone (?). Formed together with glucose by boiling morindin with dilute mineral acids. Red needles, resembling alizarin. Insol water, v sol alcohol and ether. Its solution in potash is reddish-purple, but fades on standing (unlike alizarin). Conc H_2SO_4 gives a dark-blue solution (alizarin gives an orange solution). FeCl_3 gives a green colour. On distillation with zinc-dust it yields methyl anthracene.

MORINGIC ACID An acid supposed by Walter to exist as a glyceride in oil of ben (from

Moringa aptera) but shown by Zaleski (*B* 7, 1013) to be oleic acid

MORITANNIC ACID $C_{15}H_{12}O_7$ (?) If, after the aqueous extract of fustic has deposited morin, the filtrate be shaken with $EtOAc$, the acetic ether evaporated, and the residue dissolved in cold water and pptd by $NaCl$, brown amorphous moritannic acid separates first, followed by maclurin (Löwe, *Fr* 14, 127) V sol water. The solution is pptd by albumen, gelatin, and alkaloids $FeCl_3$ gives a brownish-black pp. Alcoholic lead acetate gives a pp ($C_{15}H_{12}O_7$), 5PbO



(Knorr, *B* 22, 1117) Mol w 285 [230°] (Hesse, *Ph* [3] 18, 801) S G 1.32 (Schröder, *B* 13, 1074) S 01 at 10°, 04 at 40°, 22 at 100° (Chastaing, *Bl* [2] 37, 477) S (absolute alcohol) 1.132 at 10.6°, 8.623 at 78° S (alcohol of 90 p.c.) 3.77 at 10.6°, 2.991 at 78° S (amyl alcohol) 2.68 at 11°, 2.247 at 78° (Florio, *G* 13, 496) S (ether) 0.23 at 10° (F) S (chloroform) 0.4 at 9.4°, 1.235 at 56° (F) S (amyl alcohol) 2.61 in the cold (Kubly, *J* 1866, 823) The solubility of crystallised morphine in boiling liquids is given by Prescott (*Ph* [3] 6, 404) as follows S (ether) 0.163, S (chloroform) 0.23, S (amyl alcohol) 1.1, S (benzene) 0.112 The solubility of amorphous morphine in boiling liquids is given by Prescott as S (ether) 0.473, S (chloroform) 0.506 The solubility of freshly pptd morphine in boiling solutions is given by Prescott as S (ether) 0.94, S (chloroform) 1.16, S (amyl alcohol) 1.1, S (benzene) 0.5 According to Burg (*Fr* 19, 222), the solubility of morphine in pure chloroform is 0.1, and in a mixture of 9 pts chloroform and 1 pt alcohol it is 9 [a]_D (in a 2 p.c. solution containing 2 mols $NaOH$) = -70.23° (Hesse, *A* 176, 190) [a]_D (in a solution of the hydrochloride) = -100.67° - 1.14*p*, where *p* = percentage of salt present (Hesse, Grimbert, *J* *Ph* [5] 16, 295)

Occurs in opium, being known in an impure state as *Magisterium opii* in the seventeenth century, and was first isolated in 1816 by Sertürner (*Gül. Ann* 55, 61, 57, 192, 59, 50) Dott (*Pr E* 12, 189) found in a sample of opium that half the morphine was present as meconate and half as sulphate It also occurs in *Eschscholtzia californica* (Baudet & Adrian, *C. C* 1889, 197), and in hops

Preparation—1 Opium is macerated with water at 38°, the extract evaporated with $CaCO_3$ to a small bulk, and then mixed with $CaCl_2$. The liquid is filtered from pptd calcium meconate and evaporated. It first deposits calcium meconate, and afterwards a mixture of the hydrochlorides of morphine and codeine. These crystals are dissolved in water and treated with ammonia, which ppt morphine, leaving codeine in solution (Robiquet & Gregory, *A* 5, 87, 7, 261)—2 The aqueous extract of opium is boiled with lime, and the filtrate boiled with NH_4Cl , which ppt morphine (Mohr, *A* 35, 120)

Properties—Small trimetric prisms (containing aq), v sl sol water, alcohol, and ether. It has a very bitter taste, and is a powerful narcotic poison. It dissolves readily to the extent of one equivalent in aqueous KOH , lime, baryta, and $NaOH$, but scarcely at all in ammonia and alkaline carbonates. It readily dissolves in acids. It is levorotatory. It loses its water of crystallisation at 100° (Dott, *Ph* [3] 18, 701), and begins to sublime at 150° in dots (Blyth)

Reactions—1 Very readily oxidised, thus 0.14 p.c. of the base is sufficient to liberate iodine from iodic acid (Serullas, *B J* 11, 238) Morphine is oxidised by alkaline K_2FeO_4 to pseudomorphine $C_{17}H_{17}N_2O_6$ (Hesse, *A* 235, 229), and the same substance is formed by the action of nitrous acid on morphine (Schutzenberger, *A* 108, 346, *Bl* 4, 176), and by allowing morphine to stand in aqueous NH_3 exposed to the air (Polstorff, *B* 13, 86) Morphine slowly reduces silver nitrate in the cold—2 On oxidation with alkaline $KMnO_4$, it yields an acid, apparently pyridine tricarboxylic acid (Barth & Weidel, *M* 4, 700)—3 Potash fusion yields protocatechuic acid (B & M) and methylamine (Wertheim, *A* 73, 208) Alcoholic potash at 180° forms methyl ethyl amine (Skraup & Wiegmann, *M* 10, 110)—4 Nitrous acid forms nitroso morphine, pseudomorphine, and a base $C_{17}H_{19}NO_3$ —5 Nitric acid, diluted with ($\frac{1}{2}$ mols of) water, forms at 100° an acid $C_{16}H_{15}NO_3$, which can be converted by conc HNO_3 into picric acid, and which yields methylamine when heated with KOH (Chastaing, *C R* 94, 44, *J Ph* [5] 4, 338)—6 Converted into codeine by heating with $NaOEt$ and MeI the yield being small owing to the formation of morphine methylo iodide (Grimaux, *A Ch* [5] 27, 273, Dott, *Ph* [3] 12, 1009)—7 On distillation with zinc dust it yields pyridine, NH_3 , pyrrole, and a base which may be quinoline, as chief products, together with 4 p.c. of phenanthrene and $\frac{1}{2}$ p.c. of phenanthraquinoline (Gerichten & Schrotter, *A* 210, 396)—8 When evaporated with dilute H_2SO_4 , till white fumes appear, morphine yields a residue turned brown by alkalis. Morphine (1 pt), heated with H_2SO_4 ($\frac{1}{2}$ pts) and oxalic acid (2 pts), at 120° forms a product which, on diluting with water, leaves a yellowish white amorphous body undissolved. This substance, $C_{17}H_{19}NO_3$, is soluble in hot water and in alkalis. It turns green in air, and its alkaline solution turns red in air. The redened alkaline solution gives with HCl a blue pp, 'morphine blue', $C_{17}H_{19}NO_3$, which can be crystallised from ether. Malonic and succinic acids behave like oxalic acids, forming the compounds $C_{17}H_{19}NO_5$ and $C_{17}H_{19}NO_6$. All these formulae should perhaps be doubled (Chastaing & Barillot, *C R* 105, 941, 1012)—9 Heating with oxalic acid forms trimorphine—10 Heating with conc HCl yields apomorphine (Matthiessen & Wright, *Pr* 17, 455, 460), and the bases $C_{17}H_{17}ClN_2O_6$, $C_{17}H_{17}ClN_2O_7$, $C_{17}H_{17}ClN_2O_8$, and $C_{17}H_{17}Cl_2N_2O_8$ —11 With $ClCH_2OAc$ and H_2SO_4 it gives an intense violet colour, and after dilution with water and addition of NH_3 , an amorphous yellow base, possibly $CH_3(C_{17}H_{17}NO_3)_2$, which gives a violet colour with H_2SO_4 (Grimaux, *C R* 93, 271)—12 Aqueous *alloxan* and SO_2 form $C_{17}H_{17}N_2O_7$, $C_{17}H_{17}NO_8$, H_2SO_4 (Pelizzari, *A* 248, 161)—13 Bleaching-powder, acting on a solution

of morphine hydrochloride, forms $C_{17}H_{19}ClNO_3$, a crystalline body (Mayer, *B* 4, 121)

Colour tests—1 Liberates iodine from iodic acid solution The brown colour is increased by adding ammonia (Lefert, *J Ph* [3] 40, 97). If a layer of very dilute ammonia be poured upon a very dilute solution of morphine containing iodic acid and starch, a brown ring below a blue ring will appear at the junction of the liquids (Dupré, *C N* 8, 267)—2 *Chloride of gold* colours the solution blue, being reduced to gold—3 $KMnO_4$ is reduced, becoming green—4 *Ferric salts* give a blue colour, destroyed by acids, by heat, and by alcohol (Robinet a Pelletier, *A* 5, 172)—5 $FeCl_3$, added to a previously heated solution of morphia in H_2SO_4 , gives a deep red colour, changing to dirty green (Husemann, *A* 128, 305) According to Lindo (*C N* 37, 158), morphine, gently warmed with H_2SO_4 and $FeCl_3$, gives an indigo blue colour—6 A solution of *molybdic acid* in conc H_2SO_4 gives, with solid morphine, a violet colour, changing through blue to dirty green (Frohde, *Fr* 5, 214)—7 A solution obtained by adding ammonia to cupric sulphate till the pp cupric hydrate is just re dissolved is coloured green by salts of morphine (Nadler, *Fr* 13, 235, Lindo, *Fr* 19, 359)—8 Nitric acid gives an orange red colour, gradually changing to yellow—9 H_2SO_4 , containing a little nitric acid, gives a violet red solution (Erdmann, *A* 120, 188, Husemann, *A* 128, 305)—10 When morphine is boiled with aqueous phosphorous or hypophosphorous acid, and conc H_2SO_4 is added, the liquid becomes reddish violet On now adding a little nitric acid the solution becomes yellowish, and on warming with copper it assumes a splendid blue colour (Dragendorff, *J* 1864, 727)—11 $KClO_4$ (free from chlorate), added to morphine mixed with H_2SO_4 , gives a dark brown colour (Grove, *Fr* 13, 324, Siebold, *ibid*)—12 Conc H_2SO_4 , followed by water and some oxidising agent (iodic acid, K_2FeCy_4 , K_2CrO_4 , KNO_3 , MnO_2 , or PbO), best in the solid state, gives a deep red colour on gentle warming (Lindo, *C N* 36, 228, Dott, *C N* 36, 255)—13 A mixture of K_2FeCy_4 , ferric chloride, and HCl is coloured blue This reagent may be used to detect morphine in quinine salts (Kieffer, *A* 103, 271, Hager, *Fr* 12, 220, Armitage, *Ph* 18, 761) According to Hesse (*Ph* 18, 801), the blue colour is due to pseudomorphine, not to Prussian blue—14 Chlorine water and ammonia give a red colour in a 1 p.c solution (Flückiger, *Ar Ph* [3] 1, 117)—15 Aqueous $K_2Cr_2O_7$ gives a dirty brown pp (Plugge, *Ar Ph* [3] 25, 793)—16 Conc H_2SO_4 , containing a little K_2AsO_4 , gives on warming a blue violet colour, changing to dark brown red Water turns this green, and chloroform on shaking with it becomes violet, while ether becomes violet red, the liquid below it being brown (Tattersall, *C N* 41, 63, Donath, *J pr* [2] 33, 563)—17 H_2SO_4 and a little $KClO_4$ gives in the cold a grass green colour, the edge of the liquid being pale rose colour (Vitali, *B* 14, 1583)—18 A mixture of morphine (1 pt) and powdered sugar (7 pts) turns purple on addition of a drop of H_2SO_4 , the mass gradually changing in colour through bluish green to yellow (Schneider, *J* 1872, 747)—19 If morphine is heated with H_2SO_4 at 200° , and the

mass poured into water, the liquid becomes blue If chloroform be shaken with this liquid it acquires a blue colour, while if ether be used the ethereal layer becomes purple (Jorisson, *Fr* 20, 422)—20 On warming morphine with H_2SO_4 and sodium phosphate a violet colour appears (Vulpus, *Ar Ph* [3] 25, 256)

Estimation in Opium—Opium (200 pts) is exhausted with warm water and the extract evaporated to a thin syrup, mixed with alcohol (40 pts of S G 82), ether (380 pts), and ammonia (45 pts of S G 935) The mixture is occasionally shaken, and after 18 hours it is filtered and washed with alcohol-ether The pp is dried, digested with benzene, dried, and weighed After weighing the base may be titrated with standard HCl (Teschemacher a Smith, *C N* 57, 93, 103, 244, Flückiger, *J* 1885, 1962, *Ar Ph* [3] 27, 721, Venturini, *G* 16, 239, Stillwell, *Am* 8, 307, R Williams, *C N* 57, 134)

Another method consists in exhausting 7 g of opium with 110 c.c of water and 4 g of slaked lime 50 c.c of the filtrate are mixed with 5 c.c of 95 p.c alcohol, 25 c.c of ether, and 3 g of NH_4Cl After standing for 12 hours the ether is filtered off, and another 10 c.c of ether added The entire liquid is then filtered and the morphine collected on the filter determined by direct weighing or by titration (Goebel, *Chem Zeit* 11, 508) The morphine may also be extracted from the opium by baryta water and ppd by CO_2 (Von Perger, *J pr* [2] 29, 97) Indeed, a great many processes for estimating morphine in opium have, from time to time, been proposed (Guillemont, *J Ph* [3] 16, 17, [4] 6, 102, Riegel, *Jahrb pr Pharm* 23, 202, Gumbourt, *J Ph* [3] 41, 97, 177, F Mayer, *Am J Pharm*, 35, 28, Arnoldi, *Russ Zeit Pharm* 1873, 641, Lynn, *Amer J Pharm* [4] 6, 358, *C J* 34, 612, Flewry, *J Ph* [4] 6, 99, Schachtrupp, *Fr* 7, 284, 509, Merck, *A* 18, 79, 21, 202, 24, 46, Steen, *Ar Ph* [2] 148, 150, *C J* 25, 180, J T Miller, *Ph* [3] 2, 465, *C J* 25, 181, Bevel, *Monit Scient* 13, 812, *J* 1871, 821, Kremel, *C C* 1887, 1529, Adrian a Gallois, *J Ph* [5] 15, 193, Schlikum, *Ar Ph* [3] 25, 13, Dott, *Ph* [3] 18, 82, Plugge, *Ar Ph* [3] 25, 343)

Methods for separating morphine from other alkaloids have been given by Dragendorff (*Ph*, [3] 7, 24), Hager (*C C* 1872, 727), Neubauer (*Fr* 9, 134), and others

Salts— $B'HCl$ 3aq S 5 in the cold, 100 at 100° (Regnault, *A Ch* [2] 68, 131, *A* 26, 24), 4 at 15° (Hesse, *A* 179, 190) $[a]_D = -94^\circ$ in a 2 p.c solution containing excess (10 mols) of HCl Silky fibres (from water)— $B'HCl$ S (MeOH) 2 at 15° Minute prisms (from MeOH) The salt $B'HCl$ 3aq is partially rendered anhydrous by solution in alcohol (Hesse, *A* 203, 151)— $B'H_2PtCl_4$, yellow curdy pp, m sol water (Liebig, *A* 26, 46) Decomposed by boiling water (O de Coninck, *Bl* [2] 45, 131)— $B'H_2PtCl_4$ 6aq (Hesse)— $B'HCl(HgCl_2)$ Crystalline pp (Hinterberger, *A* 77, 201)— $B'ZnCl_2$ 2aq— $B'ZnCl_2$ 7aq (Graffinghoff, *Z* 1866, 600)— $B'HEr$ 2aq needles (Schmidt, *B* 10, 194)— $B'HI$ 2aq long silky needles grouped in rosettes, sl sol cold water (Schmidt, *B* 10, 194)— $B'HI$ 3aq (Bauer, *Ar Ph* [3] 5, 289)— $B'HI$ 3aq hair like needles (Kunz, *Ar Ph* [3]

26, 807) — B_2I_2 , crystalline mass (B) — BHI , obtained by ppg a salt of morphine with iodine dissolved in $HIAq$ Brown pp, sol $KIAq$, from which it crystallises in nearly black prisms, sol alcohol and boiling ether (Bauer, Jørgensen, *J pr* [2] 2, 487, *Z* [2] 5, 673) — $BHHgI_2$, crystalline pp, m sol alcohol (Groves, *C J* 11, 97) — $B_2H_2SO_4$, 5aq [a] = -100.47 - 96p in a p p c solution at 22.5° (Hesse) *S* 4 in the cold (Dott) — $BHClO_2$, 2aq needles (Bödeker, *A* 71, 88) — $BHVO$, (Ditte, *A Ch* [6] 13, 237) — $BHCyS$ 4aq [100°] Needles (Dollfus, *A* 65, 214) — $B_2H_2PtO_2$, groups of minute needles (Schwarzenbach, *C C* 1860, 304) — $BHOAc$ *S* 50 in the cold (Dott) Tufts of needles, v sol water, m sol alcohol (Merck, *A* 24, 48) — $B'CHCl$, CO_2H (Florio, *J* 1833, 1343) — $B'CHCl$, CO_2H 4aq — $B'CCl$, CO_2H 14aq — Butyrate $B'C_4H_7O_2$ *S* 14 at 12° (Decharme, *J* 1863, 444) — Valerate $B'C_5H_9O_2$, Large trimetric crystals (Pasteur, *A Ch* [3] 58, 455) *S* 28 in the cold (Dott) — Lactate $B'C_3H_5O_2$, monoclinic crystals *S* 10 at 13° (Decharme) — Trichloro-lactate $B'CCl_3H_5O_2$, 5aq (F) — Oxalate $B'_2C_2H_2O_4$, aq prisms *S* 48 at 12° (D) — Tartrates — $B'_2C_2H_2O_4$, 3aq Nodular groups of needles, efflorescent at 20° (Arppe, *J pr* 3, 332) *S* 11 in the cold (Dott) — $B'_2C_2H_2O_4$, 3aq Tufts of long rectangular flattened prisms (A) — Mellitate $B'_2C_2H_2O_4$, minute needles (Karmrod, *A* 81, 171) — Phenyl acetate $B'CH_2Ph$, CO_2H [92°] — Nitroprusside (Davy, *Ph* [3] 11, 756)

Metallic derivatives $C_17H_{15}KNO_2$, 24aq Obtained in a crystalline state by evaporating a solution of morphine in $KOHAq$ in a desiccator over H_2SO_4 (Chastaing, *J Ph* [5] 4, 19) — $(C_17H_{15}KNO_2)_2K_2CO_3$, 2aq — $Ba(C_17H_{15}NO_2)_2$, 3aq white crystalline solid — $Ca(C_17H_{15}NO_2)_2$, 5aq — $Ca(C_17H_{15}NO_2)_2$, 2aq

(a) **Acetyl derivative** $C_17H_{15}AcNO_2$ Obtained, together with the (8) modification, by boiling morphine (1 pt) with $HOAc$ (2 pts) for several hours (Wright, *C J* 27, 1033, Beckett a Wright, *C J* 28, 315) Crystallises from ether, either with aq or in the anhydrous state Gives no colour with $FeCl_3$ — $B'HCl$ 3aq crystals, sl sol cold water — $B_2H_2PtCl_4$ amorphous — $B'EtI$ 4aq crystals (from 85 p c alcohol)

(8) **Acetyl derivative** $C_17H_{15}AcNO_2$ The chief product obtained by heating morphine (1 mol) with acetic anhydride (1 mol) at 100° When less acetic anhydride is used a compound $(C_17H_{15}AcNO_2)(C_17H_{15}NO_2)$ is obtained (Wright) Amorphous — $B'HCl$ amorphous, v sol water Readily converted into morphine by boiling water, and hence usually gives a blue colour with $FeCl_3$ — $B'EtI$ amorphous

(7) **Acetyl derivative** $C_17H_{15}AcNO_2$ Accompanies the (8)-isomeride to the extent of 25 p c Anhydrous crystals (from ether) — $B'HCl$ crystallises with difficulty, v sol water — $B'EtI$ 4aq crystals (from 85 p c alcohol)

Di-acetyl derivative $C_{17}H_{15}Ac_2NO_2$ [169°] Formed by heating morphine with excess of Ac_2O at 85° (Wright, Hesse, *A* 222, 205) The product is treated with water and dilute NH_4aq , and extracted with ether Colourless prisms (from ether), v sol cold alcohol, m sol ether Gives no colour with $FeCl_3$ Converted by boiling water first into the (a) acetyl deriva-

tive, and then into morphine — $B'HCl$ crystals, extremely sol water — $B_2H_2PtCl_4$ — $B'MeCl$ Formed by heating morphine methylchloride with Ac_2O at 85° (Hesse) Needles, v e sol water — $B'MeI$ Heated with $AgOAs$ it yields the crystalline di-acetyl derivative of a compound $C_{17}H_{15}O_2$ [143°] (Fischer, *B* 19, 792) — $B'EtI$ 4aq Crystals (B a W)

Di-propionyl derivative

$C_{17}H_{15}(C_2H_3O)_2NO_2$ Formed by heating morphine with propionic anhydride at 85° (Hesse, *A* 222, 207) White amorphous powder, v e sol ether, alcohol, and chloroform, sl sol water $FeCl_3$ gives no colouration — $B'HCl$ — $B_2H_2PtCl_4$

(a) **Butyryl derivative** $C_{17}H_{15}(C_4H_7O)_2NO_2$ Formed by boiling morphine (1 pt) with butyric acid (2 pts) for 6 hours (Beckett a Wright, *C J* 28, 16) Crystals (from ether) Gives no colour with $FeCl_3$ — $B'HCl$ more soluble and less readily crystallisable than the hydrochloride of the (a) acetyl derivative — $B_2H_2PtCl_4$ — $B'EtI$ amorphous

(8) Butyryl derivative

$C_{17}H_{15}(C_4H_7O)_2NO_2$ An amorphous base, accompanying the preceding isomeride Gives a blue colour with $FeCl_3$

Di butyryl derivative $C_{17}H_{15}NO_2$, *ie* $C_{17}H_{15}(C_4H_7O)_2NO_2$ Formed by heating morphine (1 pt) with butyric anhydride (2 pts) for 3 hours at 140° (B a W) Amorphous Resolved by boiling water or boiling dilute alcohol into butyric acid and butyryl morphine — $B'HCl$ amorphous Gives no blue colour with $FeCl_3$ — $B'EtI$ amorphous By boiling morphine with a mixture of acetic and butyric acids, Beckett and Wright obtained a product which formed a crystalline double salt with the formula $(C_{17}H_{15}AcNO_2)(C_{17}H_{15}(C_4H_7O)_2NO_2)H_2Cl_2$, 8aq, resolved into its components by boiling water

Benzoyl derivative $C_{17}H_{15}BzNO_2$ Formed by heating morphine with benzoic acid at 160° (B a W), and by boiling the di benzoyl derivative with dilute alcohol Amorphous — $B'HCl$ crystalline, v sl sol water Not coloured by $FeCl_3$

Di benzoyl derivative $C_{17}H_{15}Bz_2NO_2$ [188°-191° cor] Formed from morphine and Bz_2O at 130° (B a W) or $BzCl$ at 105° (Wright a Rennie, *C J* 37, 609, Dott a Stockman, *Ph* [3] 18, 231, cf Brookmann a Polstorff, *B* 13, 96) Prisms, insol water, sol hot alcohol Gives no colour with $FeCl_3$ — $B'HCl$ amorphous — $B_2H_2PtCl_4$ — $B'EtI$ 4aq crystals

Acetyl benzoyl derivative

$C_{17}H_{15}AcBzNO_2$ Formed from the (a) acetyl derivative and Bz_2O at 130° (B a W) Crystalline — $B'HCl$ amorphous, v sol water — $B_2H_2PtCl_4$ — $B'EtI$ 4aq crystals (from alcohol).

Succinozoyl derivative

$(C_{17}H_{15}NO_2)CO C_2H_5CO_2H$ Formed by heating morphine (1 pt) with succinic acid (2 pts) at 180° (Beckett a Wright, *C J* 28, 689) Separates from alcohol in crystals (containing 4aq), insol water and ether — $B_2H_2PtCl_4$

Methylo-chloride $C_{17}H_{15}NO_2MeI$ 4aq Formed by warming morphine with MeI and alcohol (How, *C J* 6, 125, *A* 88, 338) If $NaOEt$ is also present the product is codeine (Grimaux, *A Ch* [5] 27, 273, *C R* 92, 1140)

Methylo-chloride $C_{17}H_{15}NO_2MeCl$ 2aq.

Formed from the methylo-iodide and AgCl (Hesse, A 222, 207) Long needles Its aqueous solution is coloured blue by FeCl_3 , Ac_2O forms $\text{C}_1\text{H}_7\text{I}_2\text{Ac}_2\text{NO}_2\text{MeCl} - (\text{C}_1\text{H}_7\text{NO}_2\text{MeCl})_2\text{PtCl}_4\text{aq}$

Methylo-hydroxide $\text{C}_1\text{H}_7\text{NO}_2\text{MeOH} \cdot 5\text{aq}$ Formed from the sulphate and baryta (Broockmann a Polstorff, B 13, 96) Colourless needles Reduces Ag_2O

Ethylo-iodide $\text{C}_1\text{H}_7\text{NO}_2\text{EtI} \cdot 3\text{aq}$ Formed by heating morphine with EtI and alcohol at 100° (How) Needles (from water).

Methyl-morphine v CODEINE

Di-methyl-morphine $\text{C}_1\text{H}_7\text{Me}_2\text{NO}$, *Methylmorphumethine* [118°] $[\alpha]_D^{20} = -208.6^\circ$ in a 4 pc solution in 97 pc alcohol at 16° Formed by heating codeine methylo-iodide (1 mol.) with aqueous KOH (1 mol), adding benzene, and shaking with acetic acid The acid liquid is saturated with NaCl, and the hydrochloride of di methyl morphine then crystallises out The free base is obtained by adding NaOH and extracting with ether (Hesse, A 222, 220) Prisms, in sol ether, v sol water and alcohol Conc H_2SO_4 gives a bluish violet colour Conc HNO_3 forms a yellow solution

Salts — $\text{B}^+\text{HCl} \cdot 2\text{aq}$ S 92 at 18° — $\text{B}_2\text{H}_2\text{PtCl}_6\text{aq}$

Methylo - hydroxide

$\text{C}_1\text{H}_7\text{Me}_2\text{NO}_2\text{MeOH}$ Formed from the iodide and moist Ag_2O (Knarr, B 22, 181) Thick syrup Split up at 160° into NMe, and the phenanthrene derivative $\text{C}_1\text{H}_7\text{EtO}_2$, Ac_2O at 160° – 200° forms acetyl methyl dioxypheanthrene $\text{C}_1\text{H}_7(\text{OMe})(\text{OAc})$, dimethylamine, and di methyl oxyethyl amine NMe $\text{CH}_2\text{CH}_2\text{OH}$ (Fischer a Von Gerichten, B 19, 794)

(a) *Methylo iodide* $\text{C}_1\text{H}_7\text{Me}_2\text{NO}_2\text{MeI} \cdot 3\text{aq}$ Formed from di methyl morphine and MeI dissolved in alcohol (Hesse, A 222, 225) Prisms v sol hot water

(b) *Methylo - iodide* $\text{C}_1\text{H}_7\text{Me}_2\text{NO}_2\text{MeI}$ Obtained by adding KOH to a solution of the (a) isomeride, allowing the oily pp to crystallise, and adding KI to the mother liquor (Hesse, A 222, 227) Crystals

(a)-*Methylo chloride*

$\text{C}_1\text{H}_7\text{Me}_2\text{NO}_2\text{MeCl} \cdot 3\text{aq}$ Obtained from the (a) methylo iodide by treatment with AgCl With Ac_2O it forms $\text{C}_1\text{H}_7\text{AcMe}_2\text{NO}_2 \cdot 2\text{aq}$ crystallising in needles v sol hot water and alcohol — $(\text{B}^+\text{MeCl})_2\text{PtCl}_4\text{aq}$

(b) *Methylo chloride* $\text{C}_1\text{H}_7\text{Me}_2\text{NO}_2\text{MeCl}$ Formed from the (b) methylo iodide and AgCl Forms a neutral sulphate and a platinumchloride

Acetyl derivative $\text{C}_1\text{H}_7\text{AcMe}_2\text{NO}_2$

[66°] Formed from di methyl morphine and Ac_2O at 85° (Hesse, A 222, 223) V sol alcohol and ether, m sol water, insol KOH aq Gives a blue colour with conc H_2SO_4 — $\text{B}^+\text{HCl} \cdot 3\text{aq}$ — $\text{B}^+\text{H}_2\text{PtCl}_6\text{aq}$ — $\text{B}^+\text{HNO}_3\text{aq}$ — $\text{B}^+\text{H}_2\text{SO}_4\text{aq}$

Ethyl-morphine $\text{C}_1\text{H}_7\text{EtNO}_2$, *Codeithine* [83°] S $2\frac{1}{2}$ at 100° Formed by heating morphine (1 mol.) with NaOEt (1 mol.), alcohol, and EtI (Grimaux, C R 92, 1140, 1228, A Ch. [5] 27, 278) Hard brilliant laminae (containing aq), v sol alcohol and ether, m sol water Ppd from solutions of its salts by KOH and Na_2CO_3 , but not by NH_4aq Gives a blue colour with H_2SO_4 containing FeCl_3 — B^+HCl groups of slender needles — $(\text{C}_1\text{H}_7\text{EtNO}_2)_2\text{MeI}$ Hard

bulky crystals, converted by moist Ag_2O into a tertiary base [132°]

Di-ethyl-morphine $\text{C}_1\text{H}_7\text{Et}_2\text{NO}_2$ The iodide of this base appears to be formed by heating morphine with NaOH (1 mol.) and EtI (2 mols.) It is crystalline, and is converted by successive treatment with Ag_2O and MeI into $\text{C}_1\text{H}_7\text{Et}_2\text{NO}_2\text{MeI}$, which crystallises from alcohol, and is converted by Ag_2O into an hydroxide which is split up on distillation into NMeEtPr and a derivative of phenanthraquinone $\text{C}_1\text{H}_7\text{EtO}_2$, (Gerichten a Schrötter, B 15, 2182)

Ethylene di-morphine $\text{C}_1\text{H}_7(\text{C}_1\text{H}_7\text{NO}_2)_2$, *Di-codethine* *Dicodethylene* Prepared by boiling morphine with alcoholic soda and ethylene bromide (Grimaux, C R. 93, 67, A Ch. [5] 27, 281) White needles (from alcohol), insol ether, v sol alcohol Blackens when heated, and decomposes above 200° With H_2SO_4 and FeCl_3 it gives a blue colouration — $\text{B}^+\text{H}_2\text{Cl}_2$ Hard prisms

Morphinyl-sulphuric acid

$\text{C}_1\text{H}_7(\text{OSO}_3\text{H})\text{NO}_2 \cdot 2\text{aq}$ *Morphine sulphonic acid* Formed by shaking morphine (20 g) with KOH (8 g), water (25 cc), and $\text{K}_2\text{S}_2\text{O}_8$ (15 g) (Stolnikoff, H 8, 242) Silvery needles (from water), not decomposed at 160° v sl sol cold water, alcohol, and ether Hot dilute HCl aq hydrolyses it to morphine and H_2SO_4 Does not give a blue colour with FeCl_3 Hot conc H_2SO_4 gives a rose colour, changing to violet Less poisonous than morphine, producing tetanus

Nitroso-morphine $\text{C}_1\text{H}_7(\text{NO})\text{NO}_2 \cdot \text{aq}$ Formed by passing nitrous acid gas into morphine suspended in water (E L Mayer, B 4, 121) Reddish yellow powder, turned black by FeCl_3

Trimorphine $(\text{C}_1\text{H}_7\text{NO}_2)_3$ Formed by heating morphine (30 g) with H_2SO_4 (30 cc) and water (30 cc) at 100° (Wright, C J 26, 220) Formed also by heating morphine (1 pt) with dry oxalic acid (3 pts) at 145° (Beckett a Wright, C J 28, 698) Amorphous, sol ether Gives a purple colour with FeCl_3 — $\text{B}^+\text{H}_2\text{Cl}_2$ Amorphous Converted by conc hydrochloric acid at 100° into $\text{C}_{10}\text{H}_{11}\text{Cl N}_3\text{O}_6\text{HCl}$, a brittle gum

Tetramorphine $(\text{C}_1\text{H}_7\text{NO}_2)_4$ Formed, together with trimorphine, by heating morphine with dilute H_2SO_4 at 100° – 140° (Wright, C J 26, 220) Amorphous On heating with HCl aq it forms the salt $(\text{C}_1\text{H}_7\text{NO}_2)_4(\text{HCl})_4$ (?) On adding HCl to an aqueous solution of tetra morphine a flocculent pp of $(\text{C}_1\text{H}_7\text{NO}_2\text{HCl})_4$ is got — $(\text{C}_1\text{H}_7\text{NO}_2)_2\text{H}_2\text{SO}_4$, *Sulphomorphine*. Obtained by heating morphine with dilute H_2SO_4 at 160° (Arppe, A 55, 96, Matthiessen a Wright, Pr 17, 455, Wright, C J 26, 220)

Apomorphine $\text{C}_1\text{H}_7\text{NO}_2$ Formed by heating morphine or codeine with HCl aq at 145° for 2 or 3 hours (Matthiessen a Wright, Pr 17, 455) Formed also by heating morphine with aqueous phosphoric acid till the temperature reaches 185° (Wright, C J 25, 652) It is also one of the products when morphine is heated with aqueous ZnCl_2 for 20 minutes at 125° (Mayer a Wright, C J 26, 211) Amorphous mass, which turns green in moist air Sl sol water, especially if it contains CO_2 , sol alcohol, ether, and chloroform Emetic, 0.2 g of the hydrochloride being sufficient to produce vomiting. A solution of apomorphine hydrochloride gives with caustic alkalis a white pp, quickly becoming black, with Na_2CO_3 a white pp, turning green, with

HNO_3 , a blood-red colour, with FeCl_3 an amethyst colour, with $\text{K}_2\text{Cr}_2\text{O}_7$ an orange pp, with KI an amorphous pp, quickly becoming green, and with platinum chloride a yellow pp, decomposing on warming. It reduces AgNO_3 . AcCl yields a mono acetyl derivative (Danckworth, *Ar Ph* 228, 572) — B^+HCl crystals, sl sol cold water

Polymeride of apomorphine $\text{C}_{138}\text{H}_{138}\text{N}_6\text{O}_{18}$ (?) Formed, together with apomorphine and several bases, which form the hydrochlorides $\text{C}_{138}\text{H}_{138}\text{ClN}_6\text{O}_{18}$, $\text{C}_{138}\text{H}_{138}\text{N}_6\text{O}_{18}\text{H}_2\text{Cl}_2$ (?), and $\text{C}_{138}\text{H}_{138}\text{Cl}_2\text{N}_6\text{O}_{18}\text{H}_2\text{Cl}_2$ (?) by heating morphine with aqueous ZnCl_2 . It forms a hydrochloride $\text{B}^+\text{H}_2\text{HCl}$, insol alcohol (Mayer & Wright)

Diapotetramorphine $\text{C}_{12}\text{H}_{12}\text{N}_2\text{O}_2$ Formed, together with apomorphine, by heating morphine with aqueous phosphoric acid (Wright) Amorphous, rapidly darkening in air. Evaporation with conc hydrochloric acid converts it into $\text{C}_{12}\text{H}_{12}\text{Cl}_2\text{N}_2\text{O}_2 \cdot 8\text{HCl}$. Evaporation with HIAq and phosphorus yields the corresponding $\text{C}_{12}\text{H}_{12}\text{N}_2\text{O}_2 \cdot 8\text{HI}$ (Wright)

Pseudomorphine $\text{C}_8\text{H}_8\text{N}_2\text{O}_2$ $1\frac{1}{2}$ aq or $\text{C}_8\text{H}_8\text{N}_2\text{O}_2 \cdot 8\text{aq}$ (Hesse, *A* 235, 229) **Dehydromorphine Oxymorphine Oxymorphine Dymorphine** Discovered by Pelletier (1832) in Levant opium. Occurs in morphine hydrochloride made by Gregory's method (Hesse, *A* 141, 87)

Formation — 1 By heating a solution of morphine hydrochloride with AgNO_3 at 60° (Schutzenberger, *Bl* [2] 4, 176) — 2 By oxidising morphine with KMnO_4 , potassium nitrite, or K_2FeCy_6 , or by exposing its ammoniacal solution to the air (Polstorff, *B* 13, 86, cf Pelletier, *A* 16, 49)

Properties — White, microcrystalline powder (from NH_4Aq), insol water, alcohol, ether, and chloroform, sol KOH aq and NH_4aq , v e sol alcoholic NH_3 . Lævrotatory (Hesse, *A* 176, 195). Decomposed by heat without melting. It is not affected by reducing agents. It is tasteless and not poisonous

Colour tests — 1 When evaporated with dilute H_2SO_4 until fumes of H_2SO_4 appear it becomes bluish green, and, on adding water, rose-red, if HNO_3 , or dilute NaNO_2 , be now added the colour changes to deep violet. Under similar circumstances morphia gives a rose red colour, turned brownish red by water, and raspberry red by HNO_3 (Donath, *J pr* [2] 33, 560) — 2 Resembles morphia in its reaction with FeCl_3 , with molybdic acid in H_2SO_4 (Froehde's reagent), with HNO_3 , and with HIO_3 . For the reaction with iodic acid HCl must be absent. NH_4Cl interferes with these tests — 3 Conc H_2SO_4 forms an olive green colour — 4 Conc HNO_3 gives an intense orange red colour, changing to yellow — 5 A mixture of equal parts of cane sugar and pseudomorphine is coloured by H_2SO_4 blue, changing to dark green (Hesse, *A* 234, 255)

Salts — $\text{B}^+\text{HCl aq}$ Minute needles $\text{S } 14$ at 20° [α_D] = $-103^\circ 13'$. — $\text{B}^+\text{HCl} 2\text{aq}$ — $\text{B}^+\text{HCl} 3\text{aq}$ — $\text{B}^+\text{HCl} 4\text{aq}$ — $\text{B}^+\text{HCl} 6\text{aq}$ — $\text{B}^+\text{HCl} 8\text{aq}$ — $\text{B}^+\text{H}_2\text{PtCl}_6\text{aq}$ — B^+HIAq — $\text{B}^+\text{H}_2\text{SO}_4\text{aq}$ small colourless needles, sl sol cold water (Polstorff) — $\text{B}^+\text{H}_2\text{SO}_4\text{aq}$ laminae $\text{S } 287$ at 20° . — $\text{B}^+\text{H}_2\text{Cr}_2\text{O}_7\text{aq}$ crystalline pp $\text{S } 1$ at 18° . — $\text{B}^+\text{H}_2\text{C}_2\text{O}_4\text{aq}$ $\text{S } 05$ at 20° . — $\text{B}^+\text{H}_2\text{C}_2\text{O}_4\text{aq}$ — Tartrate $\text{B}^+\text{C}_2\text{H}_3\text{O}_4\text{aq}$ needles — $\text{B}^+\text{C}_2\text{H}_3\text{O}_4\text{aq}$. $\text{S } 24$ at 18° .

Acetyl derivative $\text{C}_{17}\text{H}_{15}\text{Ac}_2\text{NO}_2$ [276°]

Formed by heating pseudomorphine (1 pt) with Ac_2O (2 pts) at 120° (Hesse, *A* 222, 235, 234, 255) Crystallises from ether in prisms (containing 4 aq) V e sol alcohol, m sol ether and chloroform. When mixed with cane sugar and dissolved in conc H_2SO_4 , it gives a dark green colour. If the H_2SO_4 contain $\text{Fe}_2(\text{SO}_4)_3$, the colour is at first blue — $\text{B}^+\text{H}_2\text{PtCl}_6\text{aq}$

Methylo-iodide $\text{C}_{17}\text{H}_{15}\text{NO}_2\text{MeI}$ 2 aq Formed by the action of HI on the crystalline body $(\text{C}_{17}\text{H}_{15}\text{NO}_2)_2\text{Me}(\text{OH})\text{MeI}$, which is prepared by oxidising morphine methylo iodide with alkaline K_2FeCy_6 (Polstorff, *B* 13, 93) Small prisms

Methylo-hydroxide

$\text{C}_{17}\text{H}_{15}\text{NO}_2\text{Me}(\text{OH})$ 8½ aq Prepared by the action of moist Ag_2O on the iodide or of baryta on the sulphate Crystalline, v sol water, insol alcohol

Methylo sulphate

$(\text{C}_{17}\text{H}_{15}\text{NO}_2)_2\text{Me}_2\text{SO}$ 4 aq Obtained by ppg a solution of the oxyiodide with Ag_2SO_4 . Leaflets, v sol hot water

MORPHOTHEBAINE v THEBAINE

MORPHOTROPY This term is applied to the relations between the crystalline form and the composition of those bodies which are chemically similar, and which are derived from the same parent body, v this vol p 89

MORRHUIC ACID $\text{C}_{18}\text{H}_{31}\text{NO}_2$, 1 e

$\text{CH} \begin{array}{c} \text{CH}(\text{OH}) \\ \text{CH}_2 \text{NH} \end{array} \text{C}_8\text{H}_7\text{CO}_2\text{H}$ (?) **Oxypropyl pyridine dihydride carboxylic acid** Occurs in cod-liver oil in a combination with glycerin and phosphoric acid, from which it is readily set free by acids and alkalis. Obtained by extracting the oil with dilute (85 pc) alcohol containing 5 pc of HCl (Gautier & Mourgues, *C R* 107, 740) Square flattened prisms or large lanceolate plates, insol cold, sol hot, water, v sol alcohol, ether, and alkalis. The solutions have a disagreeable aromatic odour (Gautier, *Bl* [3] 2, 233) Morrhucic acid decomposes carbonates. A solution of the potassium salt gives pps with lead acetate and AgNO_3 , but not with cupric acetate. The silver salt is readily reduced, even in the cold. Morrhucic acid forms a crystalline platinumchloride, an amorphous aurochloride, and a hydrochloride which is decomposed by boiling water. On distilling with lime it yields a basic oil which forms a crystalline methylo iodide. Potassium morrhuate is oxidised by KMnO_4 to an acid which ppts cupric acetate, and appears to be a pyridine derivative

MORRHUINE $\text{C}_{18}\text{H}_{31}\text{N}$, An alkaloid occurring together with aselline $\text{C}_{18}\text{H}_{31}\text{N}$, in cod-liver oil (Gautier & Mourgues, *Bl* [3] 2, 228, *C R* 107, 626) Thick oil, smelling like *syrringa*, v sl sol water, v sol alcohol and ether. Lighter than water. Ppts copper salts. Strongly alkaline. It forms a very deliquescent hydrochloride crystallising in groups of needles, a crystalline platinumchloride and a yellow aurochloride. Morrhucine constitutes one third of the total bases in the oil, and is a diaphoretic and diuretic

MORTARS v CEMENTS, in DICTIONARY OF APPLIED CHEMISTRY

MOSAIC GOLD An alloy of Cu and Zn in equal parts, v COPPER ALLOYS OF, vol 11 p 254, v also DICTIONARY OF APPLIED CHEMISTRY.

MUCEDIN v PROTEIDS.

MUCIC ACID $C_6H_8O_7$, *tc*

$CO_2H \cdot CH(OH) \cdot CH(OH) \cdot CH(OH) \cdot CH(OH) \cdot CO_2H$
 Mol w 210 [206°–216°] (K & T) S 33 at 14°, 17 at 100° Formed by the oxidation of milk sugar, galactose, melitose, dulcitol, gum arabic, gum tragacanth, and plant mucus with dilute nitric acid (Scheele, *Opuscul* 2, 111, Laugier, *A Ch* 41, 79, Berzelius, *A Ch* 92, 141, 94, 5, 95, 31, Malaguti, *A Ch* [2] 60, 195, 63, 86, Liebig & Pelouze, *A* 19, 258, Liebig, *A* 9, 24, 26, 160, Hagen, *P* 71, 531, *A* 64, 347, Johnson, *A* 94, 225, Schwanert, *A* 116, 257, Tollens, *A* 249, 220)

Preparation —1 Milk sugar (1 pt) is heated with HNO_3 (2 pts of S G 1 4) and water (2 pts) until red fumes appear. The flame is then removed until the reaction has abated. The liquid is then evaporated, a further quantity (4 pt) of HNO_3 being added during the evaporation. The product is washed with water and dried (Klinkhardt, *J pr* [2] 25, 43). The yield is 35 p.c., cf Guckelberger, *A* 74, 348 —2 Coarsely powdered milk sugar (100 g) is heated with HNO_3 (1200 c.c. of S G 1 15) in a basin on a water bath until the volume is greatly reduced (to 150 or 200 c.c.) After cooling, water (200 c.c.) is added, and after some days the mucic acid is filtered off and washed with water (500 c.c.) The yield is good (40 g). When galactose is treated in the same way the yield is 77 p.c. (Kent & Tollens, *A* 227, 221)

Properties —Colourless tables, sl sol cold water, insol alcohol. It does not reduce Fehling's solution.

Reactions —1 When boiled with water it changes to the isomeric paramucic acid —2 When heated with water at 180° it yields pyromucic acid —3 On dry distillation it yields CO_2 , pyromucic acid and some diphenylene oxide. When cautiously heated at 280° it forms a small quantity of dehydro mucic acid $C_6H_4O(CO_2H)_2$, which condenses on the neck of the retort as small crystals —4 Boiling nitric acid oxidises it to oxalic and racemic acids —5 On oxidation with H_2SO_4 and MnO formic acid is given off —6 Potash fusion yields acetic and oxalic acids —7 $HIAg$ and phosphorus at 140° forms some adipic acid (Crum Brown, *A* 125, 19), and diphenylene oxide (Heinzelmann, *A* 193, 186) —8 PCl_5 forms di chloro muconic chloride $C_6H_4Cl_2O$ (Lièbs Bodart, *A* 100, 325, Bode & Wichelhaus, *A* 132, 95) —9 When allowed to ferment for 9 months it yields alcohol and acetic, butyric, and traces of lactic acids (Béchamp, *Bl* [3] 3, 770) —10 Ba.S at 205° forms thiophene (8)-carboxylic acid —11 Fuming $HClAq$ (or $HBrAq$) at 100°–150° forms dehydro mucic acid.

Salts — $(NH_4)_2A$ flat four sided prisms (from water), sl sol cold, v sol hot, water. Decomposed at 220°–240° into pyrrole, pyrrole carboxylic amide, NH_3 , and CO_2 — NH_4HA aq needles or thin prisms. More soluble in water than the normal salt — Na_2A 4½aq transparent triethinic crystals — Na_3A ½aq white powder — $NaHA$ 3½aq prisms — K_2A ½aq granules S 125 Insol alcohol — K_2A 2½aq (Schmidt & Cöbenzl, *B* 17, 601) — KHA aq decomposes at 150°–180° into CO_2 and pyromucate — BaA 1½aq (dried at 100°) — CaA 1½aq — MgA 2½aq (at 100°) — PbA aq white pp — $Pb_2C_6H_4O_7$: ob-

tained by ppg ammonium mucate with basic lead acetate — CuA aq (dried at 100°) bluish-white pp — FeA 2½aq (dried at 100°) yellow powder — $K(SbO)A$ — $Na(SbO)A$ (dried at 100°) Obtained by dissolving Sb_2O_3 in $NaHA$ Amorphous (Klein, *C R* 97, 1437) — AgA (dried at 100°) white pp —Methylamine salt Decomposed by distillation into methyl pyrrole and methyl pyrrole carboxylic methylamide —Ethylamine salt Decomposed by distillation into ethyl pyrrole, ethyl pyrrole dicarboxylic diethylamide, and ethyl pyrrole carboxylic ethylamide (Bell, *B* 10, 1866) —Diethylamine salt Decomposed by heat giving off NH_3 —Aniline salt $(C_6H_5N)_2HA$ Insol boiling alcohol, sol boiling water (Kottnitz, *J pr* [2] 6, 138) Decomposed on distillation into phenyl pyrrole, CO_2 , aniline, and H_2O On heating with water it forms mucic anilide.

Methyl ether Me_2A Laminæ or flattened six sided prisms (from water), v sol hot water, v sl sol hot alcohol. Decomposes at 165°

Mono-ethyl ether $EtHA$ Formed by boiling Et_2A with alcohol (Lumprecht, *A* 165, 255). Crystals (containing 3aq), sol water and alcohol. Melts below 100° — NH_4EtA

Di-ethyl ether Et_2A [158°] S 23 at 20° S (alcohol of S G 814) 64 at 15° Formed by heating mucic acid (1 pt) with H_2SO_4 (4 pts) till it turns black, leaving it to cool, and adding alcohol (4 pts) (Malaguti, *A Ch* [2] 63, 86) Crystals (from boiling alcohol) Lumprecht was not able to obtain this ether by the action of alcohol and HCl on mucic acid. Mucic ether is reduced by sodium amalgam to an acid which strongly reduces Fehling's solution, and which may be readily re-oxidised to mucic acid (E Fischer, *B* 23, 937)

Isoamyl ether $(C_4H_9)_2A$ Needles

Amide $C_6H_8O_6(NH_2)_2$ Mucamide S G 155 1589 Formed from the normal ether and ammonia (Malaguti, *C R* 22, 854) Formed also by the action of ammonia on tetra acetyl mucic amide (Ruhemann, *B* 20, 3366) Minute crystals (from water), v sl sol boiling water, insol alcohol and ether. Converted by water at 140° into ammonium mucate. On dry distillation it yields pyrrole carboxylic amide, paracyanogen, and a little pyromucic acid.

Anilide $C_6H_4O_6(NHPh)_2$ Formed by heating the aniline salt at 120°, or the ether with excess of aniline (Kottnitz) Plates, insol water, alcohol, and dilute acids

o-Toluide $C_6H_3O_6(NHC_6H_4)_2$ Formed like the preceding (K) Plates

Tetra-acetyl derivative $C_6H_4(OAc)_4(CO_2H)_2$ [266° cor] Formed by boiling mucic acid with Ac_2O and $ZnCl_2$ (Maquenne, *Bl* [2] 48, 720) Efflorescent needles (containing 2aq), sl sol water, v sol alcohol

Ethyl ether of the tetra acetyl derivative $C_6H_4(OAc)_4(CO_2Et)_2$ [177°] S (95 p.c. alcohol) 4 at 17° Formed by heating mucic ether with $AcCl$ at 100° (Werigo, *A* 129, 195) Needles, sl sol ether, v sol boiling alcohol, v sl sol boiling water

Di-phenylhydrazide $C_6H_4(OH)_2(CO N_2H_5Ph)_2$ [240°] Formed by heating mucic acid with phenyl hydrazine hydrochloride and $NaOAc$ on the water-bath (Bulow, *A* 236, 196, Maquenne, *Bl* [2] 48,

722) Pale yellow plates (from phenyl-hydrazine), v sl sol. water, alcohol, and ether Crystallises unaltered from alcoholic potash

Paramucic acid S 136 in the cold, 58 at 100° Formed by evaporating an aqueous solution of mucic acid to dryness, dissolving the residue in alcohol, and allowing the alcoholic solution to evaporate in the air (Malaguti, *A* 15, 179) Crystalline, more soluble in water than mucic acid On recrystallisation from hot water it changes into mucic acid Paramucates, with the exception of the normal ammonium salt, are more soluble than the corresponding mucates, but their boiling solutions deposit mucates

Dehydromucic acid $C_6H_6O_5$

Formation—1 By heating mucic acid in sealed tubes with $HBrAq$ (saturated at 0°) at 100° for two days The product may be crystallised from water and separated from unaltered mucic acid by crystallisation of the Ba salt (Heinzelmann, *A* 193, 184)—2 By heating mucic acid with fuming $HClAq$ at 145° (Seelg, *B* 12, 1033)—3 In small quantity by heating mucic acid at 180° (Klinkhardt)—4 By heating hydrogen potassium saccharate with conc $HClAq$ at 150° (Sohst a Tollens, *A* 245, 19)

Preparation—Mucic acid (1 pt), conc HCl (1 pt), and conc HBr (1 pt) are heated 8 hours at 150° in sealed tubes The solid contents of the tubes are collected and distilled with steam If this steam be condensed it is found to contain di-phenylene oxide The residue is neutralised by ammonia, filtered, and mixed with HCl The dehydro mucic acid is then ppd Yield 20 p c (Klinkhardt, *J pr* 133, 44)

Properties—Plates (from alcohol) or needles (from hot water) Does not melt at 320° V sl sol cold water, alcohol, and ether May be sublimed

Reactions—1 $FeCl_3$ produces, especially on warming, in a solution of dehydro-mucic acid, a transparent gelatinous pp This result is prevented by the presence of other acids—2 **Bromine water** converts it into fumaric acid, the reaction being $C_6H_6O_5(CO_2H)_2 + 3Br_2 + 3H_2O = C_4H_4(CO_2H)_2 + 2CO_2 + 6HBr$ —3 A mixture of HNO_3 and H_2SO_4 converts it into nitropyromucic acid (*q v*)—4 **Sodium amalgam** reduces it to two acids of the formula $C_6H_6O_5$, melting at 146° and 173° (Graebe a Bungener, *B* 12, 1079) The (α) acid [146°] crystallises in thin plates and forms the crystalline salts $CaA'' 3\frac{1}{2}aq$, $BaA'' 4\frac{1}{2}aq$, and $AgA'' \frac{1}{2}aq$ The (β)-acid [173°] forms large crystals (containing *aq*) and the salts $CaA'' 1\frac{1}{2}aq$ (amorphous), $BaA'' 1\frac{1}{2}aq$ (needles) and AgA'' —5 On **dry distillation** it splits up into CO_2 and pyromucic acid (*q v*)

Salts— $BaA'' 2\frac{1}{2}aq$ — $BaA'' 6aq$ Sol water— $CaA'' 8aq$ — AgA'' white pp, decomposed by boiling water

Ethylether EtA'' [47°]

Chloride $C_6H_6O_5(COCl)_2$ [80°] (α 245°) From PCl_5 on the acid (Klinkhardt, *J pr* 133, 46) Smells like $POCl_3$ Sol ether, alcohol, and $CHCl_3$ At 100° it sublimes forming flat needles Boiling water reconverts it into dehydro-mucic acid

Amide $C_6H_6O_5(CONH_2)_2$ Formed by the action of NH_3 on an ethereal solution of the chloride. Slender needles (from water) Nearly

insol cold water, alcohol, and ether Does not melt below 240°

MUCILAGE The gum of plants from which it is obtained by steeping these in hot water, which on cooling forms a jelly When linseed mucilage is strained through linen, and the filtrate mixed with alcohol and $HClAq$, a pp is obtained which after washing with alcohol and ether has the composition $C_6H_{10}O_5$ (Schmidt, *A* 51, 50, Frank, *J pr* 95, 484, Nageli a Cramer, *Pharm Cent* 1855, 426, Kirchner a Tollens, *A* 175, 215) Brittle gum, resembling bassorin (*q v*), sol cold water, insol alcohol Dextrogyrate Insoluble in ammoniacal cupric oxide Boiling dilute H_2SO_4 partially converts it into glucose

Quince mucilage is coloured blue by iodine, and gives no mucic acid on oxidation with nitric acid Boiling dilute H_2SO_4 yields a mixture of cellulose, gum, and sugar The mucilage from salep (*Orchis mascula*), from *Tamarindus indica*, and from some other plants, is also turned blue by iodine (*v* STARCH) Quince mucilage yields furfuraldehyde on distillation with dilute H_2SO_4 , indicating the presence of arabinose or xylose It contains neither glucose nor galactose, since neither saccharic nor mucic acid is formed on oxidation

Salep mucilage yields no furfuraldehyde on distillation with dilute H_2SO_4 , but on oxidation it yields saccharic (but not mucic) acid It contains glucose and mannose, but no galactose or arabinose (Gans a Tollens, *A* 249, 245)

MUCIN *v* PROTEIDS, Appendix C

MUCOBROMIC ACID $C_6H_6BrO_5$

CHO CBr CBr CO H Semi-aldehyde of di bromo-maleic acid [121°] Formed by adding bromine to pyromucic acid covered with a little water without cooling The product is boiled and evaporated, the yield being 70 p c of the theoretical (Schmelz a Beilstein, *A Suppl* 3, 276, Jackson a Hill, *B* 11, 1671 *Am* 3, 41) Formed also from (β)-di bromo pyromucic acid and hot dilute HNO_3 (Hill a Sanger, *A* 232, 89) and by oxidation of di bromo maleic aldehyde

Properties—White plates, v sol alcohol, ether, and hot water, v sl sol cold water Red dens litmus and decomposes carbonates, but its salts are very unstable

Reactions—1 When heated it partially sublimes, and the rest decomposes into HBr , di bromo maleic acid, and CO_2 (Hill, *Am* 3, 105)—2 Boiling baryta-water forms CO_2 , bromo-acetylene, malonic acid, and formic acid (Jackson a Hill, *B* 11, 289) Cold baryta water forms HBr and bromo maleic acid A cold paste of baryta and water forms bromo propionic acid and β di bromo acrylic acid Baryta water at 0° forms mucoxybromic acid (*q v*)—3 **Bromine water** forms di bromo maleic acid (Hill, *B* 13, 734) By heating with water and bromine (3 mols) in sealed tubes at 125° it is converted into penta and hexa bromo ethane, di bromo maleic acid, and tetra bromo butyric acid (Lumprecht, *A* 165, 293) Bromine (1 at) at 145° forms di-bromo maleic acid, mucobromyl bromide, and a little dibromo succinic acid—4 PBr_3 forms the bromide (*v infra*)—5 **Phenol** (25 g) dissolved in water (80 g) containing KOH (17.5 g) converts mucobromic acid (20 g) into 'phenoxy-mucobromic' acid $C_6H_5BrO_5$, which crystallises from water in prisms [105°] Phenoxy mucic

bromic acid forms the crystalline salts KA' and BaA' , 3aq, and is converted by Ag_2O in hot water into phenoxyl-bromo-maleic acid $\text{C}_6\text{H}_4\text{BrO}_2$, which crystallises in slender needles [104°], and forms a crystalline silver salt $\text{Ag}_2\text{A}'$ (Hill & Stevens, *Am* 6, 188) — 6 Moist Ag_2O oxidises it to di-bromo-maleic acid. Nitric acid acts in like manner — 7 Alcoholic potassium nitrite forms a pp of $\text{K}_2\text{C}_4\text{H}_4\text{N}_2\text{O}_6$, v sol cold water, converted by hot water into CO_2 , HCy , HNO_2 , and $\text{KC}_4\text{H}_4\text{NO}_6$. Bromine in CS_2 converts the salt $\text{K}_2\text{C}_4\text{H}_4\text{N}_2\text{O}_6$ into $\text{C}_4\text{HBr}_2\text{N}_2\text{O}_6$ (Hill & Sanger, *B* 15, 1906) — 8 KNO_2 at 50° forms the acid $\text{C}_4\text{H}_4\text{NO}_6$, which forms the crystalline salts NaA' aq, KA' aq, CaA' 4aq, BaA' 5aq, PbA' 4aq, CuA' , and AgA' .

Salts — BaA' white plates, sol cold water and alcohol — AgA' felted needles, insol water

Ethyl ether EtA' [51°] (255°–260°) Formed from the acid, alcohol, and H_2SO_4 . Large monoclinic crystals with aromatic odour. KNO_2 converts it into crystalline $\text{C}_4\text{H}_4\text{KNO}_6$.

Mucobromic acetic anhydride $\text{C}_4\text{HBr}_2\text{O}_6$ [54°] Formed by heating mucobromic acid with AcCl at 120°. Long needles, v e sol alcohol, ether, and chloroform.

Bromide $\text{C}_4\text{HBr}_2\text{O}_6$ [56°] Obtained by heating mucobromic acid (1 pt) with PBr_3 (4 pts) at 115°. Small slender prisms (from alcohol), sol alcohol, ether, benzene, chloroform, and CS_2 . Boiling water slowly reconverts it into mucobromic acid.

MUCOCHLORIC ACID $\text{C}_4\text{H}_2\text{Cl}_2\text{O}_6$, \pm e $\text{CHO Cl Cl CO}_2\text{H}$ or $\text{CHO Cl Cl C CO}_2\text{H}$ *Semi aldehyde of di-chloro-maleic acid* [125°] Formed by passing chlorine into a solution of pyromucic acid (1 pt) in water (10 pts) at 0° in presence of iodine, the yield being 40 p c of the theoretical (Beilstein & Schmelz, *A Suppl* 3, 276, Bennett & Hill, *B* 12, 655). Plates, v sol hot water, alcohol, and ether, insol ligroin and CS_2 . Split up by alkalis, even in the cold, into di-chloro-acrylic acid and formic acid.

MUCONIC ACID (of Limpricht) $\text{C}_4\text{H}_4\text{O}_6$, \pm e $\text{CO}_2\text{H CH}_2\text{C} \begin{smallmatrix} \text{CH CH}_2 \\ \text{O CO} \end{smallmatrix}$ [100°–125°] Formed

by adding silver oxide to a hot solution of di-bromo-adipic acid [175°–190°] (formed by adding bromine to a warm solution of hydromucic acid in HOAc), and decomposing the resulting silver salt with HCl or H_2S (Limpricht, *A* 165, 274). Large crystals, v sol water, alcohol, and ether. Boiling baryta water decomposes it into CO_2 , acetic acid, succinic acid, and another acid. Salts — BaA' 4aq nodules, v sol water.

Muconic acid $\text{CO}_2\text{H CH CH CH CO}_2\text{H}$ [above 260°] S 02 at 15°. Formed by the action of alcoholic potash on β -di-bromo-adipic acid (Rupe, *A* 256, 22, Ruhemann & Blackman, *C J* 57, 373). Branching white needles, sl sol. hot water, m sol hot alcohol and HOAc . Completely decomposed by KMnO_4 in presence of Na_2CO_3 . Combines with bromine forming the acid $\text{CO}_2\text{H CHBr CHBr CHBr CO}_2\text{H}$ [c 250°]. Sodium amalgam reduces it to hydro-mucic acid [195°].

Salts — KA' transparent plates, v sol water, insol alcohol. — BaA' — PbA' — $\text{Ag}_2\text{A}'$ curdy white pp. The cupric salt is a bluish green heavy amorphous pp.

Methyl ether $\text{Me}_2\text{A}'$ [154°] Fan shaped groups of needles.

Ethyl ether EtA' [64°] Plates (from dilute alcohol).

Di-chloro-mucic acid $\text{C}_6\text{H}_2\text{Cl}_2\text{O}_6$, \pm e $\text{CO}_2\text{H CH Cl Cl CH CO}_2\text{H}$ S 10 at 100°. Obtained by heating mucic acid or saccharic acid with PCl_5 and decomposing the resulting chloride with water (Liés Bodart, *A* 100, 325, Boëde, *A* 132, 95, Bell, *B* 12, 1272, Limpricht, *A* 165, 253, Rupe, *A* 256, 6). Long needles (containing 2aq); v sl sol cold water, v sol alcohol, m sol. ether. Not decomposed by boiling aqueous alkalis.

Reactions — 1 Reduced by sodium amalgam to two acids of the formula $\text{C}_4\text{H}_4\text{O}_6$, one of which melts at 195°, and yields malonic acid on oxidation, the other melts at 169°, and yields succinic acid on oxidation. Boiling with sodium amalgam reduces it to adipic acid [148°] — 2 Alcoholic potash at 190°–200° forms oxalic and acetic acids.

Salts — The Ba and Ca salts are m sol. water — $\text{Ag}_2\text{A}'$ insoluble pp.

Methyl ether $\text{Me}_2\text{A}'$ [156°] Pearly plates, v sol ether, hot alcohol, and HOAc .

Ethyl ether EtA' [96°] From the chloride and alcohol (Wichelhaus, *A* 135, 251, Bell, *B* 12, 1273). Prisms.

Chloride $\text{C}_6\text{H}_2\text{Cl}_2\text{O}_6$. Formed by the action of PCl_5 (6 mols) on mucic acid (1 mol). Large crystals (from CS_2), decomposed by moist air.

Hydromucic acid $\text{C}_4\text{H}_4\text{O}_6$, \pm e $\text{CO}_2\text{H CH}_2\text{CH CH CH}_2\text{CO}_2\text{H}$ [195°] S 58 at 15°. Formed by reducing di-chloro-mucic acid with sodium amalgam or with zinc dust and acetic acid. White needles or prisms. Converted by boiling NaOHAq into the isomeric acid [169°]. Oxidised by KMnO_4 to oxalic and acetic acids. Its Ba and Ca salts are less soluble in hot than in cold water. Bromine added to its aqueous solution forms bromohydromucic acid $\text{CO}_2\text{H CH}_2\text{CBr CH}_2\text{CO}_2\text{H}$ [183°], which yields an ether melting at 80°. Bromine without water forms unstable dibromoadipic acid of the formula $\text{CO}_2\text{H CH}_2\text{CHBr CHBr CH}_2\text{CO}_2\text{H}$.

Ethyl ether $\text{Et}_2\text{A}'$ (163° at 35 mm) Oil.

Amide $\text{C}_4\text{H}_4\text{O}_6(\text{NH}_2)_2$ [210°] (Ruhemann & Blackman, *C J* 57, 371).

Hydromucic acid $\text{C}_4\text{H}_4\text{O}_6$, \pm e $\text{CO}_2\text{H CH CH}_2\text{CH}_2\text{CH}_2\text{CO}_2\text{H}$ [169°] S 11 9 Formed by boiling the isomeric acid with aqueous NaOH (Rupe, *A* 256, 13). Nodular aggregates of plates, v sol hot, insol cold water, v sl sol ether. Yields oxalic and succinic acids on oxidation with KMnO_4 . Bromine forms a mono-bromo-derivative [160°] crystallising in branching needles, but no di-bromo-adipic acid.

Methyl ether Oil. Forms with bromine $\text{CO}_2\text{Me CH}_2\text{CH}_2\text{CHBr CHBr CO}_2\text{Me}$ [85°].

MUCOXYBROMIC ACID $\text{C}_4\text{H}_2\text{BrO}_6$, \pm e $\text{CO}_2\text{H C(OH)CBr CHO}$ [112°] Formed by the action of baryta water at 0° upon mucobromic acid (Hill & Palmer, *Am* 9, 148). Thick prisms, with bevelled ends, v sol water, alcohol, and ether, v sl sol chloroform, benzene, ligroin, and CS_2 . FeCl_3 gives an intense garnet red colour. AgNO_3 forms a white crystalline pp. On heating with baryta-water it yields oxalate and formate. Bromine water converts it into bromalhydrate.

CBr, CH(OH)₂ and oxalic acid. Aniline yields C₆H₅BrO₂(NPh)

Salts—K₂A'' aq plates, v sol cold water, decomposed on warming—BaA'' 2aq needles, sl sol cold water, decomposed on boiling—PbA'' yellow pp—Ag₂A'' crystalline pp, explodes on heating or on moistening with HNO₃

Methyl ether Me₂A'' Sticky liquid

Mono-ethyl ether EtHA'' [89°] Prisms, sol alcohol and ether

Di-ethyl ether Et₂A'' Viscous liquid

'Anilmucoxybromic' acid

CO₂H C(OH) CBr CH NPh [132°] Formed by the action of aniline hydrochloride on a dilute solution of mucoxybromic acid (Hill a Palmer, *Am* 9, 156) Pale yellow needles (containing aq), sl sol cold water, v sol alcohol, and ether FeCl₃ gives a brown pp Hot acids and alkalis set free aniline Phenyl hydrazine ppts the phenyl hydrazine salt (C₆H₅N)₂C₁₀H₇BrNO₃ aq

Salts—K₂A''—BaA'' 1aq bright yellow crystalline pp—Ag₂A'' bright orange yellow pp

MUCOXYCHLORIC ACID C₆H₅ClO₃ 1c

CO₂H C(OH) CCl CHO [115°] Formed from mucochloric acid by treatment with a very slight excess of baryta, added slowly at a low temperature (Hill a Palmer, *Am* 9, 159) The yield is 73 p.c. of the theoretical Stout prisms, v sol water, alcohol, and ether, v sl sol chloroform and benzene With FeCl₃ it gives a deep garnet red colour On heating with excess of baryta it yields oxalate and formate Bromine water forms oxalic acid and CClBr₂CHO Phenyl hydrazine yields an unstable condensation product Aniline produces anilmucoxychloric acid

Salts—K₂A'' small tables—BaA'' 2aq crystalline pp, sl sol cold water, v sl sol dilute alcohol—BaA'' aq—Ag₂A''

Mono-ethyl ether EtHA'' [95°] Prisms, which may be sublimed, sol water, alcohol, and ether, v sol boiling benzene

Di-ethyl ether Et₂A'' Viscous liquid

Anilmucoxychloric acid C₆H₅ClNO₃ 1c CO₂H C(OH) CCl CH NPh [147°] Formed from mucoxychloric acid and aniline Pale yellow needles (containing aq), becoming brilliant yellow when anhydrous, sl sol cold water, v sol alcohol and ether When it is heated with acids or alkalis aniline is split off FeCl₃ gives a deep brown pp

Salts—K₂A''—BaH₂A'' 1aq brightly yellow needles—Ag₂A'' bright orange pp—Phenylhydrazine salt PhN₂H₂H₂A'' aq white crystalline mass, sl sol cold water, v sol alcohol

MUCUS OF PLANTS v MUCILAGE

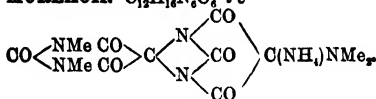
MUNJEET The root of *Rubia munjista*, used in India as a dye stuff If ground munjeet is boiled with a solution of aluminic sulphate, and the red liquid filtered and acidified with HCl, a yellow pp is formed If the pp be boiled with alcohol, pectic acid remains undissolved, and the alcohol contains purpurin and munjinstin They are separated by boiling dilute acetic acid, which dissolves munjinstin

Munjinstin is identical with purpuroxanthic acid, C₁₆H₁₂(CO₂H)₄ [231°] v Di oxy ANTHRA QUINONE CARBOXYLIC ACID (E Schunck a H Roemer, *O* J 33, 422, cf Stenhouse, *Pr* 12, 683, 13, 86, 145)

MUNTZ'S METAL An alloy of Cu and Zn, v. DICTIONARY OF APPLIED CHEMISTRY

MUREXIDE The ammonium hydrogen salt of PURPURIC ACID (q v)

MUREXOIN C₁₂H₈N₄O₂ 2c



Formed by converting caffeine into di methyl alloxan (Fischer, *B* 14, 1912), and reducing this by H₂S to tetra methyl alloxantin, which is then subjected to the action of air and ammonia (Brunn, *B* 21, 514, cf Rochleder, *J* pr 51, 405) Red prisms, sl sol water and alcohol Its aqueous solution is purple, resembling that of KMnO₄ Sublimes at about 230° Decolourised by potash (difference from murexide) Dilute HCl decomposes it, forming di methyl parabanic acid

MURIATIC ACID Synonymous with CHLORHYDRIC ACID, q v vol ii p 5

MURRAYIN C₁₈H₂₂O₁₀ [170°] A glucoside obtained from the flowers of *Murraya eratica* by extracting with water (Blas, *Z* 1869, 316, De Vrij, *Z* 1876, 850, Hoffmann, *Ar Ph* [3] 14, 139) White powder, composed of small needles, sl sol cold water, v sol boiling water and alcohol, insol ether Tastes bitter, but is not poisonous Its solutions in alkalis and in Na₂CO₃ exhibit a greenish blue fluorescence and turn brown on heating The aqueous solution is not pptd by salts of Cu or Hg FeCl₃ colours its solution blue Lead subacetate gives a pp It reduces ammoniacal AgNO₃, and Fehling's solution on warming Dilute acids split it up into glucose and murrayetin

Murrayetin C₁₂H₁₆O₈ [c 110°] Formed by boiling murrayin with dilute H₂SO₄ Prisms (containing 1/2 aq), sl sol cold, m sol boiling water, v sol alcohol, m sol ether Tasteless Its solutions exhibit strong greenish blue fluorescence, which is increased by KOH and by Na₂CO₃ FeCl₃ gives a bluish green colour Lead acetate gives, after a time, a yellow pp

MUSCARINE C₇H₁₁NO 1c CH(OH)₂ CH₂ NMe, OH An alkaloid occurring, together with neurine, in the fly agaric (*Agaricus muscarius*) (Schmiedeberg a Koppe, *J* 1870, 875, Ruckert, *N Rep Pharm* 21, 193), in the fungus *Amnaria Pantherina* (Giacosa, *J* 1883, 1488), and in putrefying flesh (Gautier, *Bl* [2] 48, 13, cf Brieger, *B* 17, 2741) Formed by oxidising neurine CH(OH)₂ CH₂ NMe, OH with conc HNO₃ (Schmiedeberg a Harnack, *C. C.* 1876, 554) Deliquescent tasteless crystals, v a. sol water and alcohol, insol ether Its solution is strongly alkaline, ppts ferric and cupric hydrates from solutions of their salts, and acts as a narcotic poison antagonistic to atropine Not affected by boiling dilute acids or alkalis Gives amorphous pps with mercury potassium iodide and bismuth-potassium iodide

Salts—(CH(OH)₂ CH₂ NMe, Cl), PtCl, 2aq—CH(OH)₂ CH₂ NMe, AuCl

Di-ethyl ether (HO)Me₂N CH₂ CH(OEt)₂ Formed by heating chloro-acetal CH₂Cl CH(OEt)₂ (from di-chloro-ether) with tri-methyl amine (Berlinerblau, *B* 17, 1139) By heating the compound C₆H₅NBr CH₂ CH(OEt)₂, formed by the action of bromo-acetal upon pyridine, with moist Ag₂O at 80° there is formed syrupy C₆H₅N(OH) CH₂ CH(OEt)₂, which is inappro

privately termed by Lochert (*Bl* [8] 8, 859) diethyl muscarine pyridine

MUSCLE Muscular tissue consists of fibres bound together into fasciculi by connective tissue. The fibres may be transversely striated (voluntary and cardiac muscle) or not (other in voluntary muscle). The plain or unstriated fibres are elongated cells with oval nuclei, and inclosed by a delicate envelope, they are singly refracting. The cardiac fibres have no sheath. The voluntary muscular fibres have a well marked sheath or sarcolemma, under which are situated numerous nuclei, the contents of the sarcolemma (muscle plasma) have a semi liquid consistency during life. Kuhne, and later Eberth, observed a nematode worm (*Myogryctes Weissmanni*) crawling up the interior of a muscular fibre (*Zeit f wiss Zool* 12, 530), the contents of the sarcolemma are not, however, homogeneous, this is denoted by the transverse striping, which is probably an optical effect produced by the presence of certain more solid structures which are described as rods, membranes, tubules, and networks of fibres by different observers, these solid bodies are isotropous and are suspended in an anisotropous (doubly refracting) viscous fluid. (For the various theories of the histological structure of striated muscle during rest and on contraction, v Quain, *Anat* 9th edit, London, 1882, p 118 *et seq*, also O Nasse, *Zur Anat u Physiol d quergestreiften Muskelsubstanz*, Leipzig, 1882, v Schafer, *Pr* 1891)

The sarcolemma is homogeneous and elastic, it is composed of an elastin like substance (Ewald, *Zeit Biol* 26, 1)

The muscle plasma, fluid during life, coagulates after death, producing the stiffening called *rigor mortis*. As blood plasma separates into a solid substance, fibrin, and a liquid residuum, serum, after it is shed, so the muscle plasma separates into a clot composed of myosin and a liquid residue muscle serum. Like the coagulation of the blood this can be hindered by cold, and by squeezing the frozen muscles of the frog Kuhne obtained a liquid muscle plasma, which set into a clot which expressed serum subsequently when it contracted. This occurred most readily at about 30°-40°C. This can also be demonstrated in the muscles of warm blooded animals, but as *rigor* occurs there more rapidly, great expedition in manipulation is required. *Rigor mortis* is also accompanied by the formation of sarcocollactic acid.

Admixture of muscle plasma with solutions of neutral salts prevents the coagulation of the latter. Dilution of such salted muscle plasma brings about coagulation, this occurs most readily at 37°-40°C. Saline extracts of rigid muscle differ from salted muscle plasma in being acid, but resemble it very closely in the way in which myosin can be made to separate from it, myosin in fact undergoes a re-coagulation. This is not a simple precipitation, it is first a jelling through the liquid, the clot subsequently contracts, squeezing out a colourless fluid or salted muscle serum. This does not take place at 0°C, it occurs most readily at the temperature of the body, and is hastened by the addition of a ferment prepared from muscle in the same way as Schmidt's ferment is prepared from blood. The ferment is not identical with

fibrin ferment, as it does not hasten the coagulation of salted blood plasma, nor does the fibrin ferment hasten the coagulation of muscle plasma. The re-coagulation of myosin is also accompanied by the formation of lactic acid.

The proteids of muscle plasma are—

1 Paramyosinogen, which is coagulated by heat at 47°C

2 Myosinogen, which is coagulated at 56°C. It is on the presence of this proteid that the power of fresh muscle juice to hasten the coagulation of blood plasma depends

3 Myoglobulin, which differs chiefly from serum globulin in its coagulation temperature (63°C)

4 Albumin, which is apparently identical with serum albumin α , coagulating at 73°C

5 Myoalbumose, this has the properties of deutoalbumose, and is identical with, or closely connected to, the myosin ferment

The first two proteids in the above list go to form the clot of myosin, paramyosinogen (called muscubin by Hammarsten) is, however, not essential for coagulation, the three last remain in the muscle serum

Paramyosinogen, myosinogen, and myoglobulin are proteids of the globulin class. They are all completely precipitated by saturation with magnesium sulphate, or sodium chloride, or by dialysing out the salts from their solutions. They can be separated by fractional heat coagulation, or by fractional saturation with neutral salts

When muscle turns acid, as it does during *rigor mortis* the pep-in which it contains is enabled to act, and at a suitable temperature (35°-40°C) albumoses and peptones are formed by a process of self digestion. It is possible that the passing off of *rigor mortis*, which is apparently due to the reconversion of myosin into myosinogen, may be the first stage in the self digestion of muscle. The usual theory with regard to the cause of the disappearance of *rigor* is that it is due to putrefaction setting in. Cossar Ewart (*Proc R Soc* 1887) has shown that the disappearance of *rigor* and the appearance of bacteria in the muscle are simultaneous. C Schipilloff's theory is that the lactic acid which is formed from the glycogen in muscle (Otto) produces *rigor* by precipitating the myosin, and the disappearance of *rigor* is due to more acid being formed, which redissolves the precipitate. R Bohm has, however, shown that lactic acid is not derived from glycogen, but from a proteid source, and Latham has been able to deduce a formula which represents the formation of the acid from a combination of cyan hydrins such as he supposes a proteid to be

For the properties of myosin v PROTEIDS

For fuller details respecting muscle plasma and the proteids of muscle consult Kuhne, *Protoplasma*, Leipzig, 1864, E Grubert, *Maly's Jahrsber* 18, p 307, J Klemptner, *ibid* p 310, E Kugler, *ibid* p 311, Demant, *Zeit physiol Chem* 3, 241, 4, 386, Halliburton, *J Physiol* 8, 133. Concerning the formation of acid during coagulation, v Kuhne, *lc*, Nasse, *lc*, Weyl, A. Seidler, *Zeit physiol Chem* 557 (W & S suppose that the acidity is partially due to the formation of acid potassium phosphate, the phos-

phoric anhydride being derived from the lecithin and nuclei of the muscle), Berzelius (*Lehrbuch*, 9, 569), Du Bois Reymond (*Gesammelte Abhandl. zur allgem. Muskel und Nervenphysik*, Leipzig, 1877, 2, p. 3), Heidenhain (*Mechanische Leistung*, p. 143), R. Böhm (*Pf* 23, 44), Hoppe-Seyler (*H* 666), Latham (*Brit Med Journal*, vol. 1, 1886, p. 680), C Schiploff (*Centralbl f d med Wissen*, 1882, 291), Chittenden (*Studies from Lab Physiol Chem Yale*, 3, 116). Concerning the digestion of myosin, see Kühne and Chittenden (*Zeit Biol* 25, 358).

Pigments of muscle—Hæmoglobin is present in small quantities in nearly all muscles, it is contained in the muscle plasma, and it is especially abundant in the slowly contracting red muscles which occur in rodents, and occasionally also in other animals. In the gastropods, *Limæna* and *Paludina*, the muscles contain hæmoglobin, but there is none in the blood (Lankester, v also HÆMOGLOBIN).

Myohæmatin is one of a group of colouring matters called histohæmatins, i.e. pigments occurring in the tissues. These substances are probably respiratory in function, they have not been definitely separated from the tissues, but are probably proteid in nature and contain iron, myohæmatin can be recognised most easily, after soaking the muscle in glycerine, by the spectroscopic, myohæmatin is contained in the muscle plasma. Myohæmatin exhibits four absorption bands—one just below δ , two between δ and ϵ , and one just below ϵ . By soaking the muscle in ether, as a result of osmotic phenomena, the liquid separates out two layers, the lower of which is watery, yellowish-red in colour, and contains myohæmatin which presents a slightly different spectrum from that just described, viz. one band between δ and ϵ and one between ϵ and δ . In both cases the bands are very feeble when the pigment is oxygenated, but become well marked on the addition of reducing agents (MacMunn, *Phil Trans* 1, 1886, p. 267, *J Physiol* 8, 51). Hoppe-Seyler believes myohæmatin is altered hæmoglobin (*Zeit physiol Chem* 13). This, however, has been shown by MacMunn to be untenable (*ibid*).

Constituents of Muscle—Muscle contains on the average 75 p.c. of water, this percentage is higher in young animals and in cold-blooded animals, of the 25 p.c. of solids, 21 p.c. consists of the proteids already described, and the remaining 4 p.c. of extractives and salts (*cf* Hofmann, *Lehrbuch der Zoochemie*, 104). The extractives are divided into (1) *nitrogenous*, viz. creatine the most abundant (0.2 to 0.3 p.c. Voit, *Z B* 4, 77, increased by starvation, Demant, *Zeit physiol Chem* 8, 387), creatinine, xanthine, hypoxanthine, and carmine, (2) *non nitrogenous* viz. fats, glycogen (C. Bernard, *C R* 48, 678, Nasse, *Pf* 2, 97, Brücke, *Sitz W* 63, 214, Abels, *Med Jahrbücher*, 1877, 551, Kulz, *Z B* 22, 161), inosite (Scherer, *Annal d Chem u Pharm* 78, 322, Gauret, a Villiers, *C R* 86, 486), sarcosolactic acid and lactic acid. In addition to the ferments already described (pepsin and myosin-ferment), muscle also contains an amolytic ferment (Nasse, *lc*). Fresh muscle yields on ignition 1 to 1.5 p.c. of mineral matters, of which the most important constituents are potas-

sum and phosphoric acid (*cf* Hofmann, *lc.*, and Bunge, *Zeit physiol Chem* 9, 60).

Contraction of muscle—The processes that occur in resting muscle are twofold—one a change of matter, this chemical tonus, as it may be called, is lessened by curare poisoning, by which the influence of the nervous system over the muscular is shut off and the other set of changes is a change of the potential energy of chemical affinity into actual energy evidenced by the production of heat. On the contraction of a muscle, there is a sudden acceleration of both these changes, viz. an increase in chemical decomposition, and in the conversion of potential into actual energy which is evidenced as heat, electrical inequality, and mechanical motion. It is with the former of these, the chemical changes, that we have here specially to deal. They may be briefly summarised as follows—

1 **Change in reaction**—The muscle ordinarily alkaline becomes acid, as it does during *rigor mortis*, and the acid produced is lactic acid. The acidity can be easily demonstrated by litmus paper (Kühne). It is the accumulation of this and other waste products, including alkaloidal substances (Mosso), in the muscle which produces fatigue—2 There is a relative increase of water (Ranke, *Tetanus*, cap. 2, p. 63). 3 The extractives soluble in water decrease, those soluble in alcohol increase (Helmholtz, *Arch f Anat u Phys* 1845 72, Ranke, *lc* 141, Heidenhain, *Pf* 3, 574)—4 Glycogen diminishes and sugar increases in amount (Ranke, Nasse, *Pf* 2, 97)—5 Creatine diminishes and creatinine increases in amount (Sarokin, *Virchow's Archiv*, 28, Voit, *Z B* 4, 77)—6 Tanned muscle is not able to oxidise pyrogallie acid as resting muscle is (Grützner, *Pf* 7, 255)—7 Nitrates are converted into nitrites (Geschleiden, *ibid* 8, 506)—8 **Gaseous changes**. The amount of oxygen used and of carbonic acid given out increases, the amount of carbonic acid exhaled is never equal in amount to that of the oxygen absorbed, and during tetanus, i.e. continuous contractions, the quotient $\frac{\text{CO}_2 \text{ exhaled}}{\text{O absorbed}}$ increases. The following

numbers from Ludwig and Schmidt illustrate the differences in the gases of the blood leaving muscle during rest and activity

Venous blood.	Oxygen less than arterial blood.	CO ₂ more than arterial blood
Muscle at rest	9 p.c.	6.71 p.c.
„ during activity	12.26 „	10.79 „

(For analyses of the gases of muscle v. Hermann, *Stoffwechsel der Muskeln*, Ludwig, Sezelkow u. A. Schmidt, *Sitz W* 45, *Sitzungsber math phys Classe der k. s. Gesellsch der Wissensch* 20, 12, *Arbeiten aus d physiol Anstalt zu Leipzig*, 1869. Full references of the literature on the effect of muscular exercise on respiration are given by Gamgee, *Physiol Chem* p. 382).

No trustworthy results exist which show that the proteids of muscle undergo any change during activity, and the effect of muscular exercise on the nitrogenous excreta is very small, the increase of urea being quite out of proportion to the amount of work done. (For experiments on dogs v. Voit, *Untersuchungen über den Einfluss*

der Kochsalzes des Kaffees und der Muskelbewegungen auf den Stoffwechsel, München, 1860 For experiments on man, v Fick u Wislizenus, *Vierteljahrsschrift d nat Gesellsch in Zurich*, 10, Parkes, *Pr* 15, 389, 16, 44, E Smith, *Phil Trans* 1862, 747, A Flint, *Journ of Anat and Physiol* 12, 91, F W Pavy, *Lancet*, 1876, North, *Journ of Physiol* 1, 171, *Pr* 39, 443)

Hermann's theory of muscular contraction.—No oxygen is obtainable from muscle *in vacuo* Hermann considers that the formation of carbonic acid is not simply the result of oxidation, but due to the splitting of a complex substance *inogen* into carbonic acid, lactic acid, and a gelatinous proteid myosin, the same occurs, but to a greater extent, in *rigor mortis*, the process of clotting especially going further Each contraction is thus the partial death of the muscle This is supported by the fact that the electrical conditions, like the chemical, are similar in dead and contracted muscle There is, however, no evidence to prove that a clot of myosin is formed at each contraction Bernstein has more recently formulated a theory in which he seeks to show that changes in form, in composition, and in electrical potential are all parts of the same mechanism (*Untersuch a d physiol Inst Halle*, 1888) See also Burdon Sanderson (*Re ports Brit Ass* 1889) W D H

MUSTARD OILS The seeds of black mustard (*Sinapis nigra*) contain potassium myronate, which, in presence of water, is decomposed by the ferment myrosin (also present in the seeds) into KHSO_4 , glucose, and allyl thiocarbimide or oil of mustard Small quantities of crotonitrile and free sulphur are formed in the hydrolysis (Forster, *L V* 1888, 209) Black mustard seed also contains a fixed oil which yields stearic and erucic acids on saponification (Darby, *A* 69, 1) The allyl thiocarbimide amounts to about 5 p c of the weight of the seeds White mustard seed (*Sinapis alba*) yields on pressure 36 p c of a fixed oil containing glyceryl erucate The seeds also contain a glucoside, sinabin $\text{C}_{20}\text{H}_{31}\text{N}_2\text{S}_2\text{O}_{16}$ which is decomposed by myrosin into sinapin sulphate $\text{C}_{16}\text{H}_{19}\text{NO}_2\text{H}_2\text{SO}_4$, glucose, and an oil $\text{C}_{17}\text{H}_{33}\text{NSO}$ (Will, *Z* [2] 7, 89, *A* 199, 150) The term 'mustard oil' has been applied not only to the fixed and volatile oils from mustard seed, but also to all compounds of the form RNCS where R denotes a hydrocarbon radicle In this dictionary these compounds are described as thiocarbimides Thus the essential oil of black mustard is described as allyl thiocarbimide

MYCOMELIC ACID $\text{C}_{17}\text{H}_{33}\text{N}_2\text{O}_4 \cdot \frac{1}{2}\text{aq}$ When alloxan is gently warmed with aqueous NH_3 , it forms a yellow solution which deposits on cooling a transparent jelly of ammonium mycomelate, from the hot aqueous solution of which salt H_2SO_4 ppts mycomelic acid (Liebig & Wöhler, *A* 26, 304) Mycomelic acid is also produced by boiling azulonic acid with water (Emmerling & Jacobsen, *B* 4, 951) and by heating uric acid with water at 180° (Wöhler, *A* 103, 118, Hlawetz, *A* 103, 211) Gelatinous pp, drying up to a loose yellow powder Reddens litmus Almost insol cold water, m sol hot water and alkalis, m sol alcohol and ether — $\text{AgC}_7\text{H}_{11}\text{N}_2\text{O}_2$.

MYCOSE v TREHALOSE and SUGARS.

MYCOTONINE $\text{C}_{27}\text{H}_{51}\text{N}_3\text{O}_5 \cdot 5\text{aq}$. [144°] Oc-

curs, together with lycaconitine, in *Aconitum lycoctonum* (Dragendorff & Salomonovitch, *C O* 1886, 861) V sol chloroform and benzene, nearly insol ether (difference from lycaconitine) Poisonous, 01 g killing a frog Produces paralysis of the extremities of the motor nerves.

MYO-HÆMATIN v MUSCLE

MYOSIN v PROTEIDS and MUSCLE

MYRICIN $\text{C}_{26}\text{H}_{40}\text{O}_2$. [72°] The portion of bees'-wax insoluble in alcohol It is myricyl palmitate (Brodie, *A* 71, 144) Light feathery crystals (from ether), readily saponified by alcoholic potash According to Nafziger (*A* 224, 251) myricin also contains an ether of oleic acid

MYRICYL ALCOHOL $\text{C}_{26}\text{H}_{42}\text{O}$ $\pm e$ $\text{C}_{26}\text{H}_{42}\text{O}$, CH_3OH [85.5°] Obtained by saponifying carnauba wax, in which it occurs both free and combined (Maskelyne, *C J* 22, 87, Von Pieverling, *A* 183, 344, Sturcke, *A* 223, 294) According to Brodie (*A* 71, 147) myricyl alcohol is obtained by saponifying the myricin of bees'-wax, but Schwallb (*A* 235, 106) considers that the myricyl alcohol so obtained has the homologous formula $\text{C}_{25}\text{H}_{40}\text{O}$

Properties — Small white needles (from ether), almost insol cold alcohol, ether, and benzene, but readily soluble in these liquids when hot On heating with soda lime at 200° it forms melissic acid $\text{C}_{25}\text{H}_{40}\text{O}_2$, CO_2H [90°]

DI MYRICYL-AMINE $\text{C}_{52}\text{H}_{84}\text{N}_2$ $\pm e$ $\text{NH}(\text{C}_{26}\text{H}_{41})_2$. [78°] Formed by passing NH_3 for 24 hours over myricyl iodide at 120° (Von Pieverling, *A* 183, 351) Crystalline, nearly insol boiling alcohol and ether, v sol boiling benzene

MYRICYL CHLORIDE $\text{C}_{26}\text{H}_{41}\text{Cl}$ [64.5°] Formed from myricyl alcohol and PCl_5 (Von Pieverling, *A* 183, 348) Waxy mass (from ether), sol alcohol, benzene, and ligroin

MYRICYL CYANIDE $\text{C}_{26}\text{H}_{39}\text{CN}$ [75°] Amorphous (Von Pieverling, *A* 183, 357)

MYRICYL IODIDE $\text{C}_{26}\text{H}_{39}\text{I}$ [70°] Formed by adding phosphorus and iodine in successive small portions to myricyl alcohol heated to 120° (Von Pieverling, *A* 183, 347) Plates (from ligroin), v sol hot alcohol and ether When heated with finely divided potassium it yields $\text{C}_{26}\text{H}_{42}$. [102°] (Hell & Hagele, *B* 22, 502)

MYRICYL MERCAPTAN $\text{C}_{26}\text{H}_{41}\text{SH}$ [94.5°] An amorphous yellow powder, formed by the action of alcoholic KSH on myricyl chloride (Von Pieverling, *A* 183, 349) Sl sol boiling ether, ligroin, and alcohol

MYRISTIC ACID $\text{C}_{14}\text{H}_{27}\text{O}_2 \pm e$ $\text{C}_{13}\text{H}_{25}\text{O}_2$, CO_2H Mol w 228 [54°] (250.5° v at 100 mm) Kraft, *B* 12, 1668, 15, 1724, 16, 1719) S G \pm 8622 H F 107,000 (Von Rechenberg) H C 2,061,712 (Lougumne, *A Ch* [6] 11, 222) Occurs as glyceryl ether (myristin) in nutmeg-butter (from *Myristica moschata*) (Playfair, *P M* [3] 18, 202, *A* 37, 153, Fluckiger, *N Rep Pharm* 24, 213), in Otoba-wax or otobite (from *Myristica otoba*) (Uricoechea, *A* 91, 369), in dika bread (prepared from the fruit of *Mangifera Gabonensis*) (Oudemans, *J pr* 81, 356) amounting to more than one half of the fatty acids contained therein, in the oil of *Cyperus esculentus* (Hell & Twerdomehoff, *B* 22, 1742), in small quantity in cocoa nut oil (Gorgey, *A* 66, 314), in common butter (Heintz, *P* 87, 267, 90, 187, 92, 429, 588, *J pr* 66, 1), in croton oil

(Schlippe, *A* 105, 1), and in Bicuhyba wax (from *Myristica Bicuhyba*). It occurs in combination with ethal in spermaceti (Heintz, *A* 92, 291). It also occurs in the seeds of *Nigella sativa* (Greenish, *Ph* [3] 11, 909, 1013) and in lycopodium spores (from *Lycopodium clavatum*) (Langer, *Ar Ph* [3] 27, 241, 289). Myristic acid is formed by fusing stearic acid with potash (Marasse, *B* 2, 361).

Preparation—By saponifying nutmeg butter or myristin and distilling the acid obtained under reduced pressure.

Properties—Shining laminae (from alcohol), insol water, v sol hot alcohol and ether. A mixture of 30 pts myristic acid and 70 pts lauric acid melts at 35°. Nitric acid (S G 1.5) readily oxidises it, forming various products (Uverdlinger, *B* 19, 1893). The dry distillation of the calcium salt yields myristone. Distillation with MeOH *in vacuo* yields tridecane (Mai, *B* 22, 2133).

Salts—KA' crystalline soap, v sol water and alcohol, insol ether (Playfair)—NaA'—BaA', minnelaminæ, v sl sol water and alcohol—MgA', Baq minute needles (Heintz)—CuA', minute bluish green needles—PbA', amorphous mass—(PbA')₂, Pb(OAc)₂, insoluble powder—AgA' amorphous powder.

Ethyl ether EtA' [11°] (Lutz, *B* 19, 1433) (295°) S G (liquid) 864 (Playfair). Formed from the acid, alcohol, and HCl. Crystals, al sol alcohol and ether, m sol ligroin.

Glyceryl ether C₁₁H₂₁O₂ or **C₁₂H₂₂O₂**, *Myristin Trimyristin* [55°] and [49°] H G 6,601,895 (Lougumine). Occurs in the cases mentioned above, and also in large quantity in the fat of the oil nut (*Myristica surinamensis*) (Reimer a Will, *B* 18, 2011), and to the extent of 1.5 or 2 p c in cochineal (Liebermann, *B* 18, 1975). It is best obtained by extracting powdered nutmeg with ether (Masino, *G* 10, 72). Brilliant leaflets, v sol warm alcohol, ether, benzene, and CHCl₃, nearly insol cold alcohol. It forms two varieties, melting at 55° and at 49°, which are interchangeable by heating 1° above the melting-point for half a minute (R a W, I).

Phenyl ether A'C₆H₅ [36°], (230° at 15 mm).

p-Tolyl ether A'C₆H₄ [39°], (240° at 15 mm) (Krafft a Burger, *B* 17, 1379).

Chloride C₁₁H₂₁COCl [−1°] (168° at 15 mm). Colourless liquid (Krafft a Burger).

Amide C₁₁H₂₁CONH₂ [102°] Formed by heating the glyceryl ether with alcoholic NH₃ at 100° (Masino, *A* 202, 173) or the ethyl ether with aqueous NH₃ at 250°. Formed also by heating the ammonium salt at 230° in a sealed tube (Reimer a Will, *B* 18, 2016), and by the action of NH₃ on the chloride (Krafft a Stauffer, *B* 15, 1730). Plates (from alcohol), v sol benzene, alcohol, and chloroform, sl sol ether, insol water. Bromine and NaOHAq forms C₁₁H₂₁NH CO NH CO C₁₁H₂₁ [103°].

Anilide C₁₁H₂₁CONHPh [84°] Prepared by boiling the acid with aniline for some days (Masino, *G* 10, 75). Silky needles, sol ether, benzene, and chloroform.

Nitrile C₁₁H₂₁CN [19°] (226.5° at 100 mm) S G $\frac{1}{2}$ 6981, $\frac{1}{2}$ 7724. Formed from the amide by distilling with P₂O₅ (Krafft a Stauffer, *B* 15, 1730).

Myristic-benzoic-anhydride

C₁₁H₂₁COOCO C₆H₅ [38°] Formed from BzCl and potassium myristate (Chiozza a Malerba, *A* 91, 102). Silky laminae, m sol ether.

Bromo-myristic acid C₁₁H₂₁BrO₂ [31°] Formed from myristic acid, amorphous P and Br (Hell a Twerdomedoff, *B* 22, 1745). Needles, insol water, sol alcohol and ether.

Tetra bromo-myristic acid C₁₁H₂₁Br₄O₄ Formed from myristic acid and Br (Masino).

Amido-myristic acid C₁₁H₂₁(NH₂)O₂ [253°] Formed from bromo myristic acid and alcoholic NH₃ (H a T). Needles, insol alcohol.

Phenyl-amido-myristic acid

C₁₁H₂₁(NHPh)O₂ [143°] Formed from bromo myristic acid and aniline (H a T). White mass, insol water, sl sol benzene, v sol alcohol. Gives a dark green pp, with cupric acetate in hot alcoholic solution.

Oxy-myristic acid C₁₁H₂₁(OH)O₂ [51.5°] Formed by boiling bromo myristic acid with excess of aqueous NaOH (H a T). Crystalline, insol hot water, v sol alcohol and benzene—BaA', flocculent pp, sl sol cold water, v sl sol hot water—AgA' white pp.

MYRISTIC ALDEHYDE C₁₁H₂₁CHO [53°] (169° at 22 mm). Crystalline solid. Prepared by dry distillation of a mixture of calcium myristate and formate (Krafft, *B* 13, 1415).

Isomeride—TETRADECIC ALDEHYDE

MYRISTICIN C₁₂H₂₁O₂ [30°] (c 145° at 10 mm) S G $\frac{25}{4}$ 1501. Occurs in oil of mace (Semmler, *B* 23, 1803). Yields benzene when distilled with zinc dust. Bromine forms C₁₂H₂₁Br₂O₂ [105°].

MYRISTICOL C₁₁H₂₁O (212°–218°). The chief constituent of the volatile oil of nutmeg (Gladstone, *C J* 25, 11, Wright, *C J* 26, 549, 686). Resinified by heat. P₂S₅ converts it into cymene. PCl₅ yields a chloride, C₁₁H₂₁Cl [100°], slowly split up on boiling into HCl and cymene.

MYRISTIN v *Glyceryl ether of MYRISTIC ACID*.

MYRISTOLIC ACID C₁₁H₂₁O₂ [12°] Formed by passing chlorine into myristic acid heated to 100°, and decomposing the product with alcoholic potash (Masino, *A* 202, 175). Oil. Not solidified by nitrous fumes. Gives Pettenkofer's reaction with sugar and H₂SO₄.

MYRISTONE C₆H₁₁CO [76°] S G $\frac{80}{4}$ = 801, $\frac{100}{4}$ = 792. Silvery plates. Formed by distilling calcium or barium myristate with lime (Overbeck, *P* 86, 591, *A* 84, 290, Krafft, *B* 15, 1713). Does not combine with NaHSO₄.

Oxim C₂₀H₃₃N [51°] Amorphous, sl sol alkalis (Spiegler, *B* 17, 1575, *M* 5, 242).

MYRISTONITRILE v *Nitrile of MYRISTIC ACID*.

MYRONIC ACID C₁₁H₂₁NSO₃ (from *μύρον*, a fragrant ointment). Occurs as potassium salt in the seeds of black mustard (Bassy, *J Ph* 16, 39, Ludwig a Lange, *Z* 1860, 430, 577, Will a Körner, *A* 125, 257), in horse radish (Winckler, *J* 1849, 436), in rape seed (*Brassica rapa*), and in turnip seed (*Brassica napus*) (Ritthausen, *J pr* [2] 24, 273). This salt may be obtained by boiling mustard seeds (1 pt) with alcohol (6 pts), macerating the residue with cold water, and evaporating the aqueous extract after addition of a little BaCO₃. The free acid may be

obtained by adding tartaric acid to a solution of the potassium salt, evaporating, and extracting with alcohol Syrup, readily decomposed by heat Its aqueous solution gives off H_2S on boiling An aqueous solution of myrosin splits it up into glucose, allyl thiocarbimide, and H_2SO_4 Boiling baryta water forms a pp of $BaSO_4$, with evolution of allyl thiocarbimide Caustic potash solution acts vigorously, yielding allyl cyanide, NH_3 , glucose, and allyl thiocarbimide Conc HCl sets free H_2SO_4 , boiling dilute H_2SO_4 yields H_2S , glucose, H_2SO_4 , and NH_3 Zinc and $HClAq$ give off H_2S

Salts— KA' Groups of silky needles (from alcohol) or glassy prisms (from water), v e sol water, nearly insol alcohol, insol ether Tastes bitter Its solution is hydrolysed by myrosin

$KC_{10}H_7NS_2O_{10} = C_6H_5O_6 + C_4H_5NCS + KHSO_4$ The hydrolysis is not brought about by emulsion, yeast, or saliva Water at 115° yields allyl cyanide, H_2S , and sulphur Silver nitrate solution gives a pp $C_6H_5NSAg_2SO_4$. — BaA'_2 (at 100°) plates, v sol water

MYEOSIN A proteid ferment contained in the seeds of black and white mustard and of many other cruciferous plants It may be obtained by exhausting white mustard with cold water, evaporating below 40° to a syrup, and ppg by alcohol (Bussy, *J Ph* 26, 44, Winckler, *Jahrb pr Pharm* 3, 93) Its aqueous solution is coagulated by heat and by alcohol, when it loses its hydrolytic power, but it recovers this after a day's immersion in water It does not hydrolyse amygdalin

MYROXOCARPIN $C_{18}H_{19}O_6$ [115°] Deposited in crystals from an alcoholic solution of

white balsam of Peru (Stenhouse, *A* 77, 306) Trimetric crystals, $a b c = 1.936 : 755$ Insol water, v sol hot alcohol and ether Does not dissolve in acids or alkalis

MYRRH A gum resin which exudes from *Balsamodendron myrrha*, a shrub growing in Arabia and Abyssinia The resin yields proto catechuic acid and pyrocatechin on potash-fusion (Hlasiwetz & Barth, *J* 1866, 630) Besides resin and gum (Buckner, *N Rep Pharm* 16, 76), myrrh contains a small quantity of an essential oil, boiling about 266° , $S G$ 15° 1.0189, μ_A 1.5196 at 75° , μ_D 1.5278, μ_H 1.5472 (Gladstone, *C J* 17, 11) The oil quickly resinifies when exposed to atmospheric oxidation It contains $C_{10}H_{16}O$ (263°) (Flückiger, *B* 9, 471) According to Kohler (*Ar Ph* 228, 291) myrrh contains a gum $C_6H_9O_3$, a resin $C_{20}H_{31}O_2(OH)$, two dibasic acids $C_{12}H_{19}O_8$ and $C_{18}H_{27}O_8$, and 7 p c of an essential oil $C_{10}H_{16}O$

MYRTLE OIL A volatile oil obtained from the berries and leaves of the myrtle (*Myrtus communis*) (Riegel, *Pharm Centr* 1850, 319) It contains a terpene $C_{10}H_{16}$ (160° – 170°), $S G$ 15° 891, μ_A 1.462 at 18° , μ_D 1.468, μ_H 1.488 (Gladstone, *J* 1863, 548) Jahns (*Ar Ph* [3] 27, 174) found in Spanish oil of myrtle pinene $C_{10}H_{16}$ (169°) [α]_D = 36.8 , and cineol (170°)

MYTILOTOXINE $C_6H_7NO_2$ Occurs in mussels (*Mytilus edulis*) and in putrid flesh (Brieger, *Die Ptomaine*, Gautier, *Bl* [2] 48, 13) Its hydrochloride crystallises in tetrahedra, and is very poisonous, but gradually decomposes, losing its poisonous properties — $B'HAuCl$, [182°] Minute cubes

N

NANDININE $C_{15}H_{17}NO$ Occurs in the root bark of *Nandina domestica* of Japan (Eijkman, *R T C* 3, 197) White amorphous powder, insol water, v sol alcohol, ether, benzene, and chloroform Poisonous Gives the alkaloidal reactions H_2SO_4 forms a reddish-violet colour, changed by a drop of HNO_3 to an intense blue Nitric acid gives a green colour changing to brown — $B'H_2PtCl_6$ turned blue by H_2SO_4

NAPHTHA v **PETROLEUM**

(β)-**NAPHTH-ACRIDINE** $C_{21}H_{15}N_3$ v e

$C_{10}H_7 \begin{array}{c} \text{CH} \\ \diagup \quad \diagdown \\ \text{N} \end{array} C_{10}H_7$ [216°] Formed by the action

of methylal, and HCl upon (β) naphthyl amine (Beed, *J pr* [2] 34, 160, 35, 317) Long, straw-yellow, needles, v sol alcohol, v al sol ether The alcoholic solution fluoresces dark blue

Nitrate $B'HNO_3$ small needles

Picrate $B'O_2C_6H_4(NO_2)_3$ amorphous

Derivative v **PHENYL-NAPHTHAACRIDINE**

NAPHTHALDEHYDE v **NAPHTHOIC ALDEHYDE**

NAPHTHALENE $C_{10}H_8$ *Naphthalin* Mol w 128 [80.2°] (Reissert, *B* 23, 2243), [80°] (Landolt, *Z P O* 4, 849), [79.5°] (Vohl), [79°] (Lössen & Zander, *A* 225, 111), [80.06°] (Mills, *P M* [5] 14, 27) (218°) at 760 mm (Vohl, *J*

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pr 102, 29, Crafts, *Bl* [2] 39, 282), (217°) at 740 mm $S G$ 15° 1.152 (V), μ_A 1.145 (Schroder, *B* 12, 1613) $S G$ (liquid) 18° 978 (Kopp, *A* 95, 329), 7° 982 (L & Z) $H C v$ 1,232,400 $H C p$ 1,233,600 (Stohmann), 1,245,000 (Berthelot & Vieille, *Bl* [2] 47, 863), 1,242,000 (Berthelot, *A Ch* [6] 13, 302, 326) $H F$ — 17,600 (Stohmann, Kleber, & Langbein, *J pr* [2] 40, 90), — 29,000 (Berthelot & Vieille, *A Ch* [6] 10, 442), — 42,000 (von Rechenberg) $S V$ 149.2 (L & Z), 148 (Lössen, *A* 254, 53), 145.46 (Ramsay), $S V S$ 130.61 (Schiff) R_p 74.12 in a 6.66 p c alcoholic solution (Kanonnikoff), 71.78 (Nasim & Bernheimer, *G* 14, 153, 15, 93) S (toluene) 32 at 16.5° , S (alcohol) 5.29 at 15° (Bech, *B* 12, 1978) Its absorption in the ultra violet spectrum has been studied by Hartley (*C J* 39, 161)

Occurrence—In petroleum from Rangoon (Warren & Storer, *Mem Amer Acad* 9, 208) In coal tar, from which it may be obtained by shaking the fraction 180° – 220° with aqueous $NaOH$ and then with dilute H_2SO_4 , and distilling the residue alone or with steam (Garden, *Thomson's Annals*, 16, 74, Faraday, *Tr* 1826, Kidd, *B. J.* 2, 186, Reichenbach, *S* 61, 175, 68, 233).

Formation.—1 A product of the passage through a red hot tube of the vapour of the following

G G

lowing substances — petroleum, alcohol (Reichenbach, *B* 12, 307), ether, acetic acid, essential oils, toluene (Ferko, *B* 20, 660), xylene, ψ -cumene, a mixture of ethylene with benzene, with styrene, with anthracene or with chrysene (Berthelot, *Bl* [2] 6, 272, 279), ethylene alone, acetylene, a mixture of benzene and acetylene (Berthelot, *Bl* [2] 7, 218, 278, 306), oil of turpentine (Schulz, *B* 9, 648), wood-tar (Letney & Atherberg, *B* 11, 1210, 1222) — 2 By passing over red hot quicklime the vapour of the bromide of phenyl butylene derived from benzyl bromide, allyl iodide and sodium (Aronheim, *B* 6, 67, *A* 171, 233) — 3 By passing the vapour of isobutyl benzene over heated lead oxide (Wreden & Znatovitch, *B* 9, 1606) — 4 By distilling colophony and gum benzoin with zinc dust (Ciamician, *B* 11, 269) — 5 By heating dimethyl aniline (1 pt) with bromine (1 pt) at 115° (Brunner & Brandenburg, *B* 11, 697) — 6 By oxidising pyrene acid and distilling the resulting naphthalene tetra-carboxylic acid with slaked lime (Bamberger & Philip, *B* 19, 1999) — 7 By hydrolysis of its sulphonic acids this takes place when superheated steam is passed through a solution of naphthalene (β) sulphonic acid in dilute H_2SO_4 at 135° (Armstrong & Miller, *C* 46, 148)

Synthesis — By dry distillation of the silver salt of tetra hydro naphthalene di carboxylic acid $C_{10}H_6$ $\begin{matrix} \text{CH}_2\text{CH}(\text{CO}_2\text{H}) \\ \text{CH}_2\text{CH}(\text{CO}_2\text{H}) \end{matrix}$ which acid is formed

by the action of *o* xylene bromide C_8H_6 $\begin{matrix} \text{CH}_3 \\ \text{CH}_3 \end{matrix}$ Br on di-sodio ethane tetra carboxylic ether $C_2Na_2(\text{CO}_2\text{Et})_4$, and boiling the product with alcoholic KOH Naphthalene is also formed by passing the tetra hydro naphthalene di carboxylic acid through a red hot tube (Baeyer & Perkin, *B* 17, 448) (*v* NAPHTHOL and NAPHTHALENE DERIVATIVES, Constitution of)

Purification — Commercial naphthalene may be purified by sublimation. It may also be purified by repeatedly heating with a little H_2SO_4 (best with MnO_2) at 180° and distilling with steam (Stenhouse & Groves, *B* 9, 683)

Properties — Monoclinic tables, insol cold, almost insol hot, water, *v* sol alcohol, ether, fatty and essential oils, and HOAc. Volatile with steam, 1 pt distilling over with about 570 pts of water (Naumann, *B* 4, 646, 10, 2014, 2100, 11, 33). Burns with smoky flame. Boiling naphthalene dissolves S, P, and the sulphides of As, Sb, and Sn, it also dissolves indigo, iodine, $HgCl_2$, HgI_2 , and As_2O_3 .

Reactions — 1 *Chlorine* forms derivatives by substitution and by addition (Laurent, *A* 49, 218, 52, 275, 69, 214) — 2 *Bromine* forms derivatives by substitution — 3 *Nitric acid* forms nitro and di nitro naphthalene — 4 The vapour of *aqua regia* in the cold forms $C_{10}H_6Cl_4$ and $C_{10}H_6Cl_2$ (Bunge, *B* 4, 289) — 5 *Phosgene* has no action (Berthelot, *Bl* [2] 13, 301) — 6 The vapour passed through a red hot tube yields carbon, methane (Kletzensky, *J* 1865, 561), and dinaphthyl (Ferko, *B* 20, 662). When passed through a red-hot tube together with hydrogen it is mainly unaltered, but yields some acetylene and benzene (Berthelot, *Bl* [2] 6, 281). When passed together with acetylene through a red-hot tube it yields much anthracene. At a white

heat it reacts with benzene forming anthracene (Berthelot). When passed through a red-hot tube containing charcoal it yields some dinaphthyl. When passed with ethylene through a red-hot tube it yields acenaphthene, phenanthrene, and dinaphthyl (Ferko, *B* 20, 662) — 7 Saturated HIAq at 280° yields the dihydride $C_{10}H_{12}$, and finally ethyl and di ethyl benzene and decane (Berthelot, *J* 1867, 709). When heated with conc HIAq and red phosphorus the products are naphthalene hexahydride, and only $C_{10}H_{18}$ (173°–180°) and $C_{10}H_{22}$ (153°–158°) (Wreden, *A* 187, 164) — 8 Boiling aqueous $KMnO_4$ oxidises it to phthalic acid. Aqueous $K_2Cr_2O_7$ and H_2SO_4 yield phthalic acid and dinaphthyl (Lossen, *A* 144, 71). Dilute HNO_3 at 130° also yields phthalic acid (Beilstein & Kurbatow, *A* 202, 215). A mixture of CrO_3 and HOAc oxidises it to naphthoquinone. Oxidation with MnO_2 and H_2SO_4 yields dinaphthyl and a resinous acid $C_{10}H_8O_4$, which forms the amorphous salts Pb_2A''' , $PbHA'''$, and Ag_2HA''' (Lossen) — 9 $KClO_4$ and H_2SO_4 form phthalic acid, di chloro-naphthalenes, and syrupy chloroxy naphthalic acid $C_{10}H_6ClO_4$ (Hermann, *A* 151, 79) — 10 CrO_3Cl_2 yields di chloro naphthoquinone — 11 Aqueous hypochlorous acid forms $C_{10}H_6(\text{HOCl})_2$, crystallising in prisms, *sl* sol water, converted by alcoholic potash into $C_{10}H_6(\text{OH})_2$, which crystallises in prisms, almost insol water, *v* sol alcohol, and forms an insoluble lead compound $Pb_2C_{10}H_6O_4$ (dried at 100°) and a sulphonic acid which yields a crystalline calcium salt $CaC_{10}H_6S_2O_{10}$ (Neuhoff, *A* 136, 342) — 12 When heated with excess of $AlCl_3$ it forms benzene and hydrides of Naphthalene. At 160° iso dinaphthyl is formed (Friedel & Crafts, *Bl* [2] 39, 195, *C* 100, 692) — 13 *Methyl chloride* in presence of $AlCl_3$ forms $C_{10}H_{12}$ [181°] crystallising in plates, *v* sol hot ether, *sl* sol cold alcohol (Bischoff, *B* 23, 1905, *C* Liebermann, *A* 163, 122, Furth, *B* 16, 2171) — 14 *Iodine* at 250° appears to form a compound $C_{10}H_8I$ (Bleunard & Vrau, *C* 94, 534) — 15 On passing a mixture of cyanogen and naphthalene vapour through a red hot tube there is formed the nitrile of (α) naphthoic acid — 16 Heating with *chloride of sulphur* yields di chloro naphthalene (Laurent, *A* 76, 298) — 17 *Nitric peroxide* forms nitro and di nitro naphthalene and, at 100°, the compounds $C_{10}H_6O_4$ [225°] and $C_{10}H_6O_2$ [131°] (Leeds, *Am* 2, 283) — 18 When heated with *potassium* it forms a black powder $C_{10}H_6K_2$, which is decomposed by water, yielding KOH and $C_{10}H_6$ (Berthelot, *Bl* [2] 7, 110) — 19 Naphthalene (2 pts) fused with *antimonious chloride* (8 pts) yields on cooling deliquescent crystals of $(C_{10}H_6)_2Sb_2Cl_4$ (W Smith, *C* 41, 411) — 20 Naphthalene taken internally appears in the urine as (β) naphthol and (β) naphthoquinone (Edlefsen, *C* 1888, 1007)

Combinations with nitro compounds — $C_{10}H_6C_6H_5(NO_2)_2$ [18] [53°] Prisms (Hepp, *A* 215, 379) — $C_{10}H_6C_6H_5(NO_2)_2$ [14] [119°] White needles, *v* *sl* sol alcohol, separated into its components by distillation with steam — $C_{10}H_6C_6H_4Cl(NO_2)_2$ [124] [78°] Long white needles (from alcohol), decomposed by heating with potash or aniline, naphthalene being set free (Willgerodt, *B* 11, 603) — $C_{10}H_6C_6H_5(NO_2)_2$ [162°] White needles, deposited from a mixture

of the alcoholic solutions of naphthalene and tri nitro benzene (Hepp, *A* 215, 377) — $C_{10}H_7Cl(NO_2)_3$ [96°] Flat canary-yellow needles (Liebemann a Palm, *B* 8, 378) — $C_{10}H_7C_2H_4Me(NO_2)_2$ [61°] Formed by mixing naphthalene and di-nitro-toluene dissolved in benzene (Hepp) — $C_{10}H_7C_2H_4Me(NO_2)_3$ [98°] Formed from (a) tri nitro toluene and naphthalene in alcoholic solution (Hepp) Needles — $C_{10}H_7C_2H_4Me(NO_2)_2$ [100°] Formed from (8)-tri nitro toluene and naphthalene Yellowish needles (from alcohol) — $C_{10}H_7C_2H_4Me(NO_2)_3$ [89°] Formed from (7) tri-nitro-toluene (H) — $C_{10}H_7C_2H_4(NO_2)_3NH_2$ [169°] Orange prisms $C_{10}H_7C_2H_4(NO_2)_3OH$ Yellow needles (Gruner, *Z* 1868, 218) — $C_{10}H_7C_2H_4(NO_2)_3OH$ [73°] Formed from (8)-tri nitro phenol and naphthalene (Henriques, *A* 215, 332) *V* e s o l alcohol — $C_{10}H_7C_2H_4(NO_2)_3OH$ [100°] Formed from naphthalene and (7) tri nitro phenol (Henriques) Golden needles (from alcohol), s l s o l alcohol — $C_{10}H_7C_2H_4Me(NO_2)_3OH$ [106°] From naphthalene and tri nitro *m*-cresol (Nöling a Collin, *B* 17, 271) Yellowish needles (from acetone) — $C_{10}H_7C_2H_4Me(NO_2)_3OH$ [127°] From naphthalene and tri nitro *m*-cresol (Nöling a Sals, *B* 15, 1862) — $C_{10}H_7C_2H_4(NO_2)_3S$ [50°] Yellow needles (from benzene) Formed from naphthalene and di nitro thiophene (Rosenberg, *B* 17, 1778)

Picric acid compound

$C_{10}H_7C_2H(NO_2)_3OH$ [149°] Golden yellow monoclinic needles, s o l alcohol, ether, and benzene Slowly separated into its components by boiling water

Naphthalene dichloride $C_{10}H_7Cl_2$ Formed by passing chlorine over naphthalene $KClO_3$ and HCl may also be used (E Fischer, *B* 11, 735, 1411) Oil, miscible with ether, m s o l alcohol Begins slowly to decompose at 40°–50° into HCl and chloro naphthalene Alcoholic potash also converts it into chloro naphthalene Sodium or sodium amalgam at 150° converts it into naphthalene

Naphthalene tetrachloride $C_{10}H_4Cl_4$ Mol w 270 [182°] R_D 105 35 in a 2 39 p c chloroform solution (kanonnikoff) Formed by passing a rapid current of chlorine over naphthalene until the product, after having become liquid, thickens again, when it is washed with ether and crystallised from benzene Formed also by treating naphthalene with a saturated solution of chlorine in $CHCl_3$ (Grimaux, *B* 5, 222, Schwarzer, *B* 10, 379), by chlorinating naphthalene in direct sunlight (Leeds a Everhart, *A* *C* *J* 2, 203), and by the action of $KClO_3$ and HCl on naphthalene (Fischer, *B* 11, 735) Large monoclinic prisms, insol water, s l s o l alcohol, m s o l ether, v s o l benzene and petroleum Sublimes at 225°–230° Decomposed on distillation into HCl and (a) and (8) di chloro naphthalenes (Krafft a Becker, *B* 9, 1088) Alcoholic potash forms (a) and some (i) di-chloro naphthalene reconverted into naphthalene on digesting with iron (Zinn, *B* 4, 288) Boiling dilute $AgNO_3$ slowly converts it into $C_{10}H_7(ClO)$ [196°] When boiled with water it yields $C_{10}H_7Cl(OH)$, which crystallises in prisms (from ether) [156°], S 3 3 at 100°, and is decomposed on distillation with HCl into water, HCl , and chloro-naphthol Zn and H_2SO_4 re-

duce it to (a) naphthol It gives rise to $C_{10}H_7Cl_2(OAc)$, [131°] and $C_{10}H_7Cl_2(OBz)$, [150°]

The existence of an isomeric naphthalene tetra-chloride [118°] has been denied by Atterberg (*B* 11, 1223, cf Fischer, *B* 11, 735)

Naphthalene tri chloro bromide $C_{10}H_4Cl_3Br$ Formed from the tetrachloride and bromine; after 48 hours the product is washed with warm alcohol and crystallised from ether Prisms

Naphthalene dihydride $C_{10}H_8$ [15 5°] (212°) VD 4 7 (calc 4 56) Occurs in heavy coal tar oil (Berthelot, *B* [2] 9, 288) Formed by heating naphthalene with conc $HIAq$ for a short time at 280° (Berthelot), and by reducing naphthalene dissolved in isoamyl alcohol with sodium (Bamberger a Lodter, *B* 20, 3073) Naphthalene dihydride is formed by distilling the bromide of naphthalene tetra hydride, or by heating it with alcoholic KOH (Graebe a Guye, *B* 16, 3032) It is likewise obtained by the action of sodium on an alcoholic solution of the nitrile of naphthoic acid (Bamberger a Lodter, *B* 20, 1704) and, as a by product, when (8) naphthylamine dissolved in isoamyl alcohol is reduced with sodium (Bamberger a Muller, *B* 21, 859)

Properties —Tables Does not combine with picric acid By treatment with bromine dissolved in chloroform it is converted into the dibromide $C_{10}H_6Br_2$, which crystallises in thick colourless prisms, v s o l alcohol and ether, and gives off HBr on heating (*B* a *L*)

Naphthalene dihydride $C_{10}H_8$ (200°) Formed by distilling the dihydride of naphthoic acid with soda lime (von Pechmann, *B* 16, 517) Liquid

Naphthalene tetrahydride $C_{10}H_{12}$ *see* $C_6H_5 \begin{matrix} < CH_2 & CH_2 \\ & CH & CH \end{matrix}$ (205°) at 716 mm SG 1 2

978 Formed by heating naphthalene with $HIAq$ at 280° (Berthelot), with PH_4I at 180° (Baeyer, *A* 155, 276), or by adding sodium to its solution in isoamyl alcohol (Bamberger a Kitschelt, *B* 23, 1561) Formed also by suspending the tetrahydride of (a) naphthyl hydrazine (1 pt) in boiling water and allowing a solution of cupric sulphate (2 pts) to drop in slowly, nitrogen is evolved, and the naphthalene tetrahydride may be separated from ppd cuprous oxide by steam distillation (Bamberger a Bordt, *B* 22, 631) Prepared by heating naphthalene (10 pts) with HI (9 pts) and amorphous phosphorus (3 pts) for 8 hours at 220°–225°, the yield is good (8 pts) (Graebe a Guye, *B* 16, 3028, cf Graebe, *B* 5, 678)

Properties —Oil, slowly turning brown in air Smells like naphthalene Its solution in chloroform absorbs bromine, giving off HBr It immediately decolorises an acid solution of $KMnO_4$, and is oxidised to $C_{10}H_7(CO_2H)CH_2CH_2CO_2H$ (Bamberger) The compound obtained by Baeyer and by Graebe yielded phthalic acid on oxidation by $KMnO_4$, and may perhaps not be identical with Bamberger's tetrahydride When passed through a red hot tube it is resolved into naphthalene and hydrogen

Naphthalene hexahydride $C_{10}H_{14}$ *Hexahydronaphthalene* (200°) (*G* a *G*), (205° at 764 mm) (*A*) SG 9 419 CE (0°–26 3°) 000817 (Lossen a Zander, *A* 225, 112) R_D 71 15 (Nasini a Bernheimer, *G* 15, 93) S V.

1712 (Lossen, *A* 254, 53) Formed by heating naphthalene with conc HIAq and phosphorus (Wieden a Znatovitch, *B* 9, 278, 1606, *A* 187, 164) Prepared by heating naphthalene (67 pts) with HI (100 pts) and amorphous phosphorus (30 pts) for 10 hours at 240°–250° (Graebe a Guye, *B* 16, 3031), or by heating naphthalene (10 g), amorphous phosphorus (3 g), and HIAq (9 g, boiling at 127°) at 235° for 7½ hours (Agrestini, *G* 12, 495) Liquid, which absorbs oxygen from the air Does not combine with picric acid Reacts with bromine, giving off HBr, and forming a product which is converted by alcoholic potash into $C_{10}H_7Br$ (270°)

Naphthalene octohydrate $C_{10}H_8 \cdot 8H_2O$ (185°–190°) SG $\frac{1}{2}$ 910, $\frac{1}{2}$ 892 Formed by heating naphthalene (5 g) with HIAq (9 g of SG 17) and red phosphorus for 15 hours at 260° (Guye, *Bn* 2, 138) Liquid, smelling like oil of turpentine Absorbs oxygen from the air

References—TRI AMIDO-, AMYL-, BROMO-, BROMO IODO-, BROMO NITRO-, CHLORO-, CHLORO NITRO-, IODO, IODO NITRO, NITRO, DI OXY, METHYL-, ETHYL-, PROPYL-, PHENYL-, and BENZYL NAPHTHALENES Also NAPHTHOL, NAPHTHYLAMINE and NAPHTHYLENE DIAMINE

NAPHTHALENE, CONSTITUTION OF, v DICTIONARY OF APPLIED CHEMISTRY

NAPHTHALENE ALDEHYDE v NAPHTHOIC ALDEHYDE

NAPHTHALENE-DIAMINE v NAPHTHYLENE-DIAMINE

NAPHTHALENE-ARSONIC ACID v Organic compounds of ARSENIC

NAPHTHALENE-AZO compounds v Azo compounds and Dis azo compounds

NAPHTHALENE CARBOXYLIC ACID v NAPHTHOIC ACID

Naphthalene Peri-dicarboxylic acid $C_{10}H_6O_4$, *ie* $C_{10}H_4(CO_2H)_2$ [11] *Naphthalic acid* Mol w 216 [266°]

Formation—1 By oxidising acenaphthene with $K_2Cr_2O_7$ and H_2SO_4 or HOAc (Behr a Dorp, *B* 6, 862, *A* 172, 266, Anselm, *B* 22, 869) —2 By oxidising pyrene ketone with $KMnO_4$ (Bamberger a Philip, *B* 19, 3040) —3 By saponification of its semi nitrile $C_{10}H_5Cy CO_2H$ which is formed by the action of cuprous cyanide upon diazotised *peri* amido naphthoic acid (Bamberger a Philip, *B* 20, 248) —4 By oxidising di-*exo*-oxy-acenaphthene with alkaline $KMnO_4$ (Ewan a Cohen, *C J* 55, 580)

Properties—Long silky hair like needles (from alcohol), almost insol water, sl sol ether Split up into water and its anhydride by heating alone at 145°, by boiling with HOAc, or even by exposure over H_2SO_4 (Blumenthal, *B* 7, 1092) Distillation with lime yields naphthalene Yields a fluorescein on heating with resorcin

Salts— $(NH_4)A \cdot EtOH$ Plates (from slcohol)— Na_2A — $K_2A \cdot EtOH$ Plates— BaA aq sparingly soluble white silvery plates— CaA aq— Al_2A aq

Di-methyl ether Me_2A [103°]

Anhydride $C_{10}H_4 \cdot \begin{smallmatrix} CO \\ \diagup \diagdown \\ O \end{smallmatrix}$ [266°]

Needles (from alcohol), sl sol alcohol and benzene Not attacked by boiling HNO_3

Imide $C_{10}H_4 \cdot \begin{smallmatrix} CO \\ \diagup \diagdown \\ CO \end{smallmatrix} NH$ [above 280°]

Formed by boiling the acid or its anhydride with aqueous ammonia White needles Sol warm KOHAq When $AgNO_3$ is added to its solution in alcoholic NH_3 , there is formed a crystalline pp $Ag_2C_{10}H_4N_3O_4$

Naphthalene dicarboxylic acid $C_{10}H_6(CO_2H)_2$ [255°] Formed by reducing di-oxy naphthalene dicarboxylic acid [162°] with HIAq and amorphous phosphorus (Claus a Meixner, *J pr* [2] 37, 8) Flocculent pp, insol water and ether, sol alcohol Yields naphthalene on distillation with lime— PbA greyish white pp

Naphthalene 'a' dicarboxylic acid $C_{10}H_6(CO_2H)_2$ [22] Obtained by digesting its nitrile with $HClAq$ at 200° (Ebert a Merz, *B* 9, 606) Long needles (from alcohol) Melts far above 300° V sl sol boiling benzene, toluene, and HOAc, m sol boiling alcohol Yields naphthalene on distillation with lime— CaA 44q minute needles, sl sol water— Ag_2A

Nitrile $C_{10}H_4(CN)_2$ [268°] Formed by distilling naphthalene (a) disulphonic acid with potassium cyanide White needles (by sublimation), m sol boiling alcohol

Naphthalene 'β' dicarboxylic acid $C_{10}H_6(CO_2H)_2$ [23] Obtained in the same way as the 'a' isomeride, from naphthalene 'β' disulphonic acid (E a M) Short needles (from alcohol) Melts far above 300° Almost insol boiling benzene, toluene, and HOAc, m sol boiling alcohol— K_2A 44q tufts of needles, v e sol water— CaA 3½q minute needles, almost insol water— Ag_2A

Nitrile $C_{10}H_4(CN)_2$ [297°] Long needles (from HOAc) Almost insol boiling ether, alcohol, and benzene

Naphthalene (γ) dicarboxylic acid $C_{10}H_6(CO_2H)_2$ Obtained from its nitrile, which is formed by distilling with KCy potassium bromo naphthalene sulphonate (obtained by sulphonating (a) bromo naphthalene) (Darmstadter a Wichelhaus, *A* 152, 809, *Z* [2] 5, 571) Small needles, v sol alcohol, insol boiling water— BaA 24q crystalline grains

Nitrile $C_{10}H_4(CN)_2$ [204°] Thin needles, sl sol alcohol and ether

Naphthalene (δ) dicarboxylic acid **Nitrile** $C_{10}H_6(CN)_2$ [236°] Obtained by distilling potassium bromo naphthalene (a) sulphonate with KCy (D a W) Needles, v sol alcohol

Naphthalene (ε) dicarboxylic acid **Nitrile** $C_{10}H_6(CN)_2$ [170°] Obtained by distilling potassium bromo-naphthalene (β) sulphonate with KCy (D a W) Small needles, v sol alcohol

Naphthalene tri-carboxylic acid $C_{10}H_5(CO_2H)_3$ Obtained by quickly heating the tetra carboxylic acid to 200°–250° (Bamberger a Philip, *B* 19, 3037)

Naphthalene tetra carboxylic acid $C_{10}H_4O_8$, *ie* $C_{10}H_4(CO_2H)_4$ [11' 44'] Formed by oxidising pyrene acid with dilute $KMnO_4$ (Bamberger a Philip, *B* 19, 1998, 20, 368, *A* 240, 182) Colourless glistening plates or needles, m sol hot HOAc and water, v sl sol alcohol and benzene Not attacked by HNO_3 , even at 160° Yields naphthalene on distillation with lime— Ba_2A (dried at 130°)— Ag_2A

Anhydride $C_{10}H_4 \cdot \begin{smallmatrix} CO \\ \diagup \diagdown \\ CO \end{smallmatrix} O$ Formed by heating the acid at 150°–170° (Bamberger a Philip, *B* 19, 3037) Needles (from HOAc)

Sublimes above 300° Gives a fluorescein when heated with resorcin

Imide $C_{10}H_6(\text{CO})_2\text{NH}$ Formed by treating the anhydride with NH_4Aq Needles and prisms Sublimes in needles above 270° V sol alcohol, ether, and benzene Aqueous NaOH colours it yellow (forming the Na salt), without dissolving it, on adding acids it turns white again

Naphthalene dihydride dicarboxylic acid $C_{10}H_8(\text{CO}_2\text{H})_2$ *Dihydronaphthalic acid* Formed by reducing naphthalic acid with sodium amalgam (Anselm, B 22, 859) Colourless crystalline solid, v sol warm alcohol, sl sol ether and water Reduces alkaline KMnO_4 It turns yellow at 150°–160°, froths at 199°, and changes to the anhydride, but is not melted at 275°

Naphthalene tetrahydride dicarboxylic acid $C_{10}H_8(\text{CH}_2\text{CH}(\text{CO}_2\text{H}))_2$ [199°] When an alcoholic solution of ethane tetra carboxylic ether $C_8H_4(\text{CO}_2\text{Et})_4$ (1 mol) and NaOEt (2 mols) is heated at 130° with di ω bromo α xylene $C_8H_4(\text{CH}_2\text{Br})_2$ (1 mol) there is formed naphthalene tetrahydride tetracarboxylic ether $C_{10}H_8(\text{CH}_2\text{C}(\text{CO}_2\text{Et}))_2$, and when this is saponified with alcoholic potash it loses CO_2 (2 mols) yielding the dicarboxylic acid The yield is 70 p c of the theoretical (Breyer & Perkin, B 17, 448, C J 53, 11) Minute tables, sl sol cold, m sol hot, water, v sol chloroform, alcoh, ether, and acetone Splits up at its melting point into H_2O and its anhydride On passing through a red-hot tube it yields naphthalene— $\text{Ag}_2\text{A}''$ white crystalline pp which yields the anhydride and naphthalene on dry distillation

Anhydride $C_{10}H_6(\text{CH}(\text{CH}_2\text{CO}))_2\text{O}$ [184°]

Four sided prisms (from ether) or needles (by sublimation), sl sol ether, m sol alcohol and chloroform, insol cold water Slowly reconverted into the acid by hot water

Naphthalene tetra hydride tetra carboxylic acid $C_{10}H_8(\text{CH}_2\text{C}(\text{CO}_2\text{H}))_2$ *Tetrahydro naphthalene tetracarboxylic acid* The ether is formed by heating an alcoholic solution of sodium chloro malonic ether $\text{CNaCl}(\text{CO}_2\text{Et})_2$ (2 mols) with α xylene bromide (1 mol), reducing the resulting $C_8H_4(\text{CH}_2\text{CCl}(\text{CO}_2\text{Et}))_2$ by means of zinc dust and HOAc to α xylene di malonic ether $C_8H_4(\text{CH}_2\text{CH}(\text{CO}_2\text{Et}))_2$, and treating the sodium derivative $C_8H_4(\text{CH}_2\text{CNa}(\text{CO}_2\text{Et}))_2$ with an ethereal solution of iodine (Baeyer & Perkin, jun, B 17, 448) The ether may also be obtained from $C_8H_4(\text{CH}_2\text{Br})_2$, sodium ethylate and $C_2H_5(\text{CO}_2\text{Et})_2$ (*v supra*) The free acid, which may be obtained by careful saponification of the ether with alcoholic potash, is a syrup which splits up at 185° into CO_2 and the dicarboxylic acid described above

Reference—BROMO NAPHTHALENE DICARBOXYLIC ACID

DI-NAPHTHALENE OXIDE v DINAPHTHYLENE OXIDE

NAPHTHALENE DI-OXIM v *Di-oxim* of NAPHTHOQUINONE.

NAPHTHALENE - DIPHENYL AZAMMONIUM HYDRATE v AZAMMONIUM COMPOUNDS

NAPHTHALENE PHOSPHINIC ACID $C_{10}H_7\text{P}(\text{OH})_2$ *Naphthyl phosphorous acid* [126°] SG 1377 (Schroder, B 12, 564) Formed by the action of water on its chloride Aggregates of small needles, sl sol cold water, almost insol HClAq Melts under water Reduces silver solution It is accompanied by an acid $(C_{10}H_7)_2\text{PO OH}$ [204°] which is insol water.

Chloride $C_{10}H_7\text{PCl}_2$ (above 360°) Formed by heating mercuric dinaphthyl $\text{Hg}(\text{C}_{10}\text{H}_7)_2$ with PCl_3 at 190° for several hours (Kelbe, B 9, 1051, 11, 1499) Liquid Combines with chlorine forming $C_{10}H_7\text{PCl}_4$

NAPHTHALENE PHOSPHONIC ACID $C_{10}H_7\text{PO}(\text{OH})_2$ [190°] SG 1440 Formed by the action of water on its tetrachloride (Kelbe) Long needles, v sol hot, sl sol cold, water When strongly heated it decomposes, with separation of carbon, into naphthalene and metaphosphoric acid— $\text{Ag}_2\text{A}''$ white pp

Chloride $C_{10}H_7\text{PCl}_4$ Formed from the compound $C_{10}H_7\text{PCl}_2$ and chlorine

NAPHTHALENE-STYRENE v NAPHTHYL ETHYLNE

NAPHTHALENE (a) SULPHINIC ACID $C_{10}H_7\text{SOH}$ Formed by digesting a solution of the chloride of naphthalene (a) sulphonic acid in ether with sodium amalgam (Gessner, B 9, 1500, cf Otto & Moraes, B 6, 860) White scales, v sol water, sl sol HClAq , m sol alcohol, sl sol ether Decomposed by HClAq at 180° into naphthalene and SO_2

Salts— KA' aq glistening scales — BA' 1' aq Slender needles S 5 at 14°, 2 at 100° — PBA' aq long branched needles — AGA' Soluble plates, not decomposed at 200°

Naphthalene (b)-sulphinic acid $C_{10}H_7\text{SO}_2\text{H}$ [105°] Formed in the same way as the preceding isomeride from naphthalene (b) sulphonic acid, and ppd by adding HCl to a solution of the Na salt as a white crystalline powder, m sol alcohol, ether, and water, almost insol HClAq Readily decomposed by HClAq at 150° into naphthalene and SO

Salts— KA' aq scales — BA' , glistening needles S 47 at 15°, 625 at 100° — CAa' 3aq white crystalline solid, v sol water and alcohol — MgA' 6aq scale, more sol alcohol than water

References—BROMO and CHLORO NAPHTHALENE SULPHONIC ACID

NAPHTHALENE (a) SULPHONIC ACID $C_{10}H_7\text{SO}_3\text{H}$ Formed, together with the (b)-isomeride, by dissolving naphthalene in H_2SO_4 At 160°–170° the chief product is the (b) acid, but at 80°–100° the product consists mainly of the (a) acid (Merz & Weith, B 3, 195, cf Faraday, P 7, 104, Berzelius, P 44, 377, Liebig & Wohler, P 24, 169, A 37, 197) Naphthalene is converted by $\text{Cl SO}_3\text{H}$ (1 mol) into a mixture of the (a) and (b) sulphonic acids (Armstrong, C J 24, 173) It may be separated from the (b) isomeride by means of the calcium or lead salts Crystalline and deliquescent, v e sol water, sol alcohol, sl sol ether Partially converted into the (b) isomeride by heating with H_2SO_4 at 100°, more completely at higher temperatures Decomposed into naphthalene and H_2SO_4 by heating with HClAq at 200° Oxidised by KMnO_4 in acid solution to phthalic acid (Beil-

stein & Kurbatoff, *C C* 1881, 359, *A* 202, 216) Alkaline KMnO_4 yields $\text{C}_6\text{H}_4(\text{CO}_2\text{H})\text{CO}_2\text{H}$ (Henriques, *B* 31, 1607) and phthalic acid. Bromine forms dibromo naphthalene and bromonaphthalene sulphonic acid

Salts— $\text{KA}'_2\text{aq}$ Plates (from alcohol) *S* 77 at 11° — $\text{CaA}'_2\text{aq}$ Plates (slowly decomposed at 80°) *S* 6 at 11° — $\text{BaA}'_2\text{aq}$ Plates *S* 115 at 10° (*M*), 113 at 15° , 476 at 100° (Regnault, *J pr* 12, 99)— $\text{PbA}'_2\text{aq}$ Plates *S* 37 at 10° — $\text{Pb}_2\text{OA}'_2$ — AgA' *S* 103 at 10°

Ethyl ether EtA' Formed by boiling the chloride (1 mol) with alcohol (2 mols) (Kimberly, *A* 114, 133) Viscid liquid, which slowly solidifies Insol water, miscible with alcohol and ether Decomposed on distillation, giving off SO_2 and naphthalene Aqueous or alcoholic potash merely saponifies it Water at 150° resolves it into alcohol, naphthalene, and H_2SO_4 Converted by PCl_5 at 160° into (a) chloro naphthalene and SOCl_2 (Carius, *A* 114, 145)

Chloride $\text{C}_{10}\text{H}_7\text{SO}_2\text{Cl}$ [66°] Glistening plates (from ether), v sol ether, CS_2 , and benzene (Maikopar, *Z* 1869, 710, Kimberly, *A* 114, 129) On passing chlorine into a solution in CS_2 , there is formed the tetrachloride $\text{C}_{10}\text{H}_6\text{Cl}_4\text{SO}_2\text{Cl}$, a thick liquid, sol CS_2 , chloroform, benzene, and ether, and converted by alcoholic potash into di chloro naphthalene sulphonic chloride (Widmann, *B* 12, 2228)

Amide $\text{C}_{10}\text{H}_7\text{SO}_2\text{NH}_2$ [150°] Prepared by warming the chloride with conc NH_4Aq Sol water, v sol alcohol and ether Its alcoholic solution gives with AgNO_3 a crystalline pp $\text{C}_{10}\text{H}_7\text{SO}_2\text{NHAg}$, v sol alcohol and ether, quickly blackening in light

Benzoyl derivative of the amide $\text{C}_{10}\text{H}_7\text{SO}_2\text{NHBz}$ [195°] Formed by heating the amide with BzCl (Kimberly) Minute four sided prisms (from alcohol) (Wolkoff, *Z* 1871, 422, *B* 5, 142) Insol water, m sol alcohol and ether Decomposed by boiling KOHaq into benzoic acid, NH_3 , and naphthalene sulphonic acid PCl_5 converts it into $\text{C}_{10}\text{H}_6\text{SO}_2\text{NCOClC}_6\text{H}_5$, which crystallises from ether in large four sided plates [94°] and is re converted by boiling water or alcohol into $\text{C}_{10}\text{H}_7\text{SO}_2\text{NHCOC}_6\text{H}_5$, and by ammonium carbonate into crystalline $\text{C}_{10}\text{H}_7\text{SO}_2\text{NC}(\text{NH}_2)_2\text{C}_6\text{H}_5$ Behaves like an acid, decomposing carbonates, and forming the salts— $\text{C}_{10}\text{H}_7\text{SO}_2\text{NKBz}$ prisms, v sol water and alcohol— $\text{CaA}'_2\text{aq}$ needles— BaA'_2 slender needles, sl sol water— AgA' minute needles

Anilide $\text{C}_{10}\text{H}_7\text{SO}_2\text{NHPH}$ [112°] Needles (Carleson, *Bl* [2] 27, 360)

(a) *Naphthalene* $\text{C}_{10}\text{H}_7\text{SO}_2\text{NHC}_6\text{H}_5$, [82°] Small needles (*C*)

Naphthalene (β)-sulphonic acid $\text{C}_{10}\text{H}_7\text{SO}_3\text{H}$ Produced by the action of H_2SO_4 upon phenyl (a) naphthyl ketone, the isomeric change resulting from the heat evolved in the reaction (Elbs & Steinike, *B* 19, 1966) Prepared by heating naphthalene (500 g) with H_2SO_4 (400 g) for 8 hours at 160° and purifying the acid by means of the Ca salt (Merz & Weith, *B* 3, 196) Non-deliquescent crystals Not decomposed by dilute HClAq at 200° Split up into naphthalene and H_2SO_4 by distilling with dilute H_2SO_4 at 135° in a current of superheated steam (Armstrong & Miller, *C J* 45, 148) Oxidised by alkaline KMnO_4 to $\text{C}_6\text{H}_4(\text{CO}_2\text{H})\text{CO}_2\text{H}$

(Henriques, *B* 21, 1607) KMnO_4 in neutral or acid solution yields phthalic acid (Beilstein & Kurbatoff, *C C* 1881, 359, *A* 202, 216) CrO_3 and dilute H_2SO_4 yield naphthoquinone sulphonic acid (*B* & *K*) The Ca salt yields phthalic acid on oxidation by CrO_3 Bromine water forms bromo naphthalene sulphonic acid

Salts— $\text{KA}'_2\text{aq}$ Plates (from water) or needles (from dilute alcohol) *S* 66 at 10° *S* (85 p.c. alcohol) 9— CaA'_2 Plates *S* 13 at 10° — $\text{BaA}'_2\text{aq}$ Plates *S* 345 at 10° — $\text{PbA}'_2\text{aq}$ Scales *S* 9 at 10°

Chloride $\text{C}_{10}\text{H}_7\text{SO}_3\text{Cl}$ [76°] Plates, less soluble in ether than the (a) isomeride (Maikopar) Reduced by HI to (β) di naphthyl disulphide [139°] (Cleve, *B* 21, 1100) Combines with chlorine (dissolved in CS_2) forming a tetrachloride $\text{C}_{10}\text{H}_6\text{Cl}_4\text{SO}_3\text{Cl}$, which separates from chloroform in colourless cubes [131°], v sol chloroform, CS_2 , and hot HOAc , insol water (Widmann, *B* 12, 959) The tetrachloride is converted by boiling alcoholic KOH into di chloro naphthalene (β) sulphonic acid

Amide $\text{C}_{10}\text{H}_7\text{SO}_3\text{NH}_2$ [212°] (Cleve, *Bl* [2] 25, 258) Small thin plates (from alcohol), sl sol water and ether

Ethylamide $\text{C}_{10}\text{H}_7\text{SO}_3\text{NHEt}$ [82.5°] (Carleson, *Bl* [2] 27, 360)

Anilide $\text{C}_{10}\text{H}_7\text{SO}_3\text{NHPH}$ [132°] Needles (a) *Naphthalene* $\text{C}_{10}\text{H}_7\text{SO}_3\text{NHC}_6\text{H}_5$ [177.5°] Needles

Naphthalene tetrahydride sulphonic acid $\text{C}_{10}\text{H}_8\text{SO}_3\text{H}$ Formed by heating naphthalene tetrahydride with H_2SO_4 for 3 hours at 40° (Graebe & Guye, *B* 16, 3030, Bamberger & Kirschelt, *B* 23, 1565) Crystals, v sol water and alcohol Decomposed by distillation with dilute H_2SO_4 and a current of superheated steam at 175° , or by dilute H_3PO_4 at 130° (Friedel & Crafts, *Bl* [2] 42, 66, *C R* 109, 95)— $\text{NaA}'_2\text{aq}$ Tables, v sol water— $\text{BaA}'_2\text{aq}$ Tables, sol hot alcohol, sl sol cold water

Naphthalene 'a'-disulphonic acid

$\text{C}_{10}\text{H}_6(\text{SO}_3\text{H})_2$ [22] Formed, together with about an equal quantity of the (β) isomeride and some of the (δ) isomeride, by heating naphthalene (1 pt) with H_2SO_4 (5 pts) for 4 hours at 160° (Ebert & Merz, *B* 9, 592, Armstrong, *B* 15, 204, cf Berzelius, *A Ch* [2] 65, 290, Laurent, *Compt Chim* 1849, 390) The acids may be partially separated by their Ca salts, that of the (β) acid crystallising out first, while that of the (a) acid is the most soluble The potassium salt of the (a) acid is more soluble than that of the (β) acid, and less soluble than that of the (δ) acid Long, very deliquescent needles, sl sol conc HClAq Gives di oxy naphthalene [186°] on fusion with KOH (cf Gress, *B* 13, 1959, Dusat, *C R* 64, 859, Darmstadter & Wichelhaus, *A* 152, 306) Water at 200° splits it up into naphthalene and H_2SO_4 Fusion with NaOH forms (β) naphthol (δ) sulphonic acid $\text{C}_{10}\text{H}_6(\text{OH})\text{SO}_3\text{H}$ (Weniger, *B* 20, 2906) PBr_3 forms (β)-di-bromo naphthalene Yields di-chloro-naphthalene [114°]

Salts— $\text{KA}'_2\text{aq}$ Transparent needles (from hot saturated solutions). *S* (of KA') 71 at 18° — $\text{NaA}'_2\text{aq}$ Glistening needles *S* (of NaA') 45.5 at 18° — $\text{CaA}'_2\text{aq}$ *S* 16 at 18° — $\text{CaA}'_2\text{aq}$ (from a rapidly-cooled hot saturated

solution) —BaA''2aq Long, broad needles S 12 at 19° —PbA''2aq Long needles, sol water

Chloride $C_{10}H_7(SO_2Cl)_2$ [158°] (E a M), [162°] (A) Obtained by heating the K salt with PCl_5 at 140° Plates (from benzene), or needles (from ether) S (benzene) 18.3 at 14°

Amide $C_{10}H_7(SO_2NH_2)_2$ [243°] Needles, m sol hot NH_4Aq and alcohol

Naphthalene β' -disulphonic acid

$C_{10}H_6(SO_3H)_2$ [231°] Almost the sole product obtained by heating a mixture of naphthalene (1 pt) and H_2SO_4 (5 pts) for 24 hours at 180° (Ebert a Merz) Formed also from its 'a'-isomeride by prolonged heating with H_2SO_4 at 180° Somewhat deliquescent plates Gives (β) naphthol 'β'-sulphonic acid and (β) di-oxy-naphthalene on fusion with potash Yields $C_{10}H_7Cl$ [135°]

Salts —NaA''aq aggregates of minute prisms S (of NaA'') 12 at 19° —K₂A'' Bushy groups of needles S 5.2 at 18° —CaA''aq S 6.2 at 18° When once dry this salt dissolves in water with great difficulty —BaA''aq —PbA''aq

Chloride $C_{10}H_6(SO_2Cl)_2$ [226°] S (benzene) 45 at 14° Tufts of small needles or thin plates, v sl sol $HOAc$ (difference from the (α)-isomeride), v sol benzene

Amide $C_{10}H_6(SO_2NH_2)_2$ Small needles, v sl sol NH_4Aq , almost insol alcohol, ether, benzene, and toluene Not melted at 305°

Naphthalene (γ) disulphonic acid

$C_{10}H_6(SO_3H)_2$ [14°] Formed by treating naphthalene (1 mol) with $ClSO_3H$ (2 mols) below 100° and also by sulphonating naphthalene with SO_3 (Armstrong, B 15, 204, Armstrong a Wynne, C J Proc 2, 230, 3, 42) Its Pb, Ca, and Ba salts resemble those of the (β) acid, dissolving very sluggishly in water On fusion with potash it yields di oxy naphthalene [260°] —NaA''2aq —K₂A''2aq plates, less soluble than the corresponding salt of the 'α' acid

Chloride $C_{10}H_6(SO_2Cl)_2$ [184°] Prisms (from benzene) Yields (γ) di chloro naphthalene [107°] on treatment with PCl_5

Naphthalene (δ) disulphonic acid

$C_{10}H_6(SO_3H)_2$ [13°] Also called (γ) Occurs in small quantity in the product of sulphonation of naphthalene by H_2SO_4 at 160° (Armstrong, B 15, 204) Formed by treating potassium naphthalene (β) sulphonate with $ClSO_3H$ (Armstrong a Wynne, C J Proc 2, 230) —BaA''4aq —NaA''7aq Formed also by heating naphthalene (β) sulphonic acid with fuming H_2SO_4 at 100° (Ewer a Pick, G P 45,229 [1887], Armstrong, C J Proc 4, 10) Long needles Gives dioxy naphthalene [135.5°] on fusion with potash Yields $C_{10}H_7Cl_2$ [49°] The potassium salt is more soluble than that of the (α) or (β) acid —K₂A''aq —Na₂A''4aq —BaA''4aq —CaA''4aq —PbA''3aq

Chloride $C_{10}H_6(SO_2Cl)_2$ [125°] Small prisms (from benzene) On treatment with PCl_5 it yields di chloro naphthalene [59°]

Naphthalene (1,2') disulphonic acid Formed from (α) naphthylamine disulphonic acid (Armstrong a Wynne, C J Proc 5, 136) Yields $C_{10}H_7Cl_2$ [62.5°] —K₂A''aq

Chloride [122.5°]

Naphthalene (1, 3) disulphonic acid Obtained from (1', 1, 3) or (2', 4') naphthylamine disulphonic acid by eliminating NH_2 (Armstrong

a Wynne, C J Proc 5, 18, 129) Yields $C_{10}H_7Cl_2$ [61°] —K₂A''2aq —BaA''4aq

Chloride [137°]

Naphthalene hexahydride (α)-di-sulphonic acid $C_{10}H_{12}(SO_3H)_2$ Formed by heating naphthalene hexahydride with H_2SO_4 and SO_3 , and separated from the (β)-isomeride by means of the K salt (Agrestini, G 12, 495) —K₂A'' (dried at 105°) Nodules, sol alcohol

Naphthalene hexahydride (β)-sulphonic acid $C_{10}H_{12}(SO_3H)_2$ Formed as above —K₂A'' 13aq Insol alcohol

Naphthalene trisulphonic acid $C_{10}H_5(SO_3H)_3$ [242°] Formed from naphthalene and $ClSO_3H$ (Wynne, C J Proc 3, 146) Yields $C_{10}H_7Cl_2$ [194°] —Na₂A''6aq

Naphthalene tetra-sulphonic acid

$C_{10}H_4(SO_3H)_4$ Formed by heating naphthalene with H_2SO_4 and P O, for three or four hours at 260° (Stenhofer, B 8, 1486, M 3, 111) Two isomeric acids are formed in the process, and when a solution of their Ba salts is slowly evaporated at 30° to 35° the salt of the acid here described crystallises out in striated prisms Prisms (containing 4aq), v e sol water, sl sol alcohol, insol ether Decomposes above 170°

Salts —K₂A''2aq non deliquescent needles —Na₂A''2aq (at 100°) —Na₂A''10aq very deliquescent prisms —Ba₂A''2aq prisms —Pb₂A''6aq crystalline pp —Cu₂A''12aq blue prisms —Ag₂A''24aq very soluble needles

References —Bromo, Chloro, Iodo- and Nitro NAPHTHALENE SULPHONIC ACIDS

NAPHTHAL-FLUORESCIN $C_{20}H_{14}O_8$..

$O < \begin{matrix} C_6H_4(OH) \\ C_6H_4(OH) \end{matrix} > C < \begin{matrix} O \\ O \end{matrix} > CO$ [308°] Formed by heating naphthalic anhydride and resorcin at 250° (Terrisse, A 227, 136) Yellow rhombic prisms Its alkaline solution is red, with splendid green fluorescence

Acetyl derivative $C_{20}H_{14}AcO_8$ [191°]

$C_{20}H_{14}AcO_8$ [120°] Needles (from acetone) Insol alkalis, but saponified slowly By treatment with PCl_5 it is converted into the chloride $C_{20}H_{14}O_8Cl_2$, which crystallises in scales, sol chloroform, acetone, or glacial acetic acid, hardly sol benzene, ether, or alcohol

Naphthal eosin $C_{20}H_{14}Br_2O_8$ [above 310°]

Formed by adding bromine to an alcoholic solution of naphthal fluorescein Its alkaline solutions are orange, with yellow fluorescence, and dye silk scarlet

NAPHTHALIC ACID v NAPHTHALENE DI CARBOXYLIC ACID

NAPHTHALIDES Alkyl derivatives of

NAPHTHYLAMINE (q v)

NAPHTHALIDINE SULPHONIC ACID v.

(1,4') NAPHTHYLAMINE SULPHONIC ACID

NAPHTHALIN v NAPHTHALENE

(β) **NAPHTHAMIDINE** $C_{10}H_9N$, C(NH) NH_2

White crystalline mass, obtained from its hydrochloride, which is prepared by the action of alcoholic NH_3 at 50° to 60° upon (β) naphthamido ether $C_{10}H_9N$, C(NH) OEt (Pinner a Klein, B 11, 1486) —B' HCl [226°] Needles

(α) **NAPHTHAMIDOXIM** $C_{11}H_{10}N_2O$.. $C_{10}H_9N$, C(NOH) NH_2 [149°] Obtained by mixing the nitrile of (α) naphthosic acid with hydroxylamine hydrochloride, Na_2CO_3 , and alcohol (Ekstrand, B 20, 223) Plates (from dilute alcohol), v sol alcohol, insol ligroin On boil-

The following are the physical properties of some of the members examined

	S.C. $\frac{16-18}{4}$	μ_a	μ_p	μ_v	μ_D	R _z
Decanaphthene .	7808	1 43066	1 43863	1 4427	1 433	77 2
Endecanaphthene .	8019	1 43883	1 4466	1 45156	1 441	84 2
Dodecanaphthene .	812	1 4438	1 45173	1 45646	1 44606	91 8
Tetradecanaphthene .	8215	1 44913	1 45706	1 4619	1 4514	107 1
Pentadecanaphthene .	8290	1 4544	1 46136	1 4661	1 4544	114 7

The naphthenes are obtained from the following fractions of Caucasian petroleum

	Boiling point
Octonaphthene $C_{18}H_{18}$	116–120°
Nonaphthene $C_{19}H_{18}$	135°–140°
Decanaphthene $C_{20}H_{18}$	152°–167°
Endecanaphthene $C_{21}H_{18}$	180°–185°
Dodecanaphthene $C_{22}H_{18}$	196°–197°
Tetradecanaphthene $C_{24}H_{18}$	240–241°
Pentadecanaphthene $C_{25}H_{18}$	246°–248°

(Markownikoff & Oglobine, *A Ch* [6] 2, 446)

Some of these hydrocarbons (e.g. octonaphthene and nonaphthene) occur in two isomeric forms. Octonaphthene on treatment with H_2SO_4 and HNO_3 yields tri nitro *m* xylene, and it is therefore probably *m* xylene hexahydride (Markownikoff & Spady, *B* 20, 1851). Nonaphthene (136°), S.G. 0.7667 is identical with pseudo cumene hexahydride, for it yields some tri nitro ψ cumene on nitration and ψ cumene sulphonic acid with excess of H_2SO_4 (Kononoff, *C C* 1887, 1133, *J R* 22, 4, 118). Heptanaphthene $C_{17}H_{14}$, also occurs in Caucasian petroleum and boils at 101° (Milkovskiy, *Bl* [2] 45, 182).

Hexanaphthene carboxylic acid $C_{16}H_{14}CO_2H$ *Hexahydro benzoic acid?* (216°) S.G. 1.84/1.84. 9503. Obtained from the oils of Baku by extracting with caustic soda, and purified by fractional distillation of its methyl ether (A-chau, *B* 23, 867). Thick colourless oil, smelling like valeric acid. Not solid at -10° . Not attacked by bromine in the cold. HNO_3 and $KMnO_4$ act slowly upon it. It slowly expels HCl from $CaCl_2$. Conc. H_2SO_4 dissolves it, and, on warming, decomposes it. Phosphoric acid decomposes it gradually in the cold.

Salts— KA . Very hygroscopic soap like mass, gradually becoming crystalline, v sol water and alcohol— NaA' . Flat hygroscopic prisms— CaA' 4aq. long pointed needles. A saturated solution becomes milky on heating and clear again on cooling— BaA' . Large thin plates, v sol alcohol, m sol water— BaA' , 2aq. Amorphous— CdA' . Pearly plates, sl sol cold water— AgA' . Curdy pp.

Methyl ether MeA' (167°) S.G. 1.84/1.84. 9055. Colourless liquid with sickly odour. **Chloride** $C_{16}H_{13}COCl$ (168°). Slowly decomposed by water.

Amide $C_{16}H_{13}CONH_2$ [123.5°]. Melts under water. Extremely thin pearly plates, m sol water, v sol other solvents.

Anilide $C_{16}H_{13}CONHPh$ [94°]. Long elastic needles.

NAPHTHENYL-AMIDINE v **NAPHTHANIDINE**

NAPHTHIDINE v **DIAMIDO DINAPHTHYL**
NAPHTHIL BENZOIN v **BENZOIN**

(β) NAPHTH IMIDO-ACETATE $C_{12}H_9NO_2$, $\pm C_{10}H_7C(NH)(OAc)$ [152°]. Prepared by boiling naphth imido isobutyl ether with Ac_2O (Pinner & Klein, *B* 11, 1487). White needles.

(β) NAPHTH-IMIDO-ISOBUTYL ETHER $C_{11}H_9NO_2$ $\pm C_{10}H_7C(NH)OC_2H_5$ [38°]. Prepared by the action of gaseous HCl on a solution of (β) naphthonitrile $C_{10}H_7CN$ in isobutyl alcohol (Pinner & Klein, *B* 11, 1487). Long white needles— $BHCl$. Decomposed by heat into isobutyl chloride and the amide of (β) naphthoic acid.

(β)-NAPHTHIMIDO ETHYL ETHER $C_{10}H_7C(NH)OEt$. Hydrochloride $BHCl$. Prepared by leading HCl into an alcoholic solution of the nitrile of (β) naphthoic acid (Pinner & Klein, *B* 11, 1485). Decomposed by heat into $EtCl$ and the amide of (β) naphthoic acid.

(α) NAPHTHINDOLE $C_{12}H_7N$ $\pm C_{10}H_5$ $\begin{smallmatrix} CH \\ \diagup \\ NH \end{smallmatrix} CH$ [175°]. Obtained by distilling its carboxylic acid (Schlieper, *A* 239, 229). Plates (from ligroin), v sol alcohol, ether, and benzene, sl sol hot water. Colours pine wood, in presence of HCl , bluish violet. Its solution in acetic acid gives a red colour with conc. HNO_3 , and a bluish green pp. on boiling with hydrogen peroxide. Its picrate crystallises in needles— $B'HCl$. Colourless pp., got by adding HCl to the solution in $HOAc$.

On boiling an alcoholic solution of (α) naphthindole with zinc dust and HCl there is formed a hydride which yields an oxalate melting at 166°.

(β) Naphthindole $C_{10}H_7$ $\begin{smallmatrix} CH \\ \diagup \\ NH \end{smallmatrix} CH$ (above 360°) at 760 mm, (222° V at 18 mm).

Formation—1. By heating ethylidene (β) naphthyl hydrazine with $ZnCl_2$, the yield being small—2. By heating the (β) naphthyl hydrazide of pyruvic ether with $ZnCl_2$ at 195°, the intermediate body being (β) naphthindole carboxylic acid (Schlieper, *A* 236, 178).

Properties—Yellow oil, v sol alcohol, ether, benzene, and $HOAc$, sl sol ligroin. Yields a picrate, crystallising in dark red needles, and a nitrosamine. Conc. HCl aq. solidifies the oil. It colours pine wood, in presence of HCl , violet. Oxidising agents yield a solid product.

(α) NAPHTHINDOLE CARBOXYLIC ACID $C_{12}H_7NO_2$, $\pm C_{10}H_5$ $\begin{smallmatrix} CH \\ \diagup \\ NH \end{smallmatrix} CO_2H$ [202°].

Formed by the action of boiling alcoholic potash upon its ether, which is obtained by heating the (α) naphthyl hydrazide of pyruvic ether with $ZnCl_2$ (Schlieper, *A* 239, 232). Silvery laminae (from water), v sl sol hot water, v sol alcohol and ether. Decomposes above 210° into CO_2 and (α)-naphthindole.

Ethyl ether EtA' [170°]

(β)-Naphthindole-carboxylic acid $C_{10}H_7NO_2$,
 $C_{10}H_6 < \begin{smallmatrix} CH \\ NH \end{smallmatrix} > CO_2H$ [226°] Obtained by

saponifying its ether, which is formed when the (β)-naphthyl hydrazide of pyruvic ether is heated with $ZnCl_2$ (Sohlhoyer, *A* 236, 181) Colourless plates, insol water, sl sol ether, v sol hot HOAc and alcohol Does not colour pine-wood Bromine gives a yellow pp

(α)-NAPHTHINDOLE SULPHONIC ACID

$C_{10}H_6 < \begin{smallmatrix} NH \\ CH \end{smallmatrix} > CSO_3H$ (?) The sodium salt of

this acid, formed by the action of (α) naphthylamine on the compound of glyoxal with $NaHSO_3$, is converted by mineral acids into (α)-naphthoxindole (Hinsberg, *B* 21, 116)

β -Naphthindole sulphonic acid

$C_{10}H_6 < \begin{smallmatrix} CH \\ NH \end{smallmatrix} > CSO_3H$ or $C_{10}H_6 < \begin{smallmatrix} N \\ CH_2 \end{smallmatrix} > CSO_3H$

The sodium salt of this acid is obtained by the action of (β)-naphthylamine on the compound of glyoxal and $NaHSO_3$ (Hinsberg, *B* 21, 113) it is stable towards alkalis, but converted by mineral acids into (β) naphthoxindole

NAPHTHIONIC ACID *v* (1, 4) NAPHTHYLAMINE SULPHONIC ACID.

NAPHTHISATIN *v* ISATIN

NAPHTHOACRIDINE *v* NAPHTHACRIDINE

NAPHTHOBENZALDOXIM so called, *v* Oxim of NAPHTHOALDEHYDE

NAPHTHOBENZYL ALCOHOL *v* NAPHTHYL AERINOL

NAPHTHOBENZYLAMINE *v* NAPHTHYL AERINYL AMINE

NAPHTHO-CINNAMIC ACID so called, *v* NAPHTHYL ACRYLIC ACID

NAPHTHOCUMARIC ACID *v* Oxy NAPHTHYL-ACRYLIC ACID

NAPHTHOCUMARIN *v* Anhydride of Oxy-NAPHTHYL ACRYLIC ACID

NAPHTHOXYAMIC ACID $C_{10}H_7N_2O_5$ The potassium salt of this acid is prepared by boiling di nitro naphthalene with alcoholic KCy (Muhlhauser, *A* 141, 214) The free acid is a brownish black mass, insol ether, nearly insol water, m sol alcohol —KA' aq Dark mass, with coppery lustre, forming a fine blue solution in hot water and hot alcohol The barium salt is a dark blue pp, and explodes when heated The silver salt is insol water, has a bronze lustre, and explodes violently when heated

NAPHTHO-HYDROQUINONE *v* HYDRO-NAPHTHOQUINONE

(α) NAPHTHOIC ACID $C_{10}H_7O_2$ *vs* $C_{10}H_6CO_2H$ Naphthalene (α) carboxylic acid Mol w 172 [162°] H C *v* 1,232,000 H C p 1,232,600 H F 77,400 (Stohmann, Kleber, a Langbein, *J pr* [2] 40, 137)

Formation —1 By saponifying its nitrile by boiling alcoholic potash (Merz, *Z* [2] 4, 34, 5, 396) or conc HClAq (Hofmann, *B* 1, 89) —2 By heating a mixture of bromo-naphthalene and $Cl CO_2Et$ with sodium amalgam at 110° for several hours, and saponifying the resulting ether with alcoholic potash (Egish, *C R* 69, 360) —3 By fusing potassium naphthalene (α)-sulphonate with sodium formate (V Meyer, *A* 156, 274) —4 By boiling with potash its amide which is formed by the action of $ClCO NH_2$ on

naphthalene in presence of $AlCl_3$ (Gattermann, *A* 244, 56)

Preparation —A mixture of 3 pts of sodium naphthalene (α) sulphonate and 2 pts of potassium ferrocyanide (both quite dry) is distilled from an iron tube, the nitrile is rectified and saponified by heating it with an equal volume of HCl to 200°, the yield is 25 p c of the sulphonate (Boessneck, *B* 16, 639)

Properties —Needles (from dilute alcohol), v sl sol water, m sol hot alcohol Yields naphthalene on distillation with lime, and phthalic acid on oxidation by CrO_3 in HOAc

Salts —BA', 4aq Needles, sl sol water —CaA', 2aq S 1.08 at 15° —AgA'

Ethyl ether EtA' (309° cor)

Chloride $C_{10}H_6COCl$ (297.5°) From the acid (4 pts) and PCl_5 (5 pts) Hydroxylamine converts it into $C_{10}H_6CO NH OH$ [187°] and ($C_{10}H_6CO$), NOH [150°] (Ekstrand, *B* 20, 1358) Reacts with potassium (β) naphthoate, forming ($\alpha\beta$) naphthoic anhydride $C_{10}H_6COOCO C_{10}H_6$ (Hausmann, *B* 9, 1515)

Amide $C_{10}H_6CONH_2$ [202°] Formed from the chloride and NH_3 , or from the nitrile and alcoholic soda (Hofmann, Ekstrand, *J pr* [2] 38, 146) Prepared by passing dry gaseous cyanic acid and HCl into a solution of naphthalene in CS, containing $AlCl_3$ heated on the water bath (Gattermann a Rossolymo, *B* 23, 1197) Colourless monoclinic tables (from alcohol) (Bamberger a Philip, *B* 20, 241)

Anilide $C_{10}H_6CONHPh$ [160°] White silky crystals

(α) Naphthalide $C_{10}H_6CONHC_6H_5$ [244°]

Anhydride ($C_{10}H_6CO$) O [145°] Obtained by distilling calcium naphthoate with naphthyl chloride (Hofmann) Prisms (from boiling benzene)

Nitrile $C_{10}H_6CN$ (α) Cyano naphthalene Mol w 153 [37°] (298° cor) *Formation* —1 By distilling naphthylamine with oxalic acid and heating the resulting formyl derivative of naphthylamine with HClAq (Hofmann, *A* 142 121, *B* 1, 39) —2 By distilling potassium naphthalene (α) sulphonate with KCy (Merz, *Bl* [2] 9, 335, 10, 47) or K_2FeCy_6 (Boessneck, *B* 16, 639, Hausmann, *B* 9, 1514) —3 By passing a mixture of cyanogen and naphthalene through a red hot tube (Merz a Weith, *B* 10, 746) —4 By heating di naphthylthiourea with copper powder (Weith, *B* 6, 967) —5 By heating tri (α) naphthyl phosphate with KCy, the yield being 20 p c of the theoretical (Heim, *B* 16, 1779) —6 By boiling the formyl derivative of (α)-naphthylamine with zinc dust in a current of hydrogen, the yield being 11 p c (Gasiorowski a Merz, *B* 18, 1006) —7 By the action of cuprous cyanide on (α) diazo naphthalene salts (Bamberger a Philip, *B* 20, 257)

Properties —Needles Reduced in alcoholic solution by sodium to the tetrahydride of naphthyl carbonylamine $C_{10}H_{11}CH_2NH_2$ On heating with hydroxylamine it yields $C_{10}H_{11}CONH_2$ and naphthamidoxim [149°]

(β)-Naphthoic acid $C_{10}H_7CO_2H$ *Isonaphthoic acid* [185° cor] (above 300°) H C *v* 1,227,800 H C p 1,228,400 H F 81,600 (Stohmann, Kleber, a Langbein, *J pr* [2] 40, 137)

Formation —1 By saponification of its

nitrile, which is formed by heating potassium naphthalene (β) sulphonate with KCl (Merz a Muhlhauser, *Z* [2] 5, 70) — 2 By boiling (β)-naphthyl naphthalene with conc HNO_3 (Giamocian, *B* 11, 272) — 3 By oxidising (β) naphthoic aldehyde with KMnO_4 (Schulze, *B* 17, 1530)

Properties — Long white needles (from hot water), sl sol hot water, v sol alcohol and ether Yields naphthalene on distillation with baryta Oxidised to phthalic acid by CrO_3 in HOAc

Salts — $\text{NaA}' \frac{1}{2}\text{aq}$ (Vieth, *A* 180, 314) Small tables, v e sol water — $\text{KA}' \frac{1}{2}\text{aq}$ — $\text{BaA}' 4\text{aq}$ — Needles (from hot water) *S* 07 at 15° — $\text{CaA}' 3\text{aq}$ *S* 055 at 15° — $\text{MgA}' 5\text{aq}$ — AgA'

Methyl ether MeA' [77°] (290°) *H.F.* 70,600 (Stohmann, *J pr* [2] 40, 353)

Ethyl ether EtA' (309°)

Chloride $\text{C}_{10}\text{H}_7\text{COCl}$ [43°] (305°) Formed from the acid and PCl_5 (Vieth, cf Grucarevic a Merz, *B* 6, 1242) Hydroxylamine solution at 100° attacks it, forming the compounds $\text{C}_{10}\text{H}_7\text{CO NH OH}$ [168°], and $(\text{C}_{10}\text{H}_7\text{CO})_2\text{NOH}$ [171°] (Ekstrand, *B* 20, 1359) The (β) naphthyl hydroxylamine reacts with (α) naphthoyl chloride forming ($\alpha\beta$) di naphthoyl hydroxylamine [160°] With urea it forms di naphthyl-urea [215°]

Anhydride $(\text{C}_{10}\text{H}_7\text{CO})_2\text{O}$ [134°] Plates composed of needles (Hausmann, *B* 9, 1515)

Amide $\text{C}_{10}\text{H}_7\text{CONH}_2$ [192°] Formed by warming the chloride with powdered ammonium carbonate Formed also by heating the nitrile with alcoholic potash (Leone, *G* 14, 120) Plates

Anilide $\text{C}_{10}\text{H}_7\text{CONHPh}$ [170°] Small plates (from benzene)

***p* Toluide** $\text{C}_{10}\text{H}_7\text{CO NHC}_6\text{H}_5$ [191°] Silky needles

(α) **Naphthalide** $\text{C}_{10}\text{H}_7\text{CO NH C}_{10}\text{H}_7$ [157°] Very small needles, sl sol ether

Nitrile $\text{C}_{10}\text{H}_7\text{CN}$ [66.5°] (305° col) Formed by reactions like those used for preparing its (α) isomeride (*v supra*) White mass, nearly insol water, v sol alcohol and ether When its alcoholic solution is saturated with HCl there is formed the hydrochloride of naphthimido ethyl ether (*q v*) (Pinner a Lohmann, *B* 11, 1485) By exhaustive chlorination with SbCl_5 it yields per chloro-benzene (Merz a Weith, *B* 16, 2887)

References — AMIDO-, BROMO, BROMO NITRO, CHLORO, CHLORO NITRO-, NITRO-, OXY, and GHIU, NAPHTHOIC ACIDS, and AMIDO-, and BROMO-AMIDO NAPHTHOIC ANHYDRIDE

(α) **NAPHTHOIC ACID TETRAHYDRIDE** $\text{C}_{10}\text{H}_{11}\text{CO}_2\text{H}$ [128°] Formed, together with the amide, by heating its nitrile with alcoholic pot ash for 5 or 6 hours at 165° (Bamberger a Bordt, *B* 23, 629) Slender dendritic prisms, v sol alcohol, sl sol cold water — CuA'_2 bluish green pp — PbA'_2 needles grouped in stars — AgA' flaky pp

Amide $\text{C}_{10}\text{H}_{11}\text{CONH}_2$ [162°] Flat needles, v sol boiling water

Nitrile $\text{C}_{10}\text{H}_{11}\text{CN}$ (278°) at 721 mm Formed from (α)-naphthylamine tetrahydride by Sandmeyer's reaction, and purified by steam-distillation

(α) **NAPHTHOIC ALDEHYDE** $\text{C}_{10}\text{H}_7\text{CHO}$ • *Naphthobenzaldehyde* Mol w 156 (292° cor)

Formed from naphthyl carbinol by oxidation with chromic acid mixture (Bamberger a Loder, *B* 21, 258) Pale yellow viscid liquid, with faint aromatic odour Conc HNO_3 at -5° yields a nitro-derivative [136°] It forms with NaHSO_3 a compound crystallising in glistering leaflets The phenyl hydrazide [185°] forms yellow leaflets, which turn red on exposure to light Aniline forms $\text{C}_{10}\text{H}_7\text{CH NPh}$ [71°], *o* toluidine yields $\text{C}_{10}\text{H}_7\text{CH NC}_6\text{H}_3$ [59°], *p* toluidine produces $\text{C}_{10}\text{H}_7\text{CH NC}_6\text{H}_4$ [93°], and (α)-naphthylamine $\text{C}_{10}\text{H}_7\text{CH NC}_6\text{H}_4$ [117°]

Oxim $\text{C}_{10}\text{H}_7\text{CH NOH}$ • *Naphthobenzald-oxim* [98°] Colourless needles (from alcohol), sl sol water (Brandes, *B* 22, 2151)

(β)-**Naphthoic aldehyde** $\text{C}_{10}\text{H}_6\text{CHO}$ [61°] Formed by distilling a mixture of calcium (β) naphthoate and calcium formate at a high temperature (Battershall, *Z* [2] 7, 292, 673, *A* 168, 116) Formed also by heating ω chloro (β) methyl-naphthalene ($\text{C}_{10}\text{H}_7\text{CH}_2\text{Cl}$) with lead nitrate (Schulze, *B* 17, 1530), and by oxidising (β)-naphthyl carbinol (Bamberger a Brekmann, *B* 20, 1118) Silvery plates, volatile with steam Reduces ammoniacal AgNO_3 , Alcoholic NH_3 converts it into the hydramide ($\text{C}_{10}\text{H}_7\text{CH}_2\text{N}_2$) [146°–150°]

(α)-**NAPHTHOL** $\text{C}_{10}\text{H}_7\text{OH}$ Mol w 144, [94°] (279°) *SG* 4 1224 (Schroder, *B* 12, 1613) *R₂₀* 76 97 in an 86 p c alcoholic solution (Kanonnikoff, *J pr* [2] 31 348), 75 25 (Nasini a Bernheimer, *G* 14, 153) Occurs in anthracene oil obtained from coal tar (K E Schulze, *A* 227, 150)

Formation — 1 By the action of nitrous acid on (α) naphthylamine (Griess, *J* 1866, 460) 2 From naphthalene (α) sulphonic acid by pot ash fusion (Eller, *A* 152, 275) — 3 By heating γ phenyl isocrotic acid $\text{CHPh CH CH}_2\text{CO}_2\text{H}$ at its boiling point for 5 or 10 minutes, water being eliminated (Fittig a Erdmann, *B* 16, 43, *A* 227, 242)

Preparation — By heating naphthalene (1 pt) with H_2SO_4 (2 pts) at 60° to 70° , and fusing the sodium salt of the (α) sulphonic acid formed with three times its weight of NaOH

Properties — Short trimetric prisms (Léger, *C R* 111, 110), sl sol hot water, v sol alcohol, ether, and benzene Volatile with steam Pine wood, moistened with an aqueous solution of (α) naphthol, and then with HClAq , turns green in sunlight, and finally reddish brown Bleaching powder gives, in a solution of (α)-naphthol, a dark violet colour, and ultimately violet flakes, which are turned brown by heating, but are decolourised by ammonia FeCl_3 gives a milky pp, soon becoming violet flakes of (α)-dinaphthol (α) Naphthol is a powerful antiseptic, more powerful in this respect than (β)-naphthol, although less poisonous (Maximovitch, *C R* 106, 366)

Reactions — 1 On boiling in the air, or on heating in sealed tubes at 350° – 400° , it yields dinaphthylene oxide ($\text{C}_{10}\text{H}_6\text{O}$) (Merz a Weith, *B* 14, 196) — 2 Heated to 270° for 8 hours with four times its weight of ammoniacal CaCl_2 (damp) at yields about 74 p c of (α) naphthylamine If ammoniacal ZnCl_2 is used instead of CaCl_2 , the product is 60 p c of ($\alpha\alpha$) dinaphthylamine (Benz, *B* 16, 14, cf Merz a Weith, *B* 14, 2844). — 3 Aniline and CaCl_2 yield, on heat-

ing, a little phenyl naphthyl-amine (M a W) — 4 *Ammonium acetate* at 270° yields the acetyl derivative of (a)-naphthylamine (Calm, B 15, 615) — 5 *Ammonium formate* at 250° yields CO and (a) naphthylamine — 6 When (a) naphthol (2 pts) is heated with anhydrous oxalic acid (1 pt) and conc H_2SO_4 (1 pt) at 130°, there is formed a compound $\text{C}_{10}\text{H}_8\text{O}_2$, crystallising from chloroform in pale rose coloured leaflets, insol alcohol and ether, converted by potash fusion into the acid $\text{C}_{10}\text{H}_6\text{O}_4$ (crystallising in golden green tables), and yielding the halogen derivatives $\text{C}_{10}\text{H}_5\text{Cl}_2\text{O}$ and $\text{C}_{10}\text{H}_4\text{Br}_2\text{O}$, both of which crystallise in the monoclinic system (Hong, M 1 251) — 7 Boiling with KClO_4 and HCl yields di chloro naphthoquinone (Wichelhaus, A 152, 801) — 8 *Chlorine*, acting on (a) naphthol dissolved in HOAc , forms $\text{C}_{10}\text{H}_7\text{Cl}_2\text{OH}$ [108°] and $\text{C}_6\text{H}_5\text{C}(\text{CO CCl})\text{CH}$ [121°] When the liquid is

kept cold the product is $\text{C}_6\text{H}_5\text{C}(\text{CO CCl})\text{CH}$ [157°] (Zincke a Kegel, B 21, 1030, 1044) — 8 PCl_5 at 100° forms di naphthyl phosphate At 150° it yields (a) chloronaphthalene (Oehler, B 15, 312) — 9 Conc HClAq at 200° yields di naphthyl oxide — 10 *Aluminium* and *iodine* yield aluminium (a)-naphthol (Gladstone a Tribe, G 41, 16) On distillation of this body there is formed a dinaphthyl [189°] — 11 *Iodine* and *potash* yield a dark-violet compound (Messinger a Fortmann, B 22, 2322) — 12 Somewhat more readily oxidised by KMnO_4 than (b) naphthol (Dreyfus, C R 105, 523) — 13 H_2SO_4 yields mono and di-sulphonic acids In presence of HOAc it also forms oxy naphthyl methyl ketone 14 HCl , acting on a mixture of (a) naphthol and aldehyde below 0° forms $\text{CH}_3\text{CH}(\text{C}_{10}\text{H}_7\text{OH})_2$ (Claus a Trauer, B 19, 3004) — 15 *Di-chloro ether* $\text{C}_6\text{H}_5\text{Cl}_2\text{OEt}$ yields amorphous $\text{C}_{10}\text{H}_8\text{O}_2$ or $\text{C}_6\text{H}_5(\text{C}_{10}\text{H}_7\text{OH})_2$ (Wischneus, A 243, 165) — 16 *Benzoic aldehyde* yields $\text{C}_6\text{H}_5\text{CH}(\text{C}_{10}\text{H}_7\text{OH})_2$ which turns brown in air (Claisen, B 19, 3316) 17 When boiled with *phthalic anhydride* it yields naphthol phthalate $\text{C}_{10}\text{H}_7\text{O}_4$ (Grabowski, B 4, 661, 725, 6, 1065) — 18 *Pyromellitic anhydride* (1 mol), heated with (a)-naphthol (4 mols), at 300° yields the compounds $\text{C}_{10}\text{H}_8\text{O}_6$ and $\text{C}_{10}\text{H}_6\text{O}_8$ [265°] When the two bodies are heated together in other proportions at 250° there are formed the compounds $\text{C}_{10}\text{H}_8\text{O}_6$, $\text{C}_{10}\text{H}_{12}\text{O}_7$, and $\text{C}_{10}\text{H}_{16}\text{O}_8$ (Grabowski) — 19 *Maleic anhydride* and ZnCl_2 at 160° forms $\text{C}_{10}\text{H}_8\text{O}_4$ and an acid $\text{CO}_2\text{HCHCHCO}_2\text{H}$, $\text{C}_{10}\text{H}_7\text{OH}$ [90°] (Burekhardt, B 18, 2868) — 20 *Benzotrifluoride* at 100° forms $\{\text{C}_6\text{H}_5\text{C}(\text{C}_{10}\text{H}_7\text{OH})_2\}_2\text{O}$, a reddish-brown powder, insol water, sol alcohol and ether (Doebner, A 257, 58)

Picric acid compound
 $\text{C}_{10}\text{H}_7\text{OC}_6\text{H}_2(\text{NO}_2)_3\text{OH}$ [190°] Orange needles (Marchetti, G 12, 503)

Compound with camphor
 $\text{C}_{10}\text{H}_7\text{OC}_6\text{H}_{10}\text{O}$ SG Δ 1 0327 $[\alpha]_D = -10.5^\circ$ Syrupy liquid, not solid at -16° Not sensibly decomposed by water (Léger, C R 111, 110)

Acetyl derivative $\text{C}_{10}\text{H}_7\text{OAc}$ [49°] Formed by heating (a) naphthol with AcCl (Schaeffer, B 2, 131) Formed also by heating (a) naphthol with HOAc at 200° (Graebe, A 209, 151), or with Ac_2O and NaOAc (Tassinari, G 10, 471) Large crystals, sol alcohol and ether

Split up into (a) naphthol and acetic acid by distillation with steam On oxidation with CrO_3 in cold HOAc it yields *c oxy phthalic acid*, and three compounds melting at 114°, 119°, and 121° respectively (Miller, B 14, 1601)

Benzoyl derivative $\text{C}_{10}\text{H}_7\text{OBz}$ [56°] Crystals, v sol ether (Maikopar, Z 1869, 216)

Methyl ether $\text{C}_{10}\text{H}_7\text{OMe}$ (270°) (V), (258°) (S), (266°) (M), (265°) (H), (269°) (V) (N a B) SG Δ 1 0974 (S), Δ 1 0964 (Nasim, G 15, 84) Prepared by the action of MeCl on sodium (a) naphthol (Vincent, Bl [2] 40, 106) or of MeI (60 g), KOH (25 g), and MeOH on (a) naphthol (60 g) (Staedel, A 217, 42) Formed also by heating (a) naphthylamine with methyl alcohol and ZnCl_2 at 200° (Hantzsch, B 13, 1347), and by heating MeOH with (a) naphthol and HCl (Manchetti, G 9, 545) Prepared by boiling (a) naphthol for several hours with MeOH and ZnCl_2 (Green, *priv com*) Liquid, smelling like orange blossom, sol alcohol, ether, and benzene With HNO_3 it gives a tri-nitro derivative [128°] With picric acid it forms a compound crystallising in red needles, decomposed by alcohol

Ethyl ether $\text{C}_{10}\text{H}_7\text{OEt}$ (281° cor) Formed from (a) naphthol, KOH , and EtI in alcoholic solution (Schaeffer, A 152, 286), or from (a) naphthol (25 g), MeOH (25 g), and H_2SO_4 (10 g) at 125° (Gattermann, A 244, 72) Heavy oil Converted by HNO_3 into a tri nitro derivative $\text{C}_{10}\text{H}_4(\text{NO}_2)_3\text{OEt}$ [148°] (Strädel, B 14, 899)

Propyl ether $\text{C}_{10}\text{H}_7\text{OC}_3\text{H}_7$ (299° cor) SG Δ 1 0447 (Nasim a Bernheimer, G 15, 84)

Ethylene ether $(\text{C}_{10}\text{H}_7\text{O})\text{CH}_2$ [126°] Plates (from alcohol) (Koelle, B 13, 190b)

(a) *Naphthyl ether v Di naphthyl oxide*
Nitroso derivatives v Oxim of (a) and (b) naphthoquinone

(b) *Naphthol* $\text{C}_{10}\text{H}_7\text{OH}$ Mol w 144 [123°] (286°) SG Δ 1 217 (Schroder, B 12, 161) n_D^{20} 76.43 in an 11.18 p.c alcoholic solution (Kanonnikoff, J pr [2] 31, 348) S 02, S (20 p.c alcohol) 2 (Bouchard, C R 105, 702) Occurs in coal tar (K Schulze, A 227, 150). Formed by the action of nitrous acid on (b)-naphthylamine (Liebermann, A 183, 268) Prepared by fusing sodium naphthalene (b) sulphinate (1 pt) with NaOH (2 pts) at 300° (Schaeffer, A 152, 282)

Properties — Plates or tables, sol hot water, v sol alcohol, ether, chloroform, and benzene Readily sublimates May be distilled by superheated steam Powerfully antiseptic (Bouchard, Maximovitch, C R 106, 1441) Pine wood dipped in an aqueous solution of (b) naphthol and then in HClAq becomes green on exposure to daylight Bleaching powder colours a solution of (b)-naphthol slightly yellow, but the colour is destroyed by excess of bleaching powder, on adding ammonia and warming, yellowish flakes separate FeCl_3 colours its solution slightly green, and, after a time, causes deposition of white flakes (of dinaphthol), which become brown on heating Its solution in NaOHaq yields with mercuric chloride a pp $(\text{C}_{10}\text{H}_7\text{O})_2\text{HgHgCl}_2.4\text{aq}$ (Pouchet, C R 106, 276). It may be estimated by adding standard iodine

solution to its solution in warm aqueous NaOH (Messinger & Vortmann, *B* 23, 2754)

Reactions —1 Alkaline potassium permanganate oxidises it to α -carboxy cinnamic acid $\text{CO}_2\text{H}\cdot\text{C}_6\text{H}_4\cdot\text{CH}=\text{CH}\cdot\text{CO}_2\text{H}$ [184°], and an acid $\text{C}_{10}\text{H}_7\text{O}_4$ [281°] which forms the salts $\text{BaA}'\cdot 7\text{aq}$ and AgA' , an ether EtA' [123°] and a dihydride $\text{C}_{10}\text{H}_7\text{O}$ [224°] (Ehrlich & Benedikt, *M* 9, 527, 10, 115) When (8) naphthol is boiled with P_2O_5 , or even by itself in presence of air, it forms di (8) naphthylene oxide —2 The dry distillation of calcium (8) naphthylate $\text{Ca}(\text{OC}_{10}\text{H}_7)_2$ yields di (8) naphthylene oxide, naphthalene, (8) naphthol, and a compound $\text{C}_{10}\text{H}_7\text{O}$ (?) [300°–305°] (Niederhäusern, *B* 15, 1122) —3 Zinc chloride forms, on heating, isodinnaphthylene oxide —4 Chloride of sulphur in presence of CS_2 , or benzene forms $(\text{HO}\cdot\text{C}_{10}\text{H}_7)\cdot\text{S}$ and $(\text{HO}\cdot\text{C}_{10}\text{H}_7)_2\cdot\text{S}_2$ (Tassinari, *G* 17, 94, Onufiovitch, *B* 23, 3356) —5 Boiling with sulphur and caustic soda solution forms $(\text{HO}\cdot\text{C}_{10}\text{H}_7)_2\cdot\text{S}_2$ [210°] and a compound melting at 170° (Lange, *B* 21, 260) —6 By heating (8) naphthol (150 g) with aluminium (10 g) as long as hydrogen comes off there is formed a mixture of (8) naphthol and aluminium (8)-naphthylate which when distilled yields di (8)-naphthyl oxide (Gladstone & Tribe, *C J* 41, 15) —7 When heated in sealed tubes with ammonia, NH_4Cl , acetamide, ammonium acetate or ammonium formate, it yields (8) naphthylamine. Heated to 270°–280° for 8 hrs with four times its weight of ammoniacal CaCl_2 (damp) it yields 80 p.c. of (8) naphthylamine and 12 p.c. of (88) dinaphthylamine. If ammoniacal ZnCl_2 is used instead of CaCl_2 , the product is 4 p.c. of (8) naphthylamine and 82 p.c. of (88)-dinaphthylamine (Benz, *B* 16, 9) —8 Reacts with nitroso-dimethyl aniline forming (8) naphthol violet $\text{C}_{10}\text{H}_7\text{N}_2\text{O}$ (Meldola, *C J* 39, 37), which is converted by heat into 'cyanamine', a blue colouring matter $\text{C}_8\text{H}_7\text{N}_2\text{O}_2$ (Witt, *B* 23, 2247) —9 Di-chloro quinonimide $\text{C}_8\text{H}_4(\text{NCl})_2$ forms a red colouring matter $\text{HN}\cdot\text{C}_8\text{H}_7\cdot\text{N}$ which forms a blue solution in conc. H_2SO_4 (Nietzki & Otto, *B* 21, 1714) The free base is a yellow pp., v. sol. alcohol and ether, and is converted by heat into a greenish blue colouring matter, greatly resembling 'cyanamine' (*v. supra*) —10 Nitrogen iodide forms iodo (8) naphthol, a substance melting at 223°, and an amorphous body (Willgerodt, *J pr* [2] 37, 446) —11 Di-chloro di-ethyl oxide $\text{CH}_3\text{CH}_2\text{CHCl}\cdot\text{OEt}$ yields $\text{C}_{12}\text{H}_{17}\text{ClO}$ [174°] crystallising in plates, insol. water, sol. alcohol and HOAc (Wislicenus, *A* 243, 169) —12 When chlorine is passed into a solution of (8) naphthol in HOAc and the product is treated with an excess of SnCl_2 there is obtained $1/2[\text{C}_{10}\text{H}_7\text{Cl}\cdot\text{OH}]$ [71°] (Zincke, *B* 21, 3284) This chloro-(8) naphthol when dissolved in HOAc or chloroform and treated with chlorine yields $\text{C}_8\text{H}_7\cdot\text{C}(\text{Cl})_2\cdot\text{CO}$ (Zincke, *B* 21, 3540) When (8) naphthol dissolved in HOAc is treated with excess of chlorine there is formed the compound $\text{C}_8\text{H}_7\cdot\text{C}(\text{Cl})_2\cdot\text{CO}$ crystallising in plates [103°] or needles [102°] and converted by Na_2CO_3 into chloro-(8) naphthoquinone [172°] (Zincke, *B* 21, 3550) When chlorine is passed into a well-cooled 10 p.c. solution of (8)-naphthol, there is

formed $\text{C}_8\text{H}_7\cdot\text{C}(\text{Cl})_2\cdot\text{CO}$ which crystallises from hot alcohol in thick yellow needles [96°] (Z) —13 Chloroform in presence of a small quantity of aqueous NaOH forms on boiling crystalline $\text{C}_{12}\text{H}_{11}\text{O}_2$ or $\text{C}_{10}\text{H}_7\cdot\text{C}(\text{OH})_2\cdot\text{C}(\text{OH})_2$, its anhydride $\text{C}_{12}\text{H}_{11}\text{O}_2$, an aldehyde $\text{C}_{10}\text{H}_7(\text{OH})\cdot\text{CHO}$, an alcohol $\text{C}_8\text{H}_7\text{O}$, and resins (Ioussseau, *C R* 94, 133, 95, 30, 232) —14 An acetic acid solution of aldehyde slowly reacts in the cold forming the di (8) naphthyl ether of acetic orthoaldehyde $\text{CH}_3\cdot\text{CH}(\text{OC}_6\text{H}_7)_2$ [201°] (Claisen, *B* 19, 3318) —15 A few drops of HCl added to a cold solution of benzoic aldehyde and (8) naphthol in HOAc forms $\text{C}_8\text{H}_7\cdot\text{CH}(\text{OC}_6\text{H}_7)_2$ [205°] converted by heating with HOAc and hydrochloric acid solution into $\text{C}_8\text{H}_7\cdot\text{CH}(\text{C}_6\text{H}_5)_2\cdot\text{O}$ [190°] (Claisen, *B* 19, 3316) On adding H_2SO_4 (2 pts) to a solution of benzoic aldehyde (3 pts) and (8) naphthol (6 pts) in alcohol (3 pts) there is formed $\text{C}_8\text{H}_7\text{O}_2$ [191°] uncr., a crystalline substance, not affected by boiling Ac_2O , and converted by fuming HNO_3 into $\text{C}_8\text{H}_7(\text{NO})\cdot\text{O}$ (Trzebinski, *B* 17, 499) —16 Benzotrichloride at 100° forms $\{\text{C}_8\text{H}_7\cdot\text{O}\cdot\text{C}_6\text{H}_5\}_2$ which melts above 350° and may be distilled. It crystallises in white needles, insol. water, alcohol, and ether, sol. nitrobenzene (Doebner, *A* 257, 59)

Combinations —With picric acid $\text{C}_{10}\text{H}_7\cdot\text{O}\cdot\text{C}_6\text{H}_3(\text{NO})_3\cdot\text{OH}$ [155°] Orange yellow needles (from alcohol), almost insol. cold water (Machetti, *G* 12, 504) —With aniline $\text{C}_{10}\text{H}_7\cdot\text{OC}_6\text{H}_5\cdot\text{N}$ [82 1/2°] Crystalline powder (from lignol) (Dyson, *C J* 43, 469) —With *p*-toluidine $\text{C}_{10}\text{H}_7\cdot\text{OC}_6\text{H}_4\cdot\text{N}$ [80 1/2°] (D) —Acetyl derivative $\text{C}_{10}\text{H}_7\cdot\text{OAc}$ [70°] (O) Miller, *B* 14, 1602) Small needles —Benzoyl derivative $\text{C}_{10}\text{H}_7\cdot\text{OBz}$ [107°] Nodular groups of needles (Maikopar, *Z* 1869, 216)

Methyl ether $\text{C}_{10}\text{H}_7\cdot\text{O}\cdot\text{Me}$ or $\text{C}_{10}\text{H}_7\cdot\text{OMe}$ [72°] (Staedel, *A* 217, 43), [70 1/2°] (Vincent, *Bl* [2] 40, 106) (274°) (Marchetti, *G* 9 545) Formed from (8) naphthol, KOH, MeOH, and MeI or MeCl. Formed also by boiling (8) naphthol with MeOH and ZnCl_2 . White plates (from alcohol), insol. cold water, v. sol. ether. Volatile with steam. With HNO_3 (S G 15) it gives at 0° a tri nitro derivative [213°]

Ethyl ether $\text{C}_{10}\text{H}_7\cdot\text{OEt}$ [33°] (275°). Formed from (8) naphthol, KOH, and EtI (Schaffer, Liebermann, *B* 15, 1428) Crystalline mass, decomposed in a sealed tube at 310° into ethylene and (8) naphthol (Bamberger, *B* 19, 1819)

Bromo ethyl ether $\text{C}_{10}\text{H}_7\cdot\text{OC}_2\text{H}_4\text{Br}$ [96°] Prepared by the action of ethylene bromide and KOH on (8) naphthol (Koelle, *B* 13, 1954) Colourless plates, sol. alcohol. Alcoholic ammonia at 100° converts it into amorphous $\text{C}_{10}\text{H}_7\cdot\text{O}\cdot\text{CH}_2\cdot\text{CH}_2\cdot\text{NH}_2$, while aniline yields crystalline $\text{C}_{10}\text{H}_7\cdot\text{O}\cdot\text{CH}_2\cdot\text{CH}_2\cdot\text{NHPH}$ [75°]

Ethylene ether $\text{C}_8\text{H}_7(\text{OC}_2\text{H}_5)_2$ [134°] **Ethylene ether** $\text{C}_8\text{H}_7(\text{OC}_2\text{H}_5)_2$ [217°] S. (benzene) 5 Plates (from benzene), insol. alcohol and ether

Ethylidene ether *v.* ALDEHYDE, vol. I, p. 105

Benzyl ether *v.* BENZYL-NAPHTHYL OXIDE

Naphthylether v Di NAPHTHOL OXIDE**Nitroso- derivative v Oxim of (β)-NAPHTHOQUINONE****References**—AMIDO, AMIDO DI IMIDO-, BROMO-, BROMO-NITRO-, CHLORO, IODO, IODO NITRO, and NITRO- NAPHTHOLS

(α) **Naphthol tetrahydride** $C_{10}H_8O$ $\frac{1}{2}e$
 $CH_3CH_2C \equiv CH$
 $CH_2CH_2C(OH)CH$ *Tetrahydro (α) naphthol* [69°] (265.5°) at 705 mm. Obtained by reducing (α) naphthol in amyl alcoholic solution by sodium. Occurs in the distillate obtained in the preparation of the nitrile of the tetrahydride of (α) naphthoic acid (Bamberger a Bordt, *B* 22, 628, 23, 215). Formed also from the tetrahydride of (α) naphthylamine by the diazo reaction (Bamberger a Althausse, *B* 21, 1892). Silvery white monoclinic tables, smelling like phenol, and turning red in air, sl sol water, v sol alcohol and ether. Bleaching powder gives only yellowish flakes in its boiling aqueous solution (whereas with (α)-naphthol it gives a deep violet colour). $FeCl_3$ gives no colouration.

Ethylether $C_{10}H_{11}OEt$ (259°) at 705 mm. Formed by means of $EtOH$, KOH , and EtI .

(β) **Naphthol tetrahydride** ('aromatic')
 $CH_3CH_2C \equiv CHCOH$ *ar Tetrahydro (β) naphthol* [58°] (275°). Formed by the action of nitrous acid on 'aromatic' (β) naphthylamine tetrahydride (Bamberger a Kitschelt, *B* 23, 884). It is also one of the products of the reduction of (β)-naphthol by sodium (*v infra*). Flat silvery needles, sl sol water, v sol alcohol and ether. Smells like creosote. Not etherified by heating with alcohol and H_2SO_4 at 100°. Not affected by heating with conc H_2SO_4 , hot dilute H_2SO_4 appears to form di naphthyl oxide octohydride $C_{10}H_{11}O$ (Bamberger a Lengfeld, *B* 23, 1129). Bleaching powder added to its aqueous solution forms a white flocculent pp. $FeCl_3$ gives a bluish green colour, and, on warming, a brownish yellow flocculent pp. Chloroform and caustic soda gave a greenish yellow colour. $NaNO_2$ added to its solution in conc H_2SO_4 gives a rose red colour. Forms a wine red compound with diazobenzene sulphonic acid. The sodium salt $C_{10}H_7ONa$ crystallises in silky scales.

(β) **Naphthol tetrahydride** ('alicyclo')
 $CHCHC \equiv CH, CHOH$
 $CHCHC \equiv CH, CH_2$ *ac Tetrahydro-(β) naphthol* (178° at 53 mm.), (264°) at 716 mm. Formed by adding sodium to a boiling solution of (β) naphthol in isoamyl alcohol (Bamberger a Lodter, *B* 23, 204), and separated from the accompanying 'aromatic' isomeride by aqueous $NaOH$, in which it is insoluble (Bamberger a Kitschelt, *B* 23, 885). V sol water, v sol alcohol and ether. On warming with KOH it yields naphthalene dihydride. HI yields an iodide, which readily splits up into HI and naphthalene dihydride. Does not react with diazo compounds.

Benzoyl derivative $C_{10}H_7OBz$ [63°] (255° at 40 mm.) Formed by heating with $HOBz$ at 170° (Bamberger a Lodter, *B* 23, 209). Tables, v e sol warm alcohol and benzene. Split up on distillation into benzoic acid and naphthalene dihydride.

Acetyl derivative $C_{10}H_7OAc$ (169° at 44 mm.) Formed by the action of $HOAc$ at

140°-150°. Decomposed at 268°-280° into $HOAc$ and naphthalene dihydride.

NAPHTHO-LACTONE v Anhydride of Oxy-NAPHTHOIC ACID

NAPHTHOL ALDEHYDE v Oxy NAPHTHOIC ALDEHYDE

NAPHTHOL-AZO- compounds v Azo compounds

DINAPHTHOL v Di Oxy DINAPHTHOL

(α) **NAPHTHOL BLUE** $C_{16}H_{16}NO$ $\frac{1}{2}e$

$NMe_2, C_6H_5N < \begin{smallmatrix} C_6H_5 \\ O \end{smallmatrix} >$ *Indophenol*. Formed by the action of nitroso dimethylaniline on (α) naphthol, and by the action of dibromo (α) naphthol on dimethyl *p* phenylene diamine in alkaline solution (Kochlin, *Bull de Mulh* 52, 532, Witt, *S C I* 1, 255). Obtained also by oxidising a mixture of di methyl *p* phenylene diamine and (α) naphthol or (α) naphthylamine with dilute $K_2Cr_2O_7$ and HCl (Pabst, *Bl* [2] 38, 161, Möhlau, *B* 16, 2851, 18, 2918). Bluish violet crystals, with bronze lustre (from alcohol), insol water. Dissolves in $HClAq$ with yellow colour, being split up into (α) naphthoquinone and dimethyl *p*-phenylene diamine.

NAPHTHOL CARBOXYLIC ACID v Oxy NAPHTHOIC ACID

(α) **NAPHTHOL-GLYCURONIC ACID**

$C_{16}H_{16}O_7$ [203°]. Occurs in urine after taking (α) naphthol (Lesnik a Nencki, *B* 19, 1537). Long needles, v sl sol chloroform. Split up by dilute $HClAq$ into (α) naphthol and glycuronic acid. Its aqueous solution is coloured a transient green by H_2SO_4 .

(β) **Naphthol-glycuronic acid** $C_{16}H_{16}O_7$ [150°] $[α] = -88°$. Occurs in the urine of dogs dosed with (β) naphthol, and can be separated through the insolubility of the lead salt (Lesnik a Nencki, *B* 19, 1534). Needles (containing 2aq), sl sol water, v sl sol chloroform. Conc H_2SO_4 gives an intense green colour. Split up by $HClAq$ into (β) naphthol and glycuronic acid— CaA' , 4aq.

(α) **NAPHTHOL-MALEIN** $C_{14}H_{10}O_4$ $\frac{1}{2}e$
 $CHCHC(C_6H_5OH)_2$ [118°-120°]. Formed,

CO—O together with a compound $C_{14}H_{10}O_4$, by heating (α) naphthol with maleic anhydride and $ZnCl_2$. Violet powder, consisting of microscopic four sided tablets. Dissolves in alcohol to a red solution, which on addition of NH_3 assumes a deep greenish red fluorescence. Sol ether, chloroform, and acetic acid, insol benzene and CS_2 (Burekhardt, *B* 18, 2867).

NAPHTHOL SULPHIDE v Di-Oxy-Di NAPHTHOL SULPHIDE

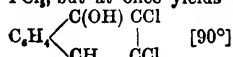
(1, 2) **NAPHTHOL SULPHONIC ACID**

[1 2] $C_{10}H_7(OH)SO_3H$ (α) *-Naphthol 'a' sulphonic acid* [101°]. Prepared by heating (α)-naphthol (1 pt) with conc H_2SO_4 (2 pts) on a water bath (Schaffer, *A* 152, 293, Claus, *B* 15, 312, Baum, *G P B* 4197 of June 30, 1883, *Monit Scient* 1883, 1122). Long, radiating, very deliquescent needles, v sol water and alcohol. Dissolves in HNO_3 , the solution slowly depositing di-nitro (α)-naphthol. $FeCl_3$ colours its solution deep blue, changing to green on warming, but becoming blue again on cooling. Gives no colour with bleaching powder. PCl_5 (2 mols) at 120° yields chloro-naphthol [57°] and di-chloro naphthalene [94°] (Claus a Oehler, *B* 15, 813).

Salts— CaA'_2 3aq Small laminae, v sol. water— PbA'_2 4aq needles, v sol water— $\text{PbC}_{10}\text{H}_7\text{SO}_3$ (dried at 100°) minute needles

Ethyl derivative $\text{C}_{10}\text{H}_7(\text{OEt})\text{SO}_3\text{H}$ Obtained from the acid by treatment with alcoholic KOH and EtI (Maikopar, *Z* 1870, 306)— $\text{K}'_2\text{A}'_2$ 3aq crystalline powder, sl sol cold water

(1, 3?) Naphthol sulphonic acid $\text{C}_{10}\text{H}_7\text{SO}_3$, i.e. $\text{C}_{10}\text{H}_6(\text{OH})\text{SO}_3\text{H}$ (a)-Naphthol 'B' sulphonic acid [90° uncor] Formed, together with other sulphonic acids by sulphonation of (a) naphthol in acetic acid solution by means of fuming sulphuric acid (80 p.c. SO_3) It is separated from concomitant products by the sparing solubility of its barium or lead salt Deliquescent, long, fine needles Very unstable, being converted into (a) naphthol and H_2SO_4 by boiling its dilute aqueous or alcoholic solutions The sodium salt cannot be converted into the chloride by PCl_5 , but at once yields di chloro (a) naphthol



Salts— NaA' easily soluble white silvery plates— KA' easily soluble needles— BaA'_2 2aq colourless plates, v sl sol cold water— PbA'_2 2aq small plates, v sl sol cold water (Baum, Claus a Knyrin, *B* 18, 2924)

Ethyl derivative $\text{C}_{10}\text{H}_7(\text{OEt})\text{SO}_3\text{H}$ [1 3?] Obtained by heating the ethyl ether of (a) naphthol (7 pts) with H_2SO_4 (5 pts) at 100° (Maikopar)— KA' 2aq large glittering plates, sl sol cold water— BaA'_2 [55°-60°] This acid is perhaps identical with the preceding

(1, 4) Naphthol sulphonic acid

[1 4] $\text{C}_{10}\text{H}_6(\text{OH})\text{SO}_3\text{H}$ [170°] Formed by diazotising the corresponding (a) naphthylamine sulphonic acid and heating the resulting diazo naphthalene sulphonic acid with dilute H_2SO_4 (Neville a Winther, *C J* 37, 632, *Monat Scient* 1884, 39, Erdmann, *A* 247, 341) Plates, v e sol water Converted by dilute HNO_3 into di-nitro (a) naphthol Chromic acid oxidises it to (a) naphthoquinone FeCl_3 colours its solution greenish blue, but on warming the colour becomes red Its sodium salt is v sol 90 p.c. alcohol, from which it crystallises in needles

(1, 1') Naphthol sulphonic acid

[1 1'] $\text{C}_{10}\text{H}_6(\text{OH})\text{SO}_3\text{H}$ [107°] Obtained from the corresponding (a) naphthylamine *peri* sulphonic acid by diazotisation, the product being boiled with water The resulting anhydride (*infra*) is heated with alcoholic ammonia for half an hour at 130° , and the ammonium salt thus obtained converted into lead salt, and thence into the free acid (Erdmann, *A* 247, 346) Deliquescent crystalline mass (containing aq), v e sol water Gives with FeCl_3 a green colour changing to red

Salts— $\text{NH}_4\text{A}'$ v sol water, but not deliquescent— KA' plates— $\text{Na}_2\text{C}_{10}\text{H}_6\text{SO}_3$ 1½aq aggregates of needles— $\text{PbC}_{10}\text{H}_6\text{SO}_3$ 3aq white crystalline powder

Anhydride $\text{C}_{10}\text{H}_6\text{SO}_3 < \text{SO}_2 >$ Naphthosultone [154°] Formed as above Prisms (from benzene), v sl sol CS_2 , sl sol alcohol, v sol chloroform, insol water Boils between 360° and 448° Not attacked by aqueous NH_3 , Na_2CO_3 , or cold NaOH aq Boiling NaOH aq converts it into the corresponding acid

(1, 4') Naphthol sulphonic acid

[1 4'] $\text{C}_{10}\text{H}_6(\text{OH})\text{SO}_3\text{H}$ [110°-120°] Obtained from the corresponding naphthylamine sulphonic acid by diazotising and boiling the product with H_2SO_4 (1 pt) diluted with water (4 pts) (Erdmann, *A* 247, 343) Hygroscopic crystalline substance

(1, 2') Naphthol sulphonic acid Formed by partial hydrolysis of heteronuclear (a) naphthol disulphonic acid (Liebmann a Studer, *E P* 7812 [1887]) Yields azo dyes

(a) Naphthol disulphonic acid (Heteronuclear) $\text{C}_{10}\text{H}_6(\text{OH})(\text{SO}_3\text{H})_2$ Obtained, together with the trisulphonic acid, and probably also the (1, 2, 4) disulphonic acid, by heating (a)-naphthol (1 pt) with H_2SO_4 (containing an additional 20 p.c. of SO_3) (5 pts) for two hours on a water bath (Claus a Mieleke, *B* 19, 1182) Best prepared by heating (a) naphthol (1 pt) with conc H_2SO_4 (3 to 4 pts) at 180° Yields a nitroso compound $\text{C}_{10}\text{H}_6(\text{OH})(\text{NO})(\text{SO}_3\text{H})_2$ (Seltzer, *G P* 20,716, Jan 30, 1882) Combines with diazo compounds On nitration it forms di nitro naphthol sulphonic acid

Chloride $\text{C}_{10}\text{H}_6(\text{OH})(\text{SO}_2\text{Cl})_2$ Liquid, v sol ether Converted by PCl_5 into tri chloro naphthalene [82°]

Ethyl derivative $\text{C}_{10}\text{H}_7(\text{OEt})(\text{SO}_3\text{H})_2$ Formed by sulphonating the ethyl ether of (a) naphthol with fuming H_2SO_4 (10 p.c. additional SO_3) (*G P* D 1501, June 7, 1883) Conc HNO_3 converts it into di nitro naphthol sulphonic acid

(a) Naphthol disulphonic acid (Homonuclear) $\text{C}_{10}\text{H}_6(\text{OH})(\text{SO}_3\text{H})_2$ [1 2 4] Formed by heating (a) naphthol with H_2SO_4 (4 pts) at 70° Differs from the preceding acid in not yielding a nitroso derivative or forming azo dyes (Schultze, *Disertation*, Freiburg, 1883, Bender, *B* 22, 993) It yields di nitro naphthol on nitration

(1, 1', 4) (a) Naphthol disulphonic acid [1 1' 4] $\text{C}_{10}\text{H}_6(\text{OH})(\text{SO}_3\text{H})_2$ (a)-Naphthol (8) sulphonic acid Formed from (a) naphthylamine *peri* sulphonic acid by sulphonation and diazotisation Formed also by sulphonating naphtho sultone (Berntsen, *B* 23, 3088) Readily forms an anhydride— NaA' plates, v sol water FeCl_3 colours its solution deep blue— BaA'

Anhydride $\text{C}_{10}\text{H}_6(\text{SO}_3\text{H})_2 < \text{SO}_2 >$ Salts NaA' 3aq, m sol water Gives no colour with FeCl_3 — BaA'_2 v sol water

Mono amide $\text{C}_{10}\text{H}_6(\text{SO}_3\text{H})(\text{OH})(\text{SO}_2\text{NH}_2)$ From the anhydride and NH_3 — NaA' 2aq Plates

(1, 1', 3) (a) Naphthol disulphonic acid [1 1' 3] $\text{C}_{10}\text{H}_6(\text{OH})(\text{SO}_3\text{H})_2$ (a) Naphthol 'ε' disulphonic acid Formed from (a) naphthylamine 'ε' disulphonic acid by the diazo reaction (Berntsen, *B* 22, 3330, cf Ewer a Pick, *Monat Scient* 1889, 604)— NaA' 6aq prisms, v e sol water Its solution is coloured deep-blue by FeCl_3

Anhydride $\text{C}_{10}\text{H}_6(\text{SO}_3\text{H})_2 < \text{SO}_2 >$ Naphthosultone sulphonic acid Formed by boiling diazonaphthalene 'ε'-disulphonic acid with dilute H_2SO_4 Needles (containing aq)

Mono-amide [1' 3] $\text{C}_{10}\text{H}_6(\text{OH})(\text{SO}_2\text{NH}_2)(\text{SO}_3\text{H})$ Naphthol sulph amide sulphonic acid Formed by the action of NH_3 on the anhydride Needles or prisms, m. sol water— NaA' 2aq needles, v. sol hot

water— NH_4NaA aq crystals, v sol water— BaA'_2 5aq crystals, m sol water

(a) **Naphthol trisulphonic acid** $\text{C}_{10}\text{H}_6\text{S}_3\text{O}_{10} \cdot 2\text{e}$ $\text{C}_{10}\text{H}_4(\text{OH})(\text{SO}_3\text{H})_3$ [124'4] Formed by heating (a) naphthol with fuming H_2SO_4 (containing 70 p.c. additional SO_3) at 50° (Seltzer, *G P* 10,786, Dec 1879, Caro, *B* 14, 2028, Claus a Mielcke, *B* 19, 1182) Slender needles Dilute HNO_3 at 50° yields di nitro naphthol sulphonic acid It does not yield azo dyes or a nitroso-derivative— $\text{K}_2\text{C}_{10}\text{H}_3\text{S}_3\text{O}_{10}$, crystals, v sol water (Lauterbach, *B* 14, 2028)

Chloride $\text{C}_{10}\text{H}_4(\text{OH})(\text{SO}_2\text{Cl})_2$ Plates (from chloroform), sl sol either, decomposed by fusion Converted by PCl_5 into tetra chloro naphthalene $\text{C}_{10}\text{H}_2\text{Cl}_4$ [140°] and C_{10}Cl_8

(2, 1')-(β) **Naphthol sulphonic acid** [21'] $\text{C}_{10}\text{H}_7(\text{OH})(\text{SO}_3\text{H})$ (Pfützing a Duisberg, *B* 22, 396, Armstrong a Wynne, *C J Proc* 1889, 60) (β)-Naphthol 'a'-sulphonic acid **Bayer's acid** **Rumpff's acid** Formed, together with the isomeric acid of Schaffer, by stirring (β) naphthol (1 pt) with conc H_2SO_4 (2 pts) at about 50° (Bayer a Co, *G P* 18,027, March 18, 1881) When (β) naphthyl sulphate $\text{C}_{10}\text{H}_7\text{O}_4\text{S}_2$ (obtained by the action of H_2SO_4 on (β) naphthol at a low temperature) is mixed with H_2SO_4 at 20° it slowly changes to (β) naphthol 'a' sulphonic acid The two isomeric acids of Schaffer and of Bayer may be separated by means of their lead salts (that of the former acid crystallising well) or by treating their sodium salts with 90 p.c. spirit, which dissolves the salt of Bayer's acid but not that of Schaffer's acid Formed also from (β) naphthylamine 'a' sulphonic acid (Badische) by diazotising and heating with dilute H_2SO_4 (Forsling, *B* 20, 2102)

Reactions—1 Converted by PCl_5 into chloro-(β) naphthol [101°] and di chloro naphthalene [61°]—2 Potash fusion yields di oxy naphthalene [175°]—3 With diazotised xylidine it yields in concentrated, but not in dilute solutions, crocein scarlet

Salts—Forms two series of salts—neutral and basic— NaA' glistening six sided plates, v sol water, sl sol alcohol— $\text{Na}_2\text{C}_{10}\text{H}_6\text{S}_2\text{O}_7$ needles, v sol alcohol, v sol water— ZnA'_2 2aq needles— PbA'_2 2aq colourless glistening rhombohedra— $\text{C}_{10}\text{H}_6\text{S}_2\text{O}_7\text{Pb}_2$ small yellow crystals— $\text{C}_{10}\text{H}_6\text{S}_2\text{O}_7(\text{PbOH})_2$ red crystals (Claus a Volz, *B* 18, 3154)

(2, 3') **Naphthol sulphonic acid**
[23'] $\text{C}_{10}\text{H}_7(\text{OH})(\text{SO}_3\text{H})$ (β) Naphthol ' β '-sulphonic acid **Schäffer's acid** [125°]

Formation—1 By heating (β) naphthol (1 pt) with conc H_2SO_4 (2 pts) on a water bath (Schaffer, *A* 152, 296, Armstrong a Graham, *C J* 39, 135)—2 By fusing naphthalene (β) disulphonic acid with potash (Ebert a Merz, *B* 9, 610, 10, 592)—3 From (β) naphthylamine sulphonic acid (Brönner's) by the diazo reaction

Properties—Small, non deliquescent laminae, v sol water and alcohol FeCl_3 turns its solution slightly green and, on warming, brown flakes are deposited Bleaching powder gives a yellow colour

Reactions—1 Potash-fusion yields di oxy-naphthalene [213°]—2 Nitrous acid forms a nitroso-derivative—3 Bromine water gives mono- and di bromo derivatives The K salt

dissolved in HOAc gives with Br yellow plates of $\text{C}_{10}\text{H}_6\text{BrSO}_4\text{K}$, which form a blood red aqueous solution—4 Conc HClAq at 200° – 210° splits it up into (β) naphthol and H_2SO_4 —5 The potassium salt (1 mol) heated with PCl_5 (2 mols) at 100° forms gelatinous anhydro naphthol sulphonic acid $\text{C}_{10}\text{H}_4(\text{OH})\text{SO}_2\text{O}\text{C}_{10}\text{H}_6\text{SO}_3\text{H}$, sol water, alcohol, and ether The K salt of this acid crystallises in colourless plates, and is decomposed by boiling alkalis with reproduction of (β) naphthol sulphonic acid (Claus a Zimmernann, *B* 14, 1481) The K salt (1 mol) heated with PCl_5 (2 mols) at 150° forms $(\text{SO}_3\text{H}\text{C}_{10}\text{H}_6\text{O}\text{SO}_2\text{C}_{10}\text{H}_6)_2\text{O}$, an amorphous mass yielding a gelatinous potassium salt KA'' (C a Z) The (β) naphthol sulphonic acid heated with PCl_5 also yields chloro (β) naphthol, and finally γ di chloro naphthalene [135°], both of which bodies yield chloro phthalic acid [148°] on oxidation (Claus a Dehne, *B* 15, 319)

Salts— KA' 2aq S 2 at 15° — $\text{NH}_4\text{A}'$ S 3 at 24° Flat prisms or plates (Meldole, *C J* 39, 41)— CaA'_2 5aq silky laminae, v sol water and alcohol, decomposing at 100° — BaA' 6aq narrow prisms (from boiling saturated solution)— PbA' 6aq small silvery laminae, v sol water

Ethyl derivative $\text{C}_{10}\text{H}_7(\text{OEt})(\text{SO}_3\text{H})$ From the (? impure) acid, KOH, and EtI (Maikopar, *Z* 1870, 366)— KA' needles, sl sol cold water

(β) **Naphthol (γ) sulphonic acid**
[24'] $\text{C}_{10}\text{H}_6(\text{OH})(\text{SO}_3\text{H})$ Formed from the corresponding (β) naphthylamine sulphonic acid by the diazo reaction (Reverdin a Nolting, *Const de la Naphthalme*) Yields di oxy naphthalene [135°] and di chloro naphthalene [48°] (Claus, *J pr* [2] 39, 315)

(2, 2') (β) **Naphthol sulphonic acid**
[22'] $\text{C}_{10}\text{H}_6(\text{OH})_2\text{SO}_3\text{H}$ (β) Naphthol ' β ' sulphonic acid **Naphthol sulphonic acid F** [89°] Formed from (β) naphthylamine ' β ' sulphonic acid by the diazo reaction (Bayer a Duisberg, *B* 20, 1431) Formed also, together with Schaffer's acid, by heating (β) naphthol (100 g) with H_2SO_4 (70 g) for 2 or 3 hours at 105° (Green, *B* 22, 723, cf Armstrong, *B* 15, 22) It is also a product of the fusion of naphthalene 'a' disulphonic acid with NaOH at 250° (Weinberg, *B* 20, 2907) Needles (containing aq), v sol water and alcohol, insol ether (W) PCl_5 (3 pts) at 165° gives di chloro naphthalene [114°], the chief product being a chloro naphthyl phosphate [215°] On fusion with potash it yields $\text{C}_{10}\text{H}_6(\text{OH})_2$ [186°] Nitrous yields a nitroso-derivative, forming the salt $\text{C}_{10}\text{H}_6(\text{NO})(\text{OH})\text{SO}_3\text{Na}$ 2aq crystallising in golden needles

Salts— NaA' 23aq S 8 at 15° Plates— Ka' aq crystals, v sol water— MgA'_2 53aq plates— BaA' , prisms, sl sol water

(β) **Naphthol sulphonic acid** **Ethyl derivative** $\text{C}_{10}\text{H}_6(\text{OEt})_2\text{SO}_3\text{H}$ Formed by warming the ethyl ether of (β) naphthol with H_2SO_4 (Maikopar, *Z* 1870, 366)— KA' aq needles, sl sol cold water— BaA'_2 needles, v sl sol water

(β) **Naphthol sulphonic acid** **Ethyl derivative** $\text{C}_{10}\text{H}_6(\text{OEt})_2\text{SO}_3\text{H}$ Two acids of this formula are obtained by treating the ethyl derivative of (β) naphthol with ClSO_3H (Amphlett a Armstrong, *C J Proc* 3, 144) The Ba salt of one acid is insol water, that of the other is sol water One of the acids is doubtless identical with the preceding acid

(2, 1' 3') Naphthol disulphonic acid [2 1' 3'] $C_{10}H_7(OH)(SO_3H)_2$ (β) Naphthol 'β'-disulphonic acid (β) Naphthol-G-disulphonic acid Formed, together with the 'R' isomeride, by heating (β)-naphthol (1 pt) with conc or fuming H_2SO_4 (2 or 3 pts) at 100°–110°, and separated from its isomeride through the greater solubility of its Ba salt (Gress, B 13, 1956)

Properties —Glistening needles, v sol water and alcohol With diazo compounds it gives scarlets of a much yellower shade than the (α)-acid PCl_5 (5 mols) at 210° forms di chloro-(β) naphthol [125°] and tri chloro naphthalene [90°], this $C_{10}H_6Cl_3$, when oxidised with CrO_3 , yields a syrupy di chloro phthalic acid and a tri chloro naphthoquinone, whence aniline forms $C_{10}H_6Cl_3(NPhH)O$, melting at 228° (Claus a Schmidt, B 19, 3173)

Salts — Na_2A'' tables or prisms, sol water and dilute alcohol — BaA'' 8aq small prisms, v. sol water

(2, 3, 3') Naphthol disulphonic acid [2 3 3'] $C_{10}H_7(OH)(SO_3H)_2$ (β) Naphthol 'α' disulphonic acid (β) Naphthol R disulphonic acid Formed as described above (Gress) White glistening needles, v sol water and alcohol, insol ether Exhibits a bluish green fluorescence in ammoniacal solution When heated in a closed vessel with ammonia it yields a naphthylamine disulphonic acid which, on removal of NH_3 , gives rise to naphthalene 'α' disulphonic acid of Ebert and Merz (Pfitzinger a Duisberg, B 22, 398)

Salts — Na_2A'' very small soluble needles — BaA'' 6aq needles, sl sol cold water, v sol hot water, insol alcohol

(β) Naphthol-β'-disulphonic acid [2 2' 3'] $C_{10}H_7(OH)(SO_3H)_2$ Formed by sulphonating (2, 2') naphthol sulphonic acid (Weinberg, B 20, 2911) Solutions of its salts exhibit green fluorescence — Na_2A'' yellowish powder, v e sol water, sl sol 80 p c alcohol — BaA'' 2½aq prisms S 56 in boiling water

(β) Naphthol-γ'-disulphonic acid $C_{10}H_7(OH)(SO_3H)_2$ Formed by the action of $ClSO_3H$ on (β) naphthol (Armstrong, B 15, 204) The Ba salt crystallises in large prisms

(β) Naphthol trisulphonic acid $C_{10}H_7(OH)(SO_3H)_3$ Formed by heating (β)-naphthol (1 pt) with H_2SO_4 (2 pts) at 70°–80°, adding H_2SO_4 (2 pts) and heating at 120°, finally adding fuming H_2SO_4 (40 p c additional SO_3) (2 pts) and heating at 150° (Levinstein, B 16, 462) Gives no colour with diazotised xylidine

References —AMIDO-, and NITRO-NAPHTHOL SULPHONIC ACIDS

(β) NAPHTHOL VIOLET $C_{10}H_7N_2O$ Mel-dola's blue Formed from nitroso-dimethyl aniline and (β) naphthol Dark flocculent powder, forming a red solution in benzene (Mel-dola, C J 89, 38)

Hydrochloride $B'HCl \cdot a$.

$ClMe_3N \cdot C_6H_5 \cdot \begin{smallmatrix} \diagup O \\ \diagdown \end{smallmatrix} C_{10}H_7$ Bronzed needles, resembling $KMnO_4$ Sol water and alcohol Its aqueous solution is violet, but turned blue by H_2SO_4 Reduces to a leuco base by taking up H_2 . Converted by heat into a blue colouring matter $NMe_2 \cdot C_6H_5 \cdot N \begin{smallmatrix} \diagup O \\ \diagdown \end{smallmatrix} C_{10}H_7 \cdot N \begin{smallmatrix} \diagup O \\ \diagdown \end{smallmatrix} C_6H_5$ (Witt, B.

23, 2247)

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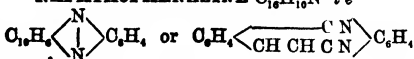
NAPHTHOL-YELLOW s v DINITRO (α)-NAPHTHOL SULPHONIC ACID

NAPHTHONITRILE v Nitrile of NAPHTHOIC ACID

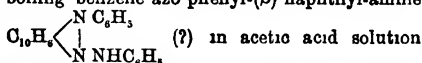
NAPHTHOPHTHALIC ACID A name for NAPHTHALENE DICARBOXYLIC ACID

NAPHTHO-OXY-QUINALDINE v OXY METYL-NAPHTHOQUINOLINE

NAPHTHOPHENAZINE $C_{10}H_7N$ s e



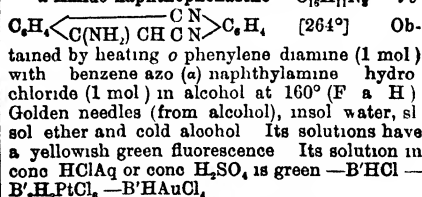
[142°] (above 360°) Prepared by boiling sulpho benzene azo phenyl-(β) naphthylamine with dilute H_2SO_4 (Witt, B 20, 574) Formed by boiling benzene azo phenyl-(β) naphthyl-amine



with conc HCl , aniline being eliminated (Zinke a. Lawson, B 20, 1169) Formed also by treating an alcoholic solution of phenyl (β) naphthyl nitrosamine with HCl (Fischer a Hepp, B 20, 2473), and by distilling with zinc dust oxy naphtho phenazine, which is itself obtained by heating with conc $HClAq$ at 190° the amido naphtho phenazine which results from the action of phenylene o diamine on benzene azo (α) naphthylamine hydrochloride (Fischer a Hepp, B 23, 846) Also got by the action of (β) naphtho quinone (1 mol) on o phenylene diamine (1 mol) in 50 p c acetic acid solution at 0°, and by oxidising a mixture of o phenylene diamine and (β) naphthol with alkaline K_2FeC_6 (Witt)

Yellowish white prisms, subliming at about 200° in long flat needles or prisms, v sol hot benzene, v sl sol alcohol and ether Conc H_2SO_4 dissolves it, forming a brownish red solution Its salts are dimorphous One sulphate crystallises in scarlet needles, the other in garnet red prisms One nitrate forms orange needles, the other brick red needles One hydrochloride forms long reddish yellow needles, and changes at 150° into the second form $SnCl_4$ reduces naphthophenazine to a hydride, crystallising in violet needles

α Amido-naphthophenazine $C_{10}H_7N_2$ s e



Acetyl derivative $C_{10}H_7AcN$, Yellow crystalline powder

Amido naphthophenazine $C_{10}H_7(NH_2)N \cdot C_6H_5$ [191°] Obtained by reducing the nitro compound with alcoholic ammonium sulphide (Zaerling, B 23, 176) Brownish-red crystals, sl sol alcohol and ether, v e sol boiling aniline — $B'HCl$

Phenyl-o-chloride of Amido-naphthophenazine $C_{10}H_7N \cdot Cl$ s e.

$C_{10}H_7 \cdot \begin{smallmatrix} \diagup N \\ \diagdown \end{smallmatrix} NPhCl \cdot C_6H_5 \cdot NH_2$ Formed by the action of quinone dichlorimide on phenyl-(β)-naphthylamine (Nietsch a. Otto, B 21, 1600).

H H

Yields a base which is a bluish-violet pp and a nitrate $C_{12}H_7N_2NO_3$, crystallising in slender needles. Yields naphthophenazine on elimination of NH_3 by the diazo reaction

Reference — EURHODINES

Nitro-naphthophenazine $C_{12}H_7(NO_2)N_2$, C_6H_5 , [222°] Formed by heating nitro (β) naphthoquinone with *o* phenylene-diamine, HOAc, and NaOAc on the water bath (Z) Greenish yellow prisms, sl sol alcohol and ether. Forms a crimson solution in conc H_2SO_4 , — $B'2H_2SO_4$, brown plates

NAPHTHOPHENAZINE CARBOXYLIC ACID $C_{12}H_7N_2(CO_2H)$ [above 300°] Obtained by heating its nitrile with KOHAq under pressure at 225° (Brunner a Witt, B 20, 2663) Needles in sol water and most solvents. Conc H_2SO_4 forms a deep red solution, becoming yellow on dilution.

Nitrile $C_{12}H_7CyN_2$ [237°] Formed from sodium naphthophenazine sulphonate and KCy. Forms a cherry red solution in conc H_2SO_4 , becoming yellow on dilution

Naphthophenazine sulphonic acid $C_{12}H_7(SO_3H)N_2$ [above 290°] Formed by sulphonating naphthophenazine with fuming H_2SO_4 (35 p c) (Brunner a Witt, B 20, 2660) Orange-red needles, sol boiling water and alcohol. KOH yields yellow flakes of a eurhodol. H_2SO_4 forms an orange red solution — NaA' 2aq

NAPHTHOPIASELENOL ν SELENIUM ORGANIC COMPOUNDS

NAPHTHOPIAZTHIOLE $C_{12}H_7\langle\begin{smallmatrix} N \\ N \end{smallmatrix}\rangle S$ [81°]. Formed by heating (αβ) naphthylene diamine (2 g) dissolved in alcohol, with a conc solution of sodium bisulphite (15 c c) at 190° for 7 hours (Hinsberg, B 23, 1393) Long needles (from MeOH), m sol water. Pleasant odour. Somewhat volatile with steam. Weak base. Gives a yellow colour with conc H_2SO_4 . Reduced by tin and HCl to H_2S and naphthylene diamine

NAPHTHOPICRIC ACID, so called, ν TRINITRO- α NAPHTHOL

NAPHTHO-QUINALDINE ν (Py 3) METHYL-NAPHTHO QUINOLINE

(a) **NAPHTHOQUINOLINE** $C_{13}H_9N$ ν $C_6H_5\langle\begin{smallmatrix} CH \\ CH \end{smallmatrix}\rangle C \begin{smallmatrix} CH \\ CH \end{smallmatrix}$ Formed by heating a mixture of (a) nitro naphthalene, (a) naphthylamine, glycerin, and H_2SO_4 (Skraup, M 2, 162, 4, 460) Formed also by distilling its dicarboxylic acid (Doebner a Peters, B 23, 1235) White prisms (from ether), insol water, sol alcohol, ether, benzene, and dilute acids. Smells like naphthylamine. On oxidation with chromic acid it yields the quinone $C_{12}H_6NO_2$ or

$C_6H_5\langle\begin{smallmatrix} CO \\ CO \end{smallmatrix}\rangle C \begin{smallmatrix} CH \\ CH \end{smallmatrix}$ which forms orange crystals, [205°], sol alcohol, benzene, ether, and HClAq, insol water. $KMnO_4$ oxidises it to phenyl pyridine dicarboxylic acid

Salts — Hydrochloride pale yellow needles — $B'H_2PtCl_4$ 2aq. bright-yellow prisms, ν sl sol water — $B'H_2SO_4$, yellowish prisms, ν sol. water — $B'H_2CrO_4$ 6aq needles, sl sol cold water — $B'CO_2H(NO_2)_2$, minute needles

Methylo-sodide B MeI 2aq. Needles.

Octohydrate $C_{13}H_9N_2 \cdot 8H_2O$

$CH_2CH_2C \begin{smallmatrix} CH \\ CH \end{smallmatrix} C \begin{smallmatrix} CH \\ CH \end{smallmatrix} CH_2$ Crystals (Bamberger, B 22, 354)

(β) **Naphthoquinoline** $C_{12}H_7N_2$

$C_6H_5\langle\begin{smallmatrix} CH \\ CH \end{smallmatrix}\rangle C \begin{smallmatrix} CH \\ CH \end{smallmatrix}$ [94°] (above 360°).

Formed by heating (β) naphthylamine (28 pts) with nitro-benzene (13 pts), glycerin (50 pts), and conc H_2SO_4 (40 pts) at 150°–160° (Skraup a Cobenzl, M 4, 436) The crude product is neutralised with alkali, extracted with ether, and distilled. In this reaction (1,2) bromo (β) naphthylamine [63°] may be substituted for (β) naphthylamine, and nitro phenol may be used instead of nitro benzene (Leimann a Schmidt, B 20, 3154) Formed also by heating (1,2) nitro (β)-naphthylamine with glycerin and H_2SO_4 (L a S) Obtained likewise by distilling its carboxylic acids

Properties — Colourless crystals, sl sol water, sol ether, alcohol, benzene, and acids. Its alcoholic solution gives a brown colouration with $FeCl_3$, and a green colour with cupric acetate

Reactions — 1. $KMnO_4$ oxidises it to (β)-phenyl pyridine dicarboxylic acid

Salts — $B'HCl$ 2aq long needles, ν sol water, sl sol alcohol — $P'H_2PtCl_4$ aq orange crystalline pp — $B'H_2CrO_4$, short golden prisms — $B'ClHCl$ [c 148°] Yellow needles, obtained by adding a solution of ICl in HClAq to a solution of the base (Dittmar, B 18, 1616) — Picrate [252°]

Methylo-sodide B MeI 2aq [200°–205°] Light yellow needles, exhibiting a blue fluorescence in aqueous solution

Octohydrate $C_{13}H_9N_2 \cdot 8H_2O$

$CH_2CH_2C \begin{smallmatrix} CH \\ CH \end{smallmatrix} C \begin{smallmatrix} CH \\ CH \end{smallmatrix} NH \begin{smallmatrix} CH_2 \\ CH_2 \end{smallmatrix}$ [60°] Obtained by boiling (β) naphthoquinoline with isoamyl alcohol, and sodium (Bamberger, B 22, 354) At the same time there is formed an isomeric $C_6H_5\langle\begin{smallmatrix} CH_2 \\ CH_2 \end{smallmatrix}\rangle C \begin{smallmatrix} CH \\ CH \end{smallmatrix} NH \begin{smallmatrix} CH_2 \\ CH_2 \end{smallmatrix}$ [91°]

Reference — Oxy, Oxy methyl, and PHENYL-NAPHTHOQUINOLINE

(β) **NAPHTHOQUINOLINE CARBOXYLIC ACID** $C_{12}H_7NO_2$ ν $C_6H_5\langle\begin{smallmatrix} CH \\ CH \end{smallmatrix}\rangle C \begin{smallmatrix} CH \\ CH \end{smallmatrix} CO_2H$

[187°] Obtained by oxidising methyl (β) naphthoquinoline with $KMnO_4$ and H_2SO_4 (Seitz, B 22, 261) White crystalline powder, insol water, m sol boiling alcohol — NaA' 2aq Crystals, sl sol cold water — BaA' 4aq flocculent pp, becoming crystalline on long boiling, insol water — $CuA' \cdot 1\frac{1}{2}$ aq crystalline, insol water — $B'HCl$ yellow needles, ν sl sol boiling dilute HClAq — $B'H_2PtCl_4$ 2aq yellow needles, ν sl sol hot dilute HClAq

(a) **Naphthoquinoline** (Py 1, 3) dicarboxylic acid $C_6H_5\langle\begin{smallmatrix} CH \\ CH \end{smallmatrix}\rangle C \begin{smallmatrix} CO_2H \\ CO_2H \end{smallmatrix}$ [278°]

Formed by oxidising (a) styryl-(a) naphthoquinoline carboxylic acid with cold aqueous $KMnO_4$ (Doebner a Peters, B 23, 1284) Groups of greenish yellow needles, insol cold water, sl sol hot water, ether, cold alcohol, and dilute HClAq, ν sol conc HClAq

Salts — The salts of the alkalis and alkaline earths are ν sol water. The Pb and Cd salts

are white pps— $\text{CuA}''2\text{aq}$ dirty green pp, sl sol water— AgA'' white flocculent pp

(β) Naphthoquinoline (*Py* 1, 3) dicarboxylic acid $\text{C}_{10}\text{H}_6\text{N}_2\text{O}_4$ $\begin{array}{c} \text{CH} \text{ CH} \text{ C} \text{ N} \\ \diagup \quad \diagdown \quad \diagup \quad \diagdown \\ \text{C} \text{ CO}_2\text{H} \end{array}$ [288°]

Formed by oxidation of (α) styryl (β) naphthoquinoline carboxylic acid by KMnO_4 (Doeberner, *Peters*, *B* 23, 1240) Very slender light yellow needles, sl sol ether, cold alcohol, and benzene, v sol HOAc and alkalis— $\text{BaA}'_2\text{aq}$ white flocculent pp— AgA'' powder, sl sol water

NAPHTHOQUINOLINE SULPHONIC ACID

$\text{C}_{10}\text{H}_6\text{NSO}_3 \cdot e$ $\text{C}_{10}\text{H}_6(\text{SO}_3\text{H})_2$ $\begin{array}{c} \text{C} \text{ CH} \text{ CH} \text{ C} \text{ N} \text{ CH} \\ \diagup \quad \diagdown \quad \diagup \quad \diagdown \\ \text{C} \text{ CH} \text{ CH} \end{array}$

Obtained from (β) naphthylamine sulphonic acid of Forsling (*B* 20, 2099) by heating with glycerin, H_2SO_4 , and nitrobenzene (Immerheiser, *B* 22, 404, cf Gentil, *B* 18, 201). Small needles, nearly insol water Very dilute solutions of the acid and its salts fluoresce blue— $\text{BaA}'_2\text{aq}$ Splendid prisms

(α)-NAPHTHOQUINONE $\text{C}_{10}\text{H}_6\text{O}_2 \cdot e$

$\text{C}_{10}\text{H}_6\text{O}_2$ $\begin{array}{c} \text{CO} \text{ CH} \\ \diagup \quad \diagdown \\ \text{CO} \text{ CH} \end{array}$ (α) Naphthoquinone Mol w 158 [125°]

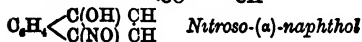
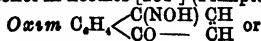
Formation—1 By oxidation of naphthalene by CrO_3 in HOAc (Groves, *C J* 26, 209, cf Hermann, *A* 151, 63)—2 By oxidising with chromic acid the following bodies naphthylene *p* diamine, and its acetyl derivative (Liebermann a Dittler, *B* 6, 945), (α) naphthylamine and its sulphonic acid (Reverdin a Nolting, *B* 12, 2305), (α) amido (α) naphthol (Liebermann, *A* 183, 242), and the acetyl derivative of (α) naphthol (Miller, *B* 14, 1600)

Preparation—1 Naphthalene (10 g) is dissolved in HOAc (60 g), to this is slowly added a solution of CrO_3 (30 g) in water (20 cc) The mixture is heated at 100° Water (30 g) is added, and the liquid, when cooled to 20°, filtered from naphthalene From the filtrate water precipitates naphthoquinone Recrystallised from light petroleum (Japp a Miller, *C J* 39, 220, cf Groves, *C J* 26, 209, Plimpton, *C J* 37, 634, Miller, *Bl* [2] 43, 125)—2 (α) Naphthol orange (Tropaeoline 000 No 1) is reduced to sulphanic acid and (14) amido naphthol, the latter is oxidised and the quinone purified by distillation with steam, the yield is 40 p c—3 By oxidation of (α)-naphthylamine (1 pt) with H_2SO_4 (6 pts), water (25 pts), and $\text{K}_2\text{Cr}_2\text{O}_7$ (2½ pts), in the cold (Monnet, Reverdin, a Nolting, *B* 12, 2306)

Properties—Yellow triclinic needles, which begin to sublime below 100° Volatile with steam V sl sol water and ligroin, v sol benzene, CS_2 , chloroform, ether, HOAc, conc H_2SO_4 , and boiling alcohol Smells like quinine Alkalis form a reddish brown solution It is not attacked by SO_2 in the cold Reacts in alcoholic solution with primary amines forming compounds represented by $\text{C}_{10}\text{H}_6\text{O}_2\text{NR}$, which may be considered to be alkyl amido-(α) naphthoquinones or more probably as oxy-(β) naphthoquinone alkylimides Secondary amines act forming compounds represented by $\text{C}_{10}\text{H}_6\text{O}_2\text{NR}'\text{R}'$, but diphenylamine acts only in presence of HCl Tertiary amines do not react in this way Hydro naphthoquinone may be used instead of naphthoquinone, being oxidised by the air Salts of amines may be used instead of amines The compounds crystallise in needles (Plimpton, *C J* 37, 638)

Reactions—1 Naphthoquinone dissolves in aqueous KOH and Na_2CO_3 , and on adding acids a red body is ppd This is also got by oxidising naphthalene with CrO_3 in acetic acid and neutralising the hot liquid with Na_2CO_3 (A. Guyard, *Bl* [2] 31, 64, Plimpton, *C J* 37, 641) It is soluble in benzene and ppd from it by light petroleum as a dark red powder This substance was named carminaphthe by Laurent (*Rev Scient* 14, 560), who ascribed to it the formula $\text{C}_{10}\text{H}_6\text{O}_2$ —2 Boiling nitric acid oxidises it to phthalic acid—3 Boiling HIAq and P reduct to hydronaphthoquinone [176°] Tin and HCl act in like manner—4 When boiled with an aqueous solution of an equivalent of hydronaphthoquinone there is deposited on cooling dark purple crystals of naphthoquinhydrone $\text{C}_{20}\text{H}_{12}\text{O}_4$ It is readily converted by oxidising agents into naphthoquinone and by reducing agents into hydronaphthoquinone—5 Bromine in presence of iodine forms di bromo naphthoquinone [218°] (Miller, *Bl* [2] 43, 125)—6. Heated with benzoic acid at 160° it forms needles of 'benzonaphthone' $\text{C}_{12}\text{H}_8\text{O}_3$? which may be purified by successive boiling with aniline, and alcohol (Japp a Miller, *C J* 39, 221) This body is insoluble in ordinary solvents and in aqueous NaOH Aqueous KMnO_4 oxidises it to phthalic acid It is not affected by ZnEt, and therefore contains no hydroxyl—7 Boiling fuming HClAq forms a green amorphous powder, insol water, alcohol, and ether, but partially dissolving in HOAc forming a blue solution (Krappe a Schultz, *A* 210, 178)—8 Phenyl hydrazine reacts forming benzene azo-(α) naphthol $\text{C}_{10}\text{H}_6(\text{OH})\text{N}_2\text{Ph}$ —9 On adding aqueous ammonia to alcoholic naphthoquinone the liquid becomes brown and, by precipitation by water, a red brown amorphous body is formed, probably Laurent's 'carminaphthe' (Plimpton, *C J* 37, 641)—10 Yellow ammonium sulphide produces a red powder melting with complete decomposition at about 300° (Willgerodt, *B* 20, 2470)—11 When a solution of methylamine (acetate) is added slowly to a small quantity of naphthoquinone dissolved in alcohol, methylamido naphthoquinone $\text{C}_{10}\text{H}_6\text{O}_2\text{NMe}$ [232° uncol], is formed It crystallises from alcohol in red needles, which are reduced by SO_2 forming a substance crystallising in white needles (Plimpton, *C J* 37, 639)—12 When an alcoholic solution of naphthoquinone is mixed with excess of aqueous dimethylamine the solution neutralised with acetic acid and the liquid allowed to evaporate spontaneously, there is deposited a substance $\text{C}_{10}\text{H}_6\text{O}_2\text{NMe}_2$, which crystallises from alcohol in bright red needles [118°] (Plimpton, *C J* 37, 642)—13 With ethylamine acetate in alcoholic solution a brown liquid is formed If this is boiled with charcoal, filtered and evaporated deposits ethyl amido naphthoquinone $\text{C}_{10}\text{H}_6\text{O}_2\text{NEt}$ as red needles [140°] It may be sublimed, and is soluble in benzene, hardly so in light petroleum (Plimpton, *C J* 37, 640)—14 Aniline in alcoholic solution forms a red liquid. If this solution is heated to boiling and poured into water, the pp washed with dilute acetic acid and crystallised from alcohol, animal charcoal being used, the product is phenyl-amido-naphthoquinone $\text{C}_{10}\text{H}_6\text{O}_2\text{NPh}$ [191°] (Plimpton, *C J* 37, 635). It crystallises in red needles, v.

sol hot alcohol, benzene, and ether Insol light petroleum It forms a crimson solution in conc H_2SO_4 from which it is ppt unchanged by water It forms a purple solution in alcoholic potash It is not attacked by acetic or benzoic anhydride It is attacked by conc HCl at 170° and by aqueous SO_2 at 125° , aniline being among the products (Plimpton, *C J* 37, 635, cf Zincke, *B* 12, 1645, 14, 92)—15 *p* Bromo aniline forms, in like manner, when heated with (a)-naphthoquinone or oxy (a) naphthoquinone, red needles [266° – 269°] (Baltzer, *B* 14, 1899)—16 *m* Nitro-aniline yields $\text{C}_{10}\text{H}_7(\text{NHC}_2\text{H}_4\text{NO})_2$ [270°] (B) 17 *p* Nitro aniline forms $\text{C}_{10}\text{H}_7(\text{NHC}_2\text{H}_4\text{NO})_2$ melting above 270° , which may be reduced to $\text{C}_{10}\text{H}_7(\text{NH C}_2\text{H}_4\text{NH}_2)_2$ [177°] (B)—18 *Ethyl aniline* produces $\text{C}_{10}\text{H}_7(\text{NPhEt})_2$ crystallising in violet needles [155°] (Elsbach, *B* 15, 1810)—19 *o* Toluidine forms $\text{C}_{10}\text{H}_7(\text{NHC}_2\text{H}_4\text{NO})_2$ crystalising in red needles (Elsbach, *B* 15, 639)—20 *p*-Toluidine forms $\text{C}_{10}\text{H}_7(\text{NHC}_2\text{H}_4\text{NO})_2$ crystallising from alcohol in red needles [200°] and forming a crimson solution in conc H_2SO_4 (Plimpton, *C J* 37, 638, Elsbach, *B* 15, 687)—21 *Diphenyl amine* forms $\text{C}_{10}\text{H}_7(\text{NPh})_2$ O , crystallising from alcohol in needles [164°] (Plimpton)



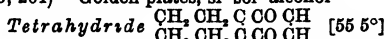
Formed, together with a greater quantity of the (β)-oxim of (β)-naphthoquinone, by the action of nitrous acid on (a) naphthol (Fuchs, *B* 8, 626) Prepared by the action of hydroxylamine hydrochloride on (a)-naphthoquinone (Goldschmidt, *B* 17, 2064) Yellowish needles, decomposing at about 190° V sol alcohol and ether, sl sol CS_2 and hot benzene Slightly volatile with steam Dissolves in alkalis, but is reppd by CO_2 Does not colour mordants (Kos tanecki, *B* 22, 1347) *Reactions*—1 *Nitric acid* converts it into di nitro (a) naphthol—2 Alcoholic K_2FeCy_4 forms nitro (a)-naphthol—3 PCl_5 forms 'β' di chloro naphthalene [68°]—4 *Phenyl cyanate* unites, forming $\text{C}_{10}\text{H}_7(\text{OH}) \text{NO CO NHPH}$, which crystallises in yellow prisms [170°] (Goldschmidt a Strauss, *B* 22, 3106)—5 Boiling with aniline and HOAc yields phenyl-(β)-amido (a) naphtho quinone anilide $\text{C}_{10}\text{H}_7(\text{NHPH}) \begin{array}{l} \text{O} \\ \text{NPh} \end{array}$ [187° cor] (Brömme, *B* 21, 893)—6 *p* Toluidine forms, in like manner, *p* tolyl-amido (a)-naphthoquinone *p* toluidide [183° cor] (B)—7 *ψ* Cumidine forms *ψ*-cumyl amido (a)-naphthoquinone *ψ* cumidide [181° cor]—8 *Naphthylamine* forms (a) naphthyl amido-(a)-naphthoquinone-(a) naphthalide [178° cor]—9 *Bromine* in HOAc forms di-bromo-(a) naphthoquinone oxim $\text{C}_{10}\text{H}_7\text{Br}_2 \begin{array}{l} \text{O} \\ \text{NOH} \end{array}$ [175°] (Brömme, *B* 21, 891) *Methyl ether* A'Me [100°] Yellow crystals, v sol alcohol, ether, and benzene, insol water, dissolves in H_2SO_4 with a yellow colour (Ilinski, *B* 17, 2589)

Di-oxim $\text{C}_{10}\text{H}_7(\text{NOH})_2$ [14°] [207°] Formed by boiling the mono oxim for two days with hydroxylamine and alcohol (Nietzki a Guter-mann, *B* 21, 433) Slender colourless needles, sol. alcohol and ether, decomposed by melting With Ac_2O it yields $\text{C}_{10}\text{H}_7(\text{NOAc})_2$ [160°] Tin and HCl reduce it to naphthylene-diamine.

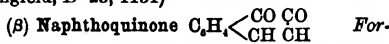
Alkaline KMnO_4 oxidises it to di-nitroso-naphthalene $\text{C}_{10}\text{H}_6(\text{NO})_2$, a yellow powder, exploding at 120°

Di-chloro-di-imide $\text{C}_{10}\text{H}_4(\text{NCl})_2$ [187°] Formed from naphthylene *p* diamine hydrochloride and a cold solution of bleaching powder (Friedlander a Böckmann, *B* 22, 591) Yellow needles (from alcohol and xylene), smelling strongly like quinone Reconverted by reducing agents to naphthylene-diamine Conc HClAq in HOAc converts it into di chloro (a) naphthoquinone [190°]

Di-phenyl di-imide $\text{C}_{10}\text{H}_4(\text{NPh})_2$ [187°] Formed by heating benzene azo (a) naphthylamine with aniline at 150° (Fischer a Hepp, *A* 256, 264) Golden plates, sl sol alcohol



Obtained by oxidising (a) naphthylamine tetrahydride with $\text{Na}_2\text{Cr}_2\text{O}_7$ and H_2SO_4 (Bamberger a. Lengfeld, *B* 23, 1131)



mation—By oxidation of amido-(β) naphthol, which is got by reducing its (a) oxim (nitroso (β) naphthol) (Stenhouse a Groves, *C J* 32, 47, 33, 415, *A* 189, 153, 194, 202, Liebermann a Jacobson, *A* 211, 40) The amido (β)-naphthol may also be obtained by reducing (β)-naphthol orange (v Azo-compounds)

Preparation—Amido-(β) naphthol hydrochloride (1 pt) is dissolved in saturated aqueous SO_2 , and when cold the solution is poured into ferric chloride solution (12 pts, containing 12 pts Fe_2O_3) The quinone is deposited as golden needles (Groves, *C J* 45, 298)

Properties—Orange needles (from alcohol) or red needles (from ether) Softens and decomposes at 115° – 120° Has no smell is not volatile with steam Dissolves in alkalis, forming a yellow solution, which absorbs oxygen from the air

Reactions—1 Boiling *nitric acid* forms phthalic acid—2 SO_2 reduces it to hydro (β) naphthoquinone HIAq acts in like manner (S a G) Tin and HClAq reduce it to 'di naphthyl dihydroquinone' $\text{C}_{10}\text{H}_8\text{O}$, crystallising in colourless needles [178°] (Korn, *B* 17, 8024) 3 Dilute H_2SO_4 forms, on warming, 'di naphthyl diquinhydrone' $\text{C}_{20}\text{H}_{12}\text{O}_4$, a black insoluble powder (S a G)—4 *Chlorine* passed into its solution in HOAc forms chloro naphthoquinone and the compound $\text{C}_6\text{H}_4 \begin{array}{l} \text{CO CO} \\ \text{CHCl CCl} \end{array}$ which crystallises in monoclinic needles (containing 2aq) [112°], and, when anhydrous, melts at 128° (Zincke, *B* 20, 2892)—5 An alcoholic solution of aniline forms the anilide $\text{C}_{10}\text{H}_7\text{NO}_2$, probably oxy (a) naphthoquinoneanilide $\text{C}_{10}\text{H}_7(\text{OH}) \begin{array}{l} \text{O} \\ \text{NPh} \end{array}$ (Liebermann a Jacobson, *A* 211, 75, Zincke, *B* 14, 1494) This substance crystallises in red or yellow needles melting above 240° insol water, sl. sol alcohol and hot benzene It is split up by boiling acids into aniline and oxy-(a)-naphthoquinone Boiling HOAc converts it into phenyl-amido-(a)-naphthoquinone With NaOEt and alkyl bromides it gives the compounds $\text{C}_{10}\text{H}_7(\text{OME})\text{O}(\text{NPh})$ [161°], $\text{C}_{10}\text{H}_7\text{Et}\text{NO}$ [104°], $\text{C}_{10}\text{H}_7\text{PrNO}$ [100°], and $\text{C}_{10}\text{H}_7\text{PrNO}$ [104°] (Zincke, *B* 15, 279). The so-called dianilide of

(β) naphthoquinone, which is formed by heating di-bromo (α)-naphthol with aniline, is probably the anilide of phenyl amido (β) naphthoquinone

$C_{10}H_7(NPhH) \begin{smallmatrix} \diagup O \\ \diagdown NPh \end{smallmatrix}$ It crystallises in orange red needles [179°] (Meldola, *C J* 45, 157, cf. Zincke, *B* 15, 481). This body has also been described as di-phenyl-di-imido-naphthol $C_{10}H_5(OH)(NPh)_2$ (Griess, *B* 13, 123). The corresponding di-*p* toluidine [c 175°] and di (β)-naphthalide [247°], prepared by the same reaction from di-bromo-(α)-naphthol, are probably constituted in a similar manner.—6 *Ethyl aniline* forms phenyl ethyl amido (β) naphthoquinone $C_{10}H_7(NPhEt)O_2$? which crystallises in thick red needles [165°], insol. NaOHAq, and is resolved by dilute HClAq into ethyl aniline and (β) oxy-(α) naphthoquinone (Elsbach, *B* 15, 691).—7 *o*-Toluidine forms $C_{10}H_7NO_2$, probably $C_{10}H_7(OH) \begin{smallmatrix} \diagup O \\ \diagdown NC_6H_5 \end{smallmatrix}$, which crystallises in red needles, sol. NaOHAq. It is not affected by HOAc at 150°. Boiling dilute HClAq splits it up into *o*-toluidine and (β) oxy (α) naphthoquinone (Elsbach, *B* 15, 689).—8 *p*-Toluidine forms the *p* toluidine $C_{10}H_7NO_2$, probably $C_{10}H_7(OH) \begin{smallmatrix} \diagup O \\ \diagdown NC_6H_4Me \end{smallmatrix}$, which crystallises in red needles [246°], sol. NaOHAq. On heating with dilute HClAq at 130° it forms (β) oxy (α) naphthoquinone, and this body is probably an intermediate product in the conversion of (β) naphthoquinone *p* toluidine into (α) naphthoquinone *p* toluidine, which takes place on heating with acetic acid at 150° (Elsbach, *B* 15, 686). It gives the following ethers— $C_{10}H_7MeNO_2$ [150°], $C_{10}H_7EtNO_2$ [137°], and $C_{10}H_7PrNO_2$ [139°] (Zincke a. Brauns, *B* 15, 1969).—9 Boiling with ammonium acetate forms (β) naphthazone

Phenyl hydrazide $C_{10}H_7 \begin{smallmatrix} \diagup O \\ \diagdown NHC_6H_5 \end{smallmatrix}$

Benzene-o azo-(α)-naphthol [138°] Formed by the action of phenyl hydrazine hydrochloride upon (β) naphthoquinone suspended in acetic acid (Zincke a. Bindewald, *B* 17, 3030). Long red needles, sol. hot alcohol and hot HOAc, insol. water. Does not combine with acids or bases. Yields amido naphthol on reduction with SnCl₂. Bromine in HOAc forms $C_{10}H_7Br_2N_2O$ [215°–219°]

o-Tolyl-hydrazide $C_{10}H_7N_2O_2$

$C_{10}H_7 \begin{smallmatrix} \diagup O \\ \diagdown NHC_6H_4 \end{smallmatrix}$ [156°] Red plates, with golden lustre, v. sol. alcohol. Reduced by SnCl₂ to amido naphthol. Nitric acid oxidises it to di-nitro-naphthol. Bromine forms $C_{10}H_7Br_2N_2O$ [254°]

p-Tolyl hydrazide $C_{10}H_7N_2O_2$

$C_{10}H_7 \begin{smallmatrix} \diagup O \\ \diagdown NHC_6H_4 \end{smallmatrix}$ [145°] Red needles (Zincke a. Bathgen, *B* 19, 2491). Forms $C_{10}H_7Br_2N_2O$ [186°]

(α) Oxim $C_{10}H_7 \begin{smallmatrix} \diagup C(NO)CO \\ \diagdown CH=CH \end{smallmatrix}$ (α) Nitroso-*(β)-naphthol* [109.5°] S (alcohol) 2.4 at 18°. Prepared by adding a concentrated aqueous solution of NaNO₂ (50 pts.) to a boiling solution of (β) naphthol (100 pts.) and ZnCl₂ (75 pts.) in spirit (600 pts.). After cooling, and standing for some hours, the brown zinc salt which separates is washed with alcohol, suspended in water

(1000 pts.), and digested for a short time with NaOH (90 pts.). After cooling, the sodium salt is filtered off, washed with a little water, and decomposed by cold HCl. The yield is 110 pts. of the naphthol (Henriques a. Ihmski, *B* 18, 704; cf. Groves, *C J* 45, 295, Stenhouse a. Groves, *C J* 32, 47, A 189, 146, Fuchs, *B* 8, 1026). *Properties*—Thin plates or short thick orange prisms. V. sol. boiling water, sl. sol. ligroin, v. sol. ether, benzene, and HOAc. Volatile with steam (when pure). *Reactions*—1

Aqueous ammonia at 100° forms a compound $C_{10}H_7N_2O$, probably the oxim of naphthoquinonimide $C_{10}H_7 \begin{smallmatrix} \diagup NH \\ \diagdown NOH \end{smallmatrix}$, which crystallises from alcohol in green needles [153°], soluble in alcohol, ether, acids, and alkalis. It is converted by nitrous acid into an unstable nitrosamine $C_{10}H_7 \begin{smallmatrix} \diagup N(NO) \\ \diagdown NOH \end{smallmatrix}$ [244°], which forms the salts $KC_{10}H_7N_2O_2$ and $AgC_{10}H_7N_2O_2$. Naphthoquinonimide oxim is reduced by ammonium sulphide to a base $C_{10}H_7N_2$ [92°–95°]. Naphthoquinonimide oxim forms the salts $KC_{10}H_7N_2O$, a red powder, $C_{10}H_7N_2OHCl$, yellow plates, ($C_{10}H_7N_2O$), H_2PtCl_4 , 2aq, orange needles, and $C_{10}H_7N_2OHNO_2$.—2 Dilute nitric acid forms nitro naphthol.—3 Ammonium sulphide reduces it to amido naphthol.—4 Chlorine passed into a cold solution of the oxim in chloroform yields

$C_{10}H_7 \begin{smallmatrix} \diagup C(NO)CO \\ \diagdown CH=CCl \end{smallmatrix}$ crystallising in yellowish red needles (Zincke a. Schmunk, *A* 257, 133). This body takes up more chlorine, forming

$C_{10}H_7 \begin{smallmatrix} \diagup C(NO)CO \\ \diagdown CCl=CHCl \end{smallmatrix}$ which crystallises in white needles [186°], and when treated with potash changes to the dichloro derivative $C_{10}H_7 \begin{smallmatrix} \diagup C(NO)CO \\ \diagdown CCl=CCl \end{smallmatrix}$ [166°].—5 Bromine added to its solution in chloroform unites forming $C_{10}H_7NO_2Br$, crystallising in colourless needles [131°], which dissolve in alkalis, changing to $C_{10}H_7 \begin{smallmatrix} \diagup C(NO)CO \\ \diagdown CH=CBBr \end{smallmatrix}$, crystallising in yellow needles [172°]. The last body is also formed by adding bromine to a solution of (β) naphthoquinone (α)-oxim in cold HOAc. Bromine added to a hot solution in HOAc forms a bromo naphthoquinone [201°] (Brömme, *B* 21, 386).—

6 *Phenyl cyanate* forms $C_{10}H_7 \begin{smallmatrix} \diagup O \\ \diagdown NO CO NPh \end{smallmatrix}$, crystallising in felted needles [128°] (Goldschmidt a. Strauss, *B* 22, 3106).—7 SO_2 or NaHSO₃, acting on its alcoholic solution, form amido-naphthol sulphonic acid (Schmidt, *J pr* [2] 42, 156).—8 *Aniline* in HOAc forms the same phenyl amido naphthoquinone anilide as it forms with the two other naphthoquinone oxims (Brömme, *B* 21, 893). Salts—NaA' green pp (from alcoholic solution), sl. sol. water—KA' lustrous green plates, sol. water and alcohol—NH₄A' green plates—CoA', brownish-red pp—CoA', purple-red pp, soluble without alteration in alcohol, aniline, phenol, conc. H₂SO₄, and HNO₃, it is very stable towards acids, alkalis, oxidising agents, and reducing agents (Ihmski a. Knorr, *B* 18, 699).—NiA', brownish-yellow pp, sl. sol. water and alcohol, easily decomposed by acids (difference from the cobalt salt).—FeA', black pp., insol. water, v. sol. aniline, forming a

dark-brown solution — CuA' , lustrous brown pp, sol dilute HClA' and 50 p c acetic acid — AgA' reddish-brown powder, insol water and alcohol $\text{Ag}(\text{NH}_4)\text{A}'$, slender green needles, insol water and alcohol — AgHA' , microcrystalline brown pp, sol alcohol (Ilinski, *B* 17, 2581, 18, 2728)

Methyl ether of the (a)-oxim

$\text{C}_6\text{H}_5\text{N} \begin{smallmatrix} \text{C(NOMe)} \\ \text{CH} \end{smallmatrix} \text{CO} \begin{smallmatrix} \text{CH} \\ \text{CH} \end{smallmatrix}$ [75°] Long yellow needles, v sol alcohol, ether, and benzene, m sol hot water, sl sol cold ligroin Forms a deep-red solution in H_2SO_4 . Yields (a) amido (β)-naphthol on reduction with tin and HClA' (Goldschmidt & Schmid, *B* 18, 571)

Ethyl ether of the (a) oxim

$\text{C}_6\text{H}_5\text{N} \begin{smallmatrix} \text{C(NOEt)} \\ \text{CH} \end{smallmatrix} \text{CO} \begin{smallmatrix} \text{CH} \\ \text{CH} \end{smallmatrix}$ [50°–60°] Needles (from alcohol-ligroin) (Ilinski, *B* 19, 341)

Benzoyl derivative $\text{C}_6\text{H}_5\text{N} \begin{smallmatrix} \text{C(NOBz)} \\ \text{CH} \end{smallmatrix} \text{CO} \begin{smallmatrix} \text{CH} \\ \text{CH} \end{smallmatrix}$ [114°]

(β)-Oxim $\text{C}_6\text{H}_5\text{N} \begin{smallmatrix} \text{C(NOBz)} \\ \text{CH} \end{smallmatrix} \text{CO} \begin{smallmatrix} \text{CH} \\ \text{CH} \end{smallmatrix}$ (β) Nitroso

(a)-naphthol [148°] (W), [152°] (G) This is the chief product of the action of nitrous acid on (a) naphthol (Fuchs, *B* 8, 626) It may be separated from the oxim of (a) naphthoquinone (*supra*), which accompanies it by means of their sodium salts (Worms, *B* 15, 1816) It is also obtained by boiling (β) naphthoquinone with hydroxylamine hydrochloride for half an hour (Goldschmidt, *B* 17, 215) Yellow needles (from benzene), almost insol cold water, m sol ether, v sol alcohol Somewhat volatile with steam Conc H_2SO_4 forms a deep-red solution Not decomposed by boiling alcoholic potash

Reactions — 1 Bromine acting on its solution in chloroform unites forming a dibromide $\text{C}_{10}\text{H}_6\text{N}_2\text{Br}_2$, crystallising in grey leaflets [155°] (Brömme, *B* 21, 390) Bromine acting on its solution in HOAc yields the bromo derivative $\text{C}_{10}\text{H}_6\text{N}_2\text{BrO}$, separating from alcohol in yellow crystals [175°] (B) — 2 Alkaline K_2FeC_4 oxidises it to (β) nitro (a) naphthol — 3 Nitric acid forms di nitro (a) naphthol and phthalic acid — 4 Phenyl cyanate unites with it, forming $\text{C}_{10}\text{H}_6\text{N} \begin{smallmatrix} \text{O} \\ \text{NO} \end{smallmatrix} \text{CO NHPH}$, crystallising from benzene in greenish yellow prisms (containing benzene) [120°] (Goldschmidt & Strauss, *B* 22, 3106) — 5 Aniline in HOAc forms the same phenyl-amido naphthoquinone amide as with the other two isomeric oxims (Brömme)

Salts — $\text{NH}_4\text{A}'$ lustrous green needles — NaA' (dried at 110°) reddish brown powder, insol conc NaOH , m sol water and alcohol — KA' lustrous green plates — BaA' , 2 aq bronzed plates — PbA' , dark brown scales, insol water — AgA' brownish-red pp

Methyl ether of the (β) oxim

$\text{C}_6\text{H}_5\text{N} \begin{smallmatrix} \text{C(NOMe)} \\ \text{CH} \end{smallmatrix} \text{CO} \begin{smallmatrix} \text{CH} \\ \text{CH} \end{smallmatrix}$ [98°] From the Ag salt and MeI (Fuchs) Formed also by heating (β)-naphthoquinone with the hydrochloride of the methyl ether of hydroxylamine (Goldschmidt & Schmid, *B* 18, 571, 2224) Yellowish green needles, v sol alcohol Reduced by tin and HCl to (β) amido (a)-naphthol

Ethyl ether $\text{C}_6\text{H}_5\text{N} \begin{smallmatrix} \text{C(NOEt)} \\ \text{CH} \end{smallmatrix} \text{CO} \begin{smallmatrix} \text{CH} \\ \text{CH} \end{smallmatrix}$ [101°] Flat greenish yellow needles

Benzoyl derivative $\text{C}_6\text{H}_5\text{N} \begin{smallmatrix} \text{C(NOBz)} \\ \text{CH} \end{smallmatrix} \text{CO} \begin{smallmatrix} \text{CH} \\ \text{CH} \end{smallmatrix}$ [162°]

Di-oxim $\text{C}_{10}\text{H}_6\text{N}_2(\text{NOH})_2$ [149°] (G & S), [166°] (B), [181°] (K & M) Formed by warming either the (a) or the (β) oxim in con-

centrated methyl alcoholic solution with hydroxylamine hydrochloride at 100° (Goldschmidt & Schmid, *B* 17, 2066, Brömme, *B* 21, 392) Formed also by adding an alkaline solution of hydroxylamine to a cold solution of the (a) oxim in caustic soda (Kehrmann & Messinger, *B* 23, 2816) Yellow needles, forming a reddish yellow solution in alkalis, and a dark red solution in H_2SO_4 . On warming with dilute H_2SO_4 or with alkalis it is converted into an anhydride Phenyl cyanate also brings about this change (Goldschmidt & Strauss, *B* 22, 3107) Phenyl hydrazine combines, forming $\text{C}_{10}\text{H}_6\text{N}_4\text{O}$, which crystallises from alcohol in tufts of long needles [105°–138°] (Polonovsky, *B* 21, 182) Stannous chloride reduces it to naphthylene (1,2) diamine Alkaline K_2FeC_4 oxidises it to di nitroso naphthalene $\text{C}_{10}\text{H}_6\text{N}_2(\text{NO})_2$, which crystallises in needles [127°], insol water and alkalis, v sol alcohol (Leuckart, *B* 19, 174, 349) The dioxim colours iron and cobalt solutions brown — $\text{KC}_{10}\text{H}_6\text{N}_2\text{O}$, brownish red amorphous pp, obtained by adding ether to an alcoholic solution of the dioxim and KOEt (Ilinski, *B* 19, 342) — $\text{AgC}_{10}\text{H}_6\text{N}_2\text{O}$, dark red pp, obtained by adding ammoniacal AgNO_3 to an alcoholic solution of the dioxim

(a) *Methyl ether of the dioxim*

$\text{C}_6\text{H}_5\text{N} \begin{smallmatrix} \text{C(NOMe)} \\ \text{CH} \end{smallmatrix} \text{C(NOH)} \begin{smallmatrix} \text{CH} \\ \text{CH} \end{smallmatrix}$ [159°] Formed by action of hydroxylamine on the methyl ether of the (a) oxim Insol water, sol alkalis

(β) *Methyl ether of the dioxim*

$\text{C}_6\text{H}_5\text{N} \begin{smallmatrix} \text{C(NOH)} \\ \text{CH} \end{smallmatrix} \text{C(NOMe)} \begin{smallmatrix} \text{CH} \\ \text{CH} \end{smallmatrix}$ Formed from the silver salt of the dioxim and MeI, and also by the action of hydroxylamine on the methyl ether of the (β) oxim Yellow oil, turning brown in air Sol alkalis

(a) *Ethyl ether of the dioxim*

$\text{C}_6\text{H}_5\text{N} \begin{smallmatrix} \text{C(NOEt)} \\ \text{CH} \end{smallmatrix} \text{C(NOH)} \begin{smallmatrix} \text{CH} \\ \text{CH} \end{smallmatrix}$ [153°] Formed by warming an alcoholic solution of the ethyl ether of the (a) oxim with hydroxylamine hydrochloride Green needles (from alcohol), insol water — $\text{KC}_{10}\text{H}_6\text{N}_2\text{O}_2$ brown needles

Anhydride of the dioxim

$\text{C}_{10}\text{H}_6\text{N} \begin{smallmatrix} \text{N} \\ \text{N} \end{smallmatrix} \text{O}$ [78°] Formed from the dioxim by the action of AcCl , alkalis, or acids Formed also by heating the (a) or (β) oxim with an alcoholic solution of hydroxylamine hydrochloride at 150° (Goldschmidt, *B* 17, 216, 801) Long colourless monoclinic needles (from ligroin) Insol alkalis

Peri Naphthoquinone $\text{C}_{10}\text{H}_6\text{O}_2$ i.e.

$\text{CH} \begin{smallmatrix} \text{O} \\ \text{C} \end{smallmatrix} \text{—} \text{C} \begin{smallmatrix} \text{O} \\ \text{C} \end{smallmatrix} \text{CH}$ Occurs in small quantity in the product of the oxidation of di bromo (a) naphthol by fuming nitric acid (Meldola & Hughes, *C J* 57, 632) Slender, pale yellow needles, no definite melting point observed Blackens and decomposes above 220° Sl sol boiling dilute NaOH Gives an azo compound with phenyl hydrazine Not reduced by cold aqueous SO_2 . Zinc-dust and HOAc reduces it to the corresponding dioxynaphthalene, which blackens at 205° and forms a diacetyl derivative melting at 227°.

References. — AMIDO, AMIDO OXY-, BROMO,

BROMO AMIDO, CHLORO, NITRO-, and OXY NAPHTHOQUINONE

NAPHTHOQUINONE-PHENAZINE

$C_{12}H_8 \begin{smallmatrix} \text{CO} \text{CN} \\ \text{CO} \text{CN} \end{smallmatrix} C_{12}H_8$. Formed by heating o nitro phenyl amido naphthoquinone with alcoholic ammonium sulphide (Leicester, *B* 23, 2797) Green plates (from alcohol) Its alcoholic solution is brown with green fluorescence

NAPHTHOQUINONE PHENYL-HYDRAZIDE

IDE v Benzene azo naphthol

NAPHTHOQUINONE TOLAZINE

$C_{12}H_8 \begin{smallmatrix} \text{CO} \text{CN} \text{C} \text{CH} \text{CH} \\ \text{CO} \text{CN} \text{C} \text{CH} \text{CH} \text{CM}_6 \end{smallmatrix}$ Formed by reduction of o nitro tolyl amido (a) naphthoquinone with alcoholic ammonium sulphide (Leicester, *B* 23, 2797) Steel blue plates with green lustre, forming a green powder Conc H_2SO_4 forms a green solution Its solutions in alcohol or HOAc are greenish yellow with faint moss green fluorescence

NAPHTHOQUINONE TOLYL HYDRAZIDE

v Toluene azo naphthol

NAPHTHOQUINOXALINE $C_{12}H_8N_2$ s.e

$C_{12}H_8 \begin{smallmatrix} \text{CN} \text{CH} \\ \text{CH} \text{CH} \text{CN} \text{CH} \end{smallmatrix}$ [62°] Formed by warming naphthylene (aβ) diamine with the bisulphite compound of glyoxal in presence of some HOAc (Hinsberg, *B* 23, 1394) Small colourless needles, v sol alcohol and ether, sl sol hot water Volatile with steam Conc H_2SO_4 gives a deep red colour, turned yellow by addition of water—The sulphate crystallises well The platinumchloride is sl sol water

NAPHTHOQUINOXIM v Oxim of NAPHTHOQUINONE

NAPHTHOSTYRIL v Lactone of peri AMIDO-

NAPHTHOIC ACID

NAPHTHOTOLAZINE v TOLUNAPHTHAZINE

(a) NAPHTHOKINDOLE $C_{12}H_8NO$ s.e

$C_{10}H_6 \begin{smallmatrix} \text{NH} \\ \text{CH} \end{smallmatrix} \text{CO}$ [245°] Formed by the action of mineral acids on the sodium salt of (a) naphthindole sulphonic acid (Hinsberg, *B* 21, 116) Colourless needles (from alcohol) Yields a nitroso derivative, which on reduction and subsequent oxidation forms (a) naphthosatin

(β) Naphthoxindole $C_{10}H_6 \begin{smallmatrix} \text{NH} \\ \text{CH} \end{smallmatrix} \text{CO}$

[234°] Formed by the action of mineral acids on (β) naphthindole sulphonic acid (Hinsberg, *B* 21, 114) Faint greenish needles Sl sol water, v sol alcohol, ether, HOAc Not attacked by mineral acids Conc KOHAq dissolves it without change Baryta water at 150° in sealed tubes yields the Ba salt of a strong acid $NaNO_2$ in HOAc solution gives a nitroso-derivative [c 240°], crystallising in yellowish red needles M sol alcohol, ether, HOAc, sl sol water

NAPHTHOXY-ACETIC ACID v Naphthyl derivative of GLYCOLLIC ACID, vol II p 639

NAPHTHOYL BENZOIC ACID $C_{12}H_8O_2$ s.e

$C_{10}H_6 \begin{smallmatrix} \text{CO} \text{C}_6H_5 \\ \text{CO} \text{C}_6H_5 \end{smallmatrix}$ [173.5°] Formed by the action of phthalic anhydride on naphthalene in presence of $AlCl_3$ (Ador a Crafts, *C R* 88, 1355) Small white prisms (from dilute alcohol) Its Ba salt crystallises from alcohol in very hygroscopic needles Conc H_2SO_4 converts it into naphthantraquinone $C_{12}H_8 \begin{smallmatrix} \text{CO} \\ \text{CO} \end{smallmatrix} C_{12}H_8$ (Elbs, *B* 19, 2209)

NAPHTHOYL CYANIDE v Nitrile of NAPH-

THYL-GLYOXYLIC ACID

NAPHTHOYL FORMIC ACID v. NAPHTHYL-

GLYOXYLIC ACID

(aa) **DINAPHTHYL $C_{20}H_{14}$ s.e ($C_{10}H_7$)₂** Mol. w 254. [154°] (above 360°) V D 8.67 (calc 8.77)

Formation—1 Together with phthalic acid and other bodies by heating naphthalene with MnO_2 and dilute H_2SO_4 (Lossen, *A* 144, 77)—2 By distilling (ββ) dioxy dinaphthyl (1 pt) with zinc dust (12 pts), the yield being 65 p.c. of the theoretical (Walder, *B* 15, 2170, Julius, *B* 19, 2549)—3 By distilling (β) 'dinaphthyl diquinone' $C_{20}H_{14}O_2$ with zinc dust (Korn, *B* 17, 3019)—4 By diazotising di amido dinaphthyl (naphthidine), and boiling the tetra azo dinaphthyl sulphate with alcohol (Nietzki & Groll, *B* 18, 3256)

Preparation—Naphthalene is boiled with MnO_2 and H_2SO_4 (diluted with more than its own bulk of water) The product is boiled with water, filtered, and the residue extracted with alcohol, the alcohol boiled off (in a current of air), and the residue distilled The fraction coming over above 360° is boiled with light petroleum and animal charcoal, filtered, and allowed to crystallise (Watson Smith, *C J* 35, 225) No ββ compound is formed

Properties—Colourless plates (by sublimation), v sol benzene, CS_2 , HOAc, and ether, m sol alcohol Its solutions exhibit blue fluorescence (K)

Picric acid compound

$C_{20}H_{14} \cdot 2C_6H_3(NO_2)_3OH$ [145°] Reddish-brown needles

(αβ) **Dinaphthyl $C_{20}H_{14}$** [76°] (S), [80°] (W) V D 8.78 (calc 8.77) Formed, together with a larger quantity of the (ββ) and a very little of the (aa) isomerides, by passing the vapour of naphthalene, mixed with $SbCl_3$, through a red hot tube (Watson Smith, *C J* 32, 559) Small six sided tables, m sol alcohol, ether, and benzene than the (ββ) isomeride

Picrate $C_{20}H_{14} \cdot C_6H_3(NO_2)_3OH$ [156°] (Wegscheider, *B* 23, 3199)

(ββ) **Dinaphthyl $C_{20}H_{14}$** *Isodinaphthyl* [187°] (S), [189°] (G & T) V D 8.73 (calc 8.77)

Formation—1 By passing naphthalene through a red hot tube (Watson Smith, *C J* 24, 1184)—2 By passing naphthalene and CCl_4 through a tube at dull red heat $4C_{10}H_8 + CCl_4 = 2C_{20}H_{14} + 4HCl + C$ (Watson Smith, *C J* 35, 229)—3 By passing through a red hot tube, naphthalene with chloroform, (a) bromo naphthalene, $SbCl_3$ or $SnCl_4$ —4 By heating to redness (a) bromo naphthalene with naphthalene and soda lime (S)—5 By passing $C_{10}H_8$ and $C_{10}H_7Br$ through a red hot tube packed with Fe_2O_3 (S)—6 By distilling aluminium (β) naphthol (Gladstone a Tribe, *C J* 41, 16)—7 By dropping isomyl chloride upon naphthalene (100 g) and $AlCl_3$ (25 g) at 120°, pentane being also formed (Roux, *Bl* [2] 41, 379)

Properties—Plates (from benzene), with slight blue fluorescence Much less sol alcohol, ether, and benzene than either of its isomerides On oxidation with $KMnO_4$ or dilute HNO_3 at 160° it yields phthalic acid CrO_3 in HOAc produces a quinone $C_{20}H_{10}O_4$.

Picric acid compound
 $C_{10}H_7O_6N_3$ (NO₂)₃OH. [184°] Orange prisms (Wegecheider, *B.* 23, 3200)

($\beta\beta$)-Dinaphthyl sulphonic acid $C_{20}H_{13}SO_3$ \pm $C_{10}H_7$, $C_{10}H_7SO_3H$ Prepared by heating ($\beta\beta$)-dinaphthyl (15 g) with conc H_2SO_4 (3 g) for 5 hours at 200° (Watson Smith a Takamatsu, *C* J 39, 552) —CaA', 2aq white needles, m. sol. hot water, insol. alcohol, ether, and benzene —BaA', 2aq?

($\beta\beta$) Dinaphthyl disulphonic acid
 $C_{10}H_7(SO_3H)C_{10}H_7(SO_3H)$ Two isomeric acids of this formula are formed by heating ($\beta\beta$)-dinaphthyl (10 g) with conc H_2SO_4 (7 g) at 190° for 5 hours (S a T) One gives an insoluble barium salt, the other forms BaA' 2aq, v sol water, and PbA' 2aq, a yellowish white crystalline powder

($\beta\beta$) Dinaphthyl tetrasulphonic acid
 $C_{10}H_7(SO_3H)_2C_{10}H_7(SO_3H)_2$ Formed by heating ($\beta\beta$)-dinaphthyl with an excess of Nordhausen sulphuric acid (S a T) —PbA' 6aq v sol. water, insol alcohol, ether, and benzene

References —DI AMIDO-, DI BROMO-, TETRA-CHLORO-, and NITRO- DINAPHTHYL

NAPHTHYL-ACETAMIDINE $C_{17}H_{13}N_2$ \pm $CH_3C(NH)NHC_{10}H_7$. (a) *Naphthyl-ethenyl-amidine* Prepared by the action of (a) naphthylamine hydrochloride on acetonitrile at 165° (Bernthsen a Trompeter, *B.* 11, 1758) —B'HCl soluble prisms —B'H₂PtCl₆ small yellow tables —B'H₂C₂O₄ small crystals —B'H₂SO₄ white crystals —B HNO₃ oil

Di (a)-naphthyl-acetamidine $C_{22}H_{15}N_2$ \pm $CH_3C(NO_2H)NHC_{10}H_7$ Formed from (a)-naphthylamine (6 mols), AcCl (3 mols), and PCl₅ (Hofmann, *J* 1865, 415) Resinous

Di-(β)-naphthyl acetamidine
 $CH_3C(NO_2H)NHC_{10}H_7$ [168°] Formed by heating (β) naphthylamine (6 mols) with AcCl (3 mols) and PCl₅ (1 mol) at 150° (Maschke, *O C* 1886, 824)

(a)-NAPHTHYL-ACETIC ACID $C_{17}H_{13}O_2$ \pm $C_{10}H_7CH_2CO_2H$ [131°] Prepared by heating (a)-naphthyl-glyoxylic acid with HI and P (Boessneck, *B.* 16, 641) Long silky needles Sol alcohol, ether, acetic acid, benzene, and hot water, al sol cold water

Amides $C_{10}H_7CH_2CONH_2$ [181°] (B), [154°] (W) Formed from the acid (B) and perhaps also by the action of yellow ammonium sulphide on (a) naphthyl methyl ketone (Willgerodt, *B.* 21, 534) Colourless needles, sol benzene, acetic acid, ether, CS₂, and hot water

Nitrile $C_{10}H_7CH_2CN$ (above 300°). Formed from the amide and P₂O₅ Oil

DI-NAPHTHYL-ACETYLENE

$C_{10}H_7CC_2H_5$ [225°] (above 360°) Obtained by distilling *exo* di chloro di naphthyl-ethylene or *exo*-tri chloro di naphthyl-ethane (1 pt) with soda-lime (10 pts) (Grobowski, *B.* 11, 801) Long silky needles (from alcohol)

(a) NAPHTHYL-ACRYLIC ACID $C_{17}H_{13}O_2$ \pm $C_{10}H_7CH=CHCO_2H$ (a) *Naphthocinnamic acid* [207°] (L), [212°] (B) S 014 Obtained by heating naphthoic aldehyde (2 pts) with sodium acetate (1 pt) and Ac₂O (20 pts) at 170° (Lugl, *G.* 11, 894, Brandis, *B.* 22, 2155) Needles, m. sol. hot water, v sol. alcohol and ether Oxidised by KMnO₄ to naphthoic aldehyde and naphthoic acid Combines with bromine form-

ing $C_{10}H_7CHBrCHBrCO_2H$ [189°]. HBr at 100° yields $C_{10}H_7CHBrCHBrCO_2H$ [216°] —CaA', plates —BaA', needles —CuA', —AgA': white pp, blackening on exposure

NAPHTHYL-ALLYL-THIO UREA $C_{17}H_{13}N_2S$ \pm $C_{10}H_7NHCSNHC_2H_5$ [145°] Formed from (a) naphthylamine and oil of mustard (Zinin, *A.* 84, 346, Prager, *B.* 22, 3000) Crystals, insol water, v sol alcohol and ether

Dinaphthyl-allyl ψ -thiourea v Di NAPHTHYL-IMIDO THIO CARBAMIC ETHER

(a) NAPHTHYL-AMIDO-ACETIC ACID

$C_{17}H_{13}NO_2$ \pm $C_{10}H_7NHCH_2CO_2H$ *Naphthyl-glycocol* [199°] (B a N), [198°] (J), [192°] (M) Formed from (a) naphthylamine, chloroacetic acid, and NaOAc at 100° (Bischoff a Nastvogel, *B.* 22, 1808, Jolles, *B.* 22, 2372, Mauthner, *M.* 10, 251, Forte, *G.* 19, 361) Needles, insol water, almost insol ether, al sol alcohol, v sol acetone Forms a platinumchloride and a nitrosamine The Ca salt distilled with calcium formate gives a product crystallising in colourless plates [163°] —CaA', 3aq tufts of needles —BaA', 2aq —CuA', small plates —AgA' aq silvery plates

Anhydride $(C_{10}H_7NHCH_2CO)_2O$ [269°] Formed by heating the acid at 230° Scales, insol ether, sol alcohol

Acetyl derivative [154°] —BaA', 6aq

(β) Naphthyl-amido acetic acid

$C_{17}H_{13}NHCH_2CO_2H$ [135°] Formed by heating (β) naphthylamine (2 mols) with a solution of chloroacetic acid (1 mol) at 100° (Jolles, *B.* 22, 2372, Bischoff, *B.* 23, 2005) Minute crystals (from water), v sol alcohol, ether, and HOAc Yields a nitrosamine $C_{17}H_{13}N(NO)CH_2CO_2H$ crystallising from MeOH in reddish plates [126°]

Salts —The Ag salt decomposes on drying in the air (β) Naphthylamine salt $C_{10}H_7NH_2HA'$ [116°] Needles

(a) NAPHTHYL AMIDO-CROTONIC ACID

Ethyl ether $C_{10}H_7NHCH(Me)CHCO_2Et$ [45°] Formed by the action of acetoacetic ether on (a) naphthylamine (Conrad a Lampach, *B.* 21, 531) White silky needles, sol ether and benzene Yields on heating to 240° (Py 13) oxymethyl (a)-naphthoquinoline, with elimination of alcohol

(β) Naphthyl-amido-crotonic acid *Ethyl ether* $C_{10}H_7NHCH(Me)CHCO_2Et$ [66°] Formed by the action of acetoacetic ether on (β) naphthylamine at 100° (Conrad a Lampach, *B.* 21, 532) Large prisms (from alcohol) Yields (Py 13) Oxy methyl (β)-naphthoquinoline on heating to 240°

(β)-NAPHTHYL AMIDO-ETHYL OXIDE

$C_{17}H_{13}NO$ \pm $C_{10}H_7OC_2H_4NH_2$ Formed by the action of alcoholic ammonia on the chloroethyl ether of (β) naphthol (Koelle, *B.* 13, 1955) Amorphous mass —B'HCl aq. needles —B'H₂PtCl₆ needles

(a)-NAPHTHYL-AMIDO-METHYL-MALON-

AMIC ACID $CH_2C(NHC_2H_5)(CO_2H)(CONH_2)$

Ethyl ether A'Et [159°] Formed by dissolving α -cyano- α -(a)-naphthyl-amido-propionic ether (*q v*) in conc H_2SO_4 (Gerson, *B.* 19, 2969) Long white needles, al sol cold water, more easily in hot water with a beautiful green fluorescence, v sol alcohol and ether

(a) - NAPHTHYLAMIDO - (β) - NAPHTHO-QUINONE (a)-naphthylamide $C_{10}H_7N_2O$ *ts.*

$C_{10}H_7(NHC_6H_5) < \overset{O}{\underset{||}{N}} C_6H_5$, [178° cor] Formed from (a) naphthoquinone oxim and (a) naphthylamine (Brömme, B 21, 394) Forms a violet solution in alcohol and a blue solution in conc H_2SO_4 . Gives a green colour on heating with H_2SO_4 .

(β) Naphthyl-amido (β) naphthoquinone (β)-naphthylamide $C_{10}H_7(NHC_6H_5) < \overset{O}{\underset{||}{N}} C_6H_5$

[247°] Formed by heating di bromo (a)-naphthol with (β) naphthylamine (Meldola, C J 45, 160) Dull, reddish, fibrous needles. Feebly basic. Insol alcohol, but dissolves when boiled with alcohol and HCl, forming a violet solution. Dissolves in hot toluene or chloroform, giving a reddish brown liquid.

DI-(a) NAPHTHYL DI AMIDO-OIAZTHIOLE

$C_{22}H_{14}N_4S$ *ts* $S < \overset{C(NHC_6H_5)}{\underset{C(NHC_6H_5)}{N}} N$ [136°] Obtained by oxidising (a) naphthyl thio urea with hydrogen peroxide and dilute HClAq (Hector, B 23, 359) Crystallises from alcohol in white needles containing HOEt (1 mol) and melting at 104° Insol water. Cyanogen, passed into its warm solution, forms $C_{22}H_{14}N_4SCyN_4$ [203°]

Salts — $B^+H^+PtCl_6^-$ [225°] — **Picrate** $B^+C_6H_4(NO_2)_3OH^- + EtOH$ [below 100°] Small yellow grains — $B^+AgNO_3^-$ Pp, insol alcohol

Acetyl derivative $C_{22}H_{14}AcN_4S$ [263°] Needles (from alcohol)

Benzoyl derivative $C_{22}H_{14}BzN_4S$ [270°]**Di (β) naphthyl di-amido oiazthiole**

$C_{22}H_{14}N_4S$ [100°–117°] Prepared from (β)-naphthyl thio urea and H_2O_2 . Grey powder (from alcohol), more sol alcohol than the (a)-isomeride. Cyanogen passed into its warm alcoholic solution forms $C_{22}H_{14}N_4SCyN_4$ [200°] — $B^+H^+PtCl_6^-$ Begins to decompose at 236°–240° — $B^+C_6H_4(NO_2)_3OH^-$ [178°] Yellow powder, sl sol alcohol — $B^+AgNO_3^-$ white pp

Acetyl derivative $C_{22}H_{14}AcN_4S$ [203°] Needles (from alcohol)

Benzoyl derivative $C_{22}H_{14}BzN_4S$ [247°]

(β) NAPHTHYL PHENYL-AMIDO ETHYL OXIDE $C_{18}H_{11}NO$ *ts* $C_{10}H_7OCH_2NHC_6H_5$ [75°] Formed by the action of aniline on the chloro ethyl ether of (β) naphthol (Koelle, B 13, 1955)

TRI (β) NAPHTHYL - TRI AMIDO TRI-PHENYL CARBINOL $(C_{10}H_7NH C_6H_5)_3COH$ Formed by heating para rosaniline with (β) naphthylamine (Meldola, C N 47, 133, 147) Dyes wool or silk blue

TRI - NAPHTHYL - TRI - AMIDO - TRI-PHENYL CARBINYL CHLORIDE $C_{10}H_7N_3Cl$ *ts* $(C_{10}H_7NH C_6H_5)_3CCl$ C_6H_5NHCl $C_{10}H_7$ Phenyl-

(a) naphthylamine blue. Formed by heating phenyl (a) naphthylamine with oxalic acid (Hausdörfer, B 23, 1965) Dark brown powder, v sol hot aniline, sl sol cold alcohol, insol ether and benzene

NAPHTHYL-AMIDO-ISOSUCCINAMIC ACID

(a)-NAPHTHYLAMINE $C_{10}H_7N$ *ts* $C_{10}H_7 < \overset{C(NH_2)}{CH} \overset{OH}{CH} = CH$ **Naphthalamine** **Naphthal-amine** Mol w 148 [50°]. (300°) V.D (at

194°) 72.6 (calc 71.5) (Eykmann, B 22, 2757) S. 167 in the cold (Ballo, B 3, 875)

Formation — 1 By reduction of nitro-naphthalene by alcoholic ammonium sulphide (Zinin, J pr 27, 143), by iron and acetic acid (Béchamp, A Ch [3] 42, 186, Schützenberger a Willm, C R 47, 82, Ballo, B 3, 288, 673), or by tin or zinc and HClAq (Böttger, D P J 173, 480) — 2 By heating (a) naphthol with four times its weight of ammoniacal $CaCl_2$ to 270° for 8 hours, the yield being 74 p.c (Benz, B 16, 14, cf Calin, B 15, 616)

Preparation — Granulated nitronaphthalene (600 kilos) is slowly added to a warm mixture of iron borings (800 kilos), hydrochloric acid (40 kilos), and some water. The mixture is agitated and kept at about 50° by blowing in steam for 7 hours, after which milk of lime (containing 50 kilos of CaO) is added. The mixture is distilled in a current of superheated steam (Witt, Chem Industrie, 10, 215)

Properties — White silky needles, with powerful odour. May be sublimed. V sol alcohol and ether. Colours pine wood yellow. When not quite pure it rapidly becomes coloured in the air. Does not turn red litmus blue. Aqueous solutions of its salts give a blue pp with $FeCl_3$, $AgNO_3$, auric chloride, platinum chloride, $SnCl_4$, $HgCl_2$, chromic acid, H_2O_2 , and other oxidising agents (Pina, A Ch [3] 31, 217, 78, 64, Schiff, A 101, 92, Wurster, B 22, 1910) This blue pp, 'oxynaphthylamine' $C_{10}H_7NO$, is amorphous, v sol chloroform, and not reduced by SO_2 (Schiff, A 129, 255). If to a solution of (a) naphthylamine in alcohol or HOAc there be added a small quantity of nitrous acid and a little HClAq an intense purple colour is produced (Liebermann, A 183, 265)

Reactions — 1 Heated with $ZnCl_2$ or $CaCl_2$ at 280° it splits up to some extent into NH_3 and dinaphthylamine — 2 Distillation over PbO yields naphthazine — 3 Boiling chromic acid mixture oxidises it to naphthoquinone, phthalic acid, and other products (Reverdin a Nöling, B 12, 2306) — 4 Heated with nitro naphthalene and some HClAq at 200° it yields 'tri naphthylene diamine' $C_{30}H_{18}N_4$ (Salzmänn a Wichelhaus, B 9, 1107) — 5 Urea (1 pt) heated with (a)-naphthylamine (2 pts) at 120° forms only dinaphthyl urea, but both mono and di naphthylureas are formed on heating (a) naphthylamine hydrochloride with urea at 160° (Pagliani, G 9, 28) — 6 Cyanic acid forms (a) naphthyl urea — 7 Alkyl-thio-carbimides form alkyl naphthylthio ureas — 8 Alcoholic CS_2 forms di naphthylthio urea — 9 MeOH and $ZnCl_2$ at 190° form the methyl ether of (a) naphthol (Hantzsch, B 13, 1347) — 10 (a) naphthylamine hydrochloride heated with o-amido-azo-compounds yields azine colouring matters (eurydines) — 11 Readily converted into naphthalene by diazotising, and pouring the alkaline solution of the diazo-compound into a solution of $SnCl_4$ in aqueous NaOH (Friedländer, B 22, 587) — 12 SiF_4 in benzene solution forms minute crystals of $Si(NHC_6H_5)_4$ (Reynolds, C J 55, 482) — 13 The bisulphite warmed with benzoic aldehyde forms $C_{10}H_7CH_2NO_2$, a yellow powder (Papasogli, A 171, 188) — 14 Benzoic aldehyde and pyruvic acid form the compound $C_{10}H_7 < \overset{N^+O^+}{C} \overset{N^+O^+}{C} (CO_2H) > OH$ crystallising in yellow

needles [300°] (Doebner & Kuntze, *A* 249, 109)
 15 *Benzil* reacts on heating, forming the compounds $C_6H_5C(NC_6H_5)_2$, CO , C_6H_5 , [139°] and $C_6H_5C(NC_6H_5)_2C(NC_6H_5)_2$, C_6H_5 , [219°] both crystallising in yellow needles (Bandrowsky, *M* 9, 685) — 16 *a* cone boiling aqueous solution of *alloxan* forms a compound $C_{10}H_7N_3O_4$, which separates in transparent white needles, insol water, and coloured green by H_2SO_4 . Alkalis convert it into a crystalline acid $C_{10}H_7N_3O_5$ (Pellizzari, *G* 17, 409) — 17 *Cyanuric chloride* forms $N_3C_6Cl_3(NHC_6H_5)_3$, [149°], $N_3C_6Cl_3(NHC_6H_5)_2$, [215°], or $N_3C_6(NHC_6H_5)_3$, [228°], according to the proportions used (Fries, *B*, 19, 242, *C* *J* 49, 314) — 18 *Citraconic acid* at 145° forms $C_{10}H_7N_3C_6H_5O_4$, [143°] (360°) (Morawski & Glaser, *M* 9, 284) — 19 *Itaconic acid* heated with (a) naphthylamine in aqueous solution forms $C_{10}H_7N_3COCH_2CH_2COOH$, a white crystalline powder [206°], m sol hot alcohol (Scharfenberger, *A* 254, 151) — 20 *Chloro acetic ether* in ethereal solution forms $C_{10}H_7NO_2Cl$, crystallising in prisms [75°] (Bender, *B* 20, 2750)

Salts — B^+HCl^- Needles (by sublimation) or scales (from alcohol), v e sol water, alcohol, and ether — $B^+H_2PtCl_6^-$ yellow pp, sl sol water — B^+HBr^- — $B^+H_2SO_4^-$ — $B^+H_2SO_4^-$ 2aq — $B^+HNO_3^-$ scales — $B^+H_2C_2O_4^-$ stellar groups of small laminae — $B^+H_2C_2O_4^-$ nodules Yields on distillation $C_6O_2(NHC_6H_5)_2$ and the formyl derivative $CHO.NHC_6H_5$, (Zinn, *A* 108, 228) — $B^+H_2PtCl_6^-$ crystals (Scholtz, *M* 1, 905) — $B^+H_2SO_4^-$ pearly rosettes — *Citraconate* [99°] Formed by mixing solutions of (a) naphthylamine and citraconic acid in benzene (Morawski & Glaser, *M* 9, 285) — *Phenate* $B^+C_6H_5OH^-$ [30 1°] Formed by heating phenol with (a) naphthylamine (Dyson, *C* *J* 43, 468) Needles (from light petroleum) — *Benzene sulphate* [225°] (Norton & Westenhof, *Am* 10, 129) — *Toluene p-sulphonate* [239°] (Norton & Otten, *Am* 10, 140)

Formyl derivative $C_{10}H_7NH(COH)$ [139°] White silky needles (Tobias, *B* 15, 2447)

Acetyl derivative $C_{10}H_7NHAO$ [159°] Formed from the base by treatment with $AcCl$, Ac_2O , or $HOAc$ (Rother, *B* 4, 850, Tommasi, *C* *R* 76, 1267, Liebermann, *A* 183, 229) Formed also by heating (a) naphthol with ammonium acetate White silky needles, sl sol boiling water, v sol alcohol On heating with sulphur the products are ethenyl amido-naphthyl mercaptan and $C_{10}H_7<N>C<S>C<S>C_{10}H_7$, which crystallises in yellow plates [above 300°] (Hofmann, *B* 20, 1801) The only products of the nitration of acetyl-(a)-naphthylamine are the o- and p-nitro-derivatives of melting points [199°] and [190°] respectively, the supposed isomeride of melting-point [170°] is a molecular compound of the o- and p- bodies, and the so called 'β- and δ-nitroacetonaphthalides' are diacetyl derivatives of the same two nitro-(a) naphthylamines (Lellmann & Remy, *B* 19, 796) Chlorine passed into its solution in $HOAc$ forms $C_{10}H_7Cl_2NHAO$ [214°] (Cleve, *B* 20, 448)

Chloro-acetyl derivative $C_{10}H_7NClO$ *ts.* $C_{10}H_7NHCOCH_2Cl$ [121°] Formed from

naphthylamine and chloro acetyl chloride (Tommasi, *Bl* [2] 20, 21, Abenius, *J* pr [2] 40, 437) Silky needles

Thioacetyl derivative $C_{10}H_7NHCSCH_3$, [96°] (B & T), [111°] (J) Formed by heating (a) naphthyl acetamidine with CS_2 for several hours at 100° (Berntsen & Trompeter, *B* 11, 1760) Formed also by heating acetyl (a) naphthylamine with P_2S_5 (Jacobsen, *B* 20, 1897) White tables Gives $C_{10}H_7NHCH_2CH_2$ on reduction Oxidised by K_2FeCy_6 to ethenyl amido naphthyl mercaptan $C_{10}H_7<N>C<S>COH$

Benzoyl derivative $C_{10}H_7NHBz$ [156°] (W), [162°] (Kühn, *B* 18, 1477), [160°] (Hofmann, *B* 20, 1798) Colourless needles, v sol dilute alcohol, sl sol absolute alcohol and water (Ebell, *B* 7, 1317, 8, 562, Worms, *B* 15, 1814) Yields on nitration $C_{10}H_7(NO_2)NHBz$ [224°] PCl_5 converts it into $C_{10}H_7NCCl_2CH_3$, [60°] (Just, *B* 19, 984)

Thiobenzoyl derivative $C_{10}H_7NHSC_6H_5$, [148°] Formed from the benzoyl derivative by heating with P_2S_5 , or from (a) naphthyl acetamidine by heating with CS_2 (B & T, J) Yellowish needles or plates. Oxidised by K_2FeCy_6 to benzenyl amido naphthyl mercaptan

(a) Naphthylamine ar-tetrahydride $C_{10}H_{11}NH_2$, *ts.* $CH_2CH_2C(NH_2)CH$ CH_2CH_2COH — CH (275°) at 713 mm SG d_{20}^{20} 1.0625 Formed by the action of sodium on a solution of (a) naphthylamine in isomyl alcohol (Bamberger, *B* 20, 2916, 21, 1789) Thick colourless oil, sl sol water, v sol alcohol and ether, insol $NaOHAq$ Less basic than the (β)-isomeride Reduces Ag from hot solutions, but does not reduce Fehling's solution Readily diazotised Aqueous $NaNO_2$ (1 mol) acting on its hydrochloride (1 mol) at 0° forms $C_{10}H_{11}N_2C_6H_5NH_2$ (Bamberger & Lengfeld, *B* 23, 1134) Yields dyes with diazo-compounds $KMnO_4$ oxidises it to adipic acid

Salts — B^+HCl^- dimetric plates, v sol water and alcohol — $B^+HClHgCl_2^-$ flat white plates, sl sol cold, v sol hot water — $B^+H_2SO_4^-$ 1aq — *Picrate* yellow needles

Acetyl derivative $C_{10}H_{11}NHAO$ [158°] Needles, v sol ether, chloroform, and alcohol

(a)-Naphthylamine ac tetrahydride

$C_6H_5<CH(NH_2)CH_2$ — CH_2 (246.5°) at 714 mm. Formed by adding a 10 p.c solution of $CuSO_4$ slowly at 100° to a solution of amido naphthylhydrazine tetrahydride derived from (1,4') naphthylene diamine (Bamberger & Bammann, *B* 22, 968) Liquid, sol cold water, v sol hot water, alcohol, and ether Unlike its 'aromatic' isomeride it turns red litmus blue It cannot be diazotised Does not yield dyes with diazo-compounds $FeCl_3$ has no effect in the cold, but on warming produces a reddish-brown colour $K_2Cr_2O_7$ and H_2SO_4 give no colour. $KMnO_4$ oxidises it, in cold dilute solution, to o-carboxy β phenyl-propionic acid [166°] — B^+HCl^- needles, v e. sol water — $B^+H_2PtCl_6^-$ 2aq. [190°] Orange prisms, sol water — $B^+HNO_3^-$ [139°] Needles, v e sol water — $B^+H_2CO_3^-$ needles — *Picrate* needles, sol hot water.

Acetyl derivative $C_{10}H_{11}NHAO$, [146°] Prisms or needles, sl sol cold water

(β) Naphthylamine $C_{10}H_7N$ α
 $C_{10}H_7 \begin{smallmatrix} \text{CH} \text{CH} \text{NH}_2 \\ \text{CH} \text{CH} \end{smallmatrix}$ [112°] (294°) (Liebemann
 a. Jacobson, *A* 211, 41)

Formation—1 From the acetyl derivative of (α) naphthylamine by successive bromination, nitration, elimination of Ac, removal of NH_2 by the diazo reaction, and reduction of the resulting bromo nitro naphthalene by tin and HCl_{aq} (Liebemann & Scheiding, *A* 183, 258)—2 By heating (β)-naphthol with ammonia zinc chloride at 200°, di (β) naphthylamine being also formed (Merz & Weith, *B* 13, 1300)—3 By passing dry ammonia over strongly heated (β)-naphthol (Graebe, *B* 13, 1850)—4 By heating (β) naphthol with four times its weight of ammoniacal $CaCl_2$ (prepared by passing NH_3 over ordinary granulated $CaCl_2$ containing about 18 p.c. of H_2O) for 8 hours at 270°–280°, the yield being 80 p.c. on the naphthol, together with 12 p.c. of (β)-dinaphthylamine (Benz, *B* 16, 8)—5 In small quantity, together with a large quantity of dinaphthylamine, by heating (β)-naphthol with ammoniacal $ZnCl_2$ (*B*)

Preparation—1 By passing NH_3 under pressure into (β) naphthol at 150°–160°—2 By heating (β) naphthol (10 pts) with $NaOH$ (4 pts) and NH_4Cl (4 pts) (*G P* 14,612 [1880])

Properties—White plates (from water) with out odour Volatile with steam Gives no colouration with $FeCl_3$, chromic acid, or bleaching powder Its alcoholic solution is not coloured by nitrous acid and HCl

Reactions—1 Heated with PbO it gives azo naphthalene (Volodkevitch, *Bl* [2] 45, 178)—2 $SiCl_4$ forms $SiCl_2(NHC_{10}H_7)_2$ (Horden, *C J* 51, 40) A benzene solution of $SiCl_4$ forms $Si(NH C_{10}H_7)_4$ (Reynolds, *C J* 55, 481)—3 With *paraldehyde*, *acetone*, and HCl it gives dimethyl (β) naphthoquinoline (Reed, *J pr* [2] 35, 298)—4 With *methylal*, *acetone*, and HCl it forms methyl (β)-naphthoquinoline, (β) naphthoacridine and a base, $C_{10}H_7N_2$, which is probably methyl amido naphthyl naphthoquinoline dihydride $C_{10}H_7 \begin{smallmatrix} \text{CMe} \text{CH} \\ \text{NH} \text{CH} \text{C}_6\text{H}_4\text{NH}_2 \end{smallmatrix}$ This base [203°

uncor] forms the following derivatives $B''C_6H_4(NO_2)_2OH$ — $B''Et_2I_2$ — $C_6H_5N_2O$, [α 238°] (*R*)—5 *Cyanuric chloride* forms $C_6N_2Cl_2(NHC_{10}H_7)_2$ [154°], $C_6N_2Cl_2(NHC_{10}H_7)_2$ [278°], and $C_6N_2(NHC_{10}H_7)_3$ [209°] (Fries, *B* 19, 2056)—6 *Benzonic aldehyde* in hot alcoholic solution forms $C_{10}H_7CHNC_6H_5$ [103°] which may be reduced by sodium amalgam to $C_{10}H_7CH_2NHC_6H_5$ [68°] which forms a nitros amine $C_{10}H_7CH_2N(NO)C_6H_5$ [112°] (Claisen, *A* 237, 272, Kohler, *A* 241, 360)—7 *Benzil* at 215° forms $C_{10}H_7CH(OH)C(NC_{10}H_7)_2$ C_6H_5 crystallising from alcohol in yellow prisms [130°] (Voigt, *J pr* [2] 34, 22)—8 The compound of *glyoxal* with $KHSO_4$ yields $C_{10}H_7NSO_4K$ crystallising in white plates (Hinsberg, *B* 21, 110)—9 *Malic acid* reacts on heating, forming the compounds $C_{10}H_7O(CO NHC_6H_5)_2$ [263°] and $C_{10}H_7N \begin{smallmatrix} \text{CO} \text{CH} \text{OH} \\ \text{CO} \text{CH}_2 \end{smallmatrix}$ [193°] (Bischoff, *B* 23, 2046)—10 *Pyruvic acid* and *benzoic aldehyde* form $N C_{10}H_7 \begin{smallmatrix} \text{C} \text{CO} \text{H} \\ \text{CPh} \text{CH} \end{smallmatrix}$ (Doebner, *A* 249, 109)

11. *Quinone chlorimide* acts on an alcoholic solution forming a eurdiodine of the formula

$C_{10}H_7 \begin{smallmatrix} \text{N} \\ \text{C}_6\text{H}_4\text{NH}_2 \end{smallmatrix}$ (Nietzki & Otto, *B* 21, 1598)—12 *Chloro-acetic acid* forms $C_{10}H_7NHCH_2CO NHC_6H_5$ [170°] (Cosiner, *B* 14, 60)

Salts— $B'HCl$ colourless plates, v e sol water and alcohol, sl sol HCl_{aq} — $B''H_2PtCl_6$ yellow plates, sol water— $B''H_2SO_4$ plates, m sol cold water— $B'HNO_3$ colourless plates, sl sol cold water—*Picrate* [195°] Long yellow needles, v sol alcohol—*Citrate* $B''C_6H_5O_7$ [89°] (Hecht, *B* 19, 2616)—*Citraconate* [173°] Yellow needles (from acetone) (Morawski & Glaser, *M* 9, 285)

Compounds with metallic salts $B''CuSO_4$ (Lachovitch, *M* 9, 516) Forms also a compound with mercuric chloride

Formyl derivative $C_{10}H_7NHCHO$ [120°] (*O*, Tobias, *B* 15, 2447) [129°] (*L* & *J*) Formed by heating (β) naphthylamine with formic ether and alcohol (Cosiner, *B* 14, 58), or with formic acid (Liebemann & Jacobson, *A* 211, 42) Small plates, sl sol hot water

Acetyl derivative $C_{10}H_7NHAc$ [132°] Long needles or plates, sol hot water (Merz & Weith, *B* 13, 1300, 14, 2343) On heating with sulphur it yields $C_{10}H_7 \begin{smallmatrix} \text{N} \\ \text{C} \text{C} \end{smallmatrix} \begin{smallmatrix} \text{N} \\ \text{S} \end{smallmatrix}$ $C_{10}H_7$

crystallising in yellow plates, insol most solvents sol nitrobenzene (Hofmann, *B* 20, 1804) Bromine in $HOAc$ forms $C_{10}H_7BrNHAc$ [140°] (Lellmann & Schmidt, *B* 20, 3154)

Thio acetyl derivative $C_{10}H_7NS$ v e $C_{10}H_7NHCSCH_3$ [146°] Formed by heating the acetyl derivative with P_2S_5 (Jacobson, *B* 21, 2627) Needles or plates (from alcohol) Oxidised by K_2FeCy_4 to ethenyl amido naphthyl mercaptan $C_{10}H_7 \begin{smallmatrix} \text{N} \\ \text{S} \end{smallmatrix} \text{CMe}$ [81°]

Valeryl derivative $C_{10}H_7NHCOOC_4H_9$ [138.5°] (Bamberger & Muller, *B* 21, 1112)

Benzoyl derivative $C_{10}H_7NHBz$ [143°] (Cosiner, *B* 14, 58), [157°] (Hofmann, *B* 20, 1803) Minute needles, sol ether, benzene, and hot alcohol PCl_5 converts it into the compound $C_{10}H_7NCClC_6H_5$ [68°] (Just, *B* 19, 983)

(β) Naphthylamine *ar*-tetrahydride $C_{10}H_{11}N$ $\begin{smallmatrix} \text{CH}_2\text{CH}_2\text{CCHCNH}_2 \\ \text{CH}_2\text{CH}_2\text{CCHCNH}_2 \end{smallmatrix}$ [38°] (276°) at 713mm

Formed to the extent of 3 or 4 p.c. in the preparation of the alicyclic isomeride by reduction of (β) naphthylamine (Bamberger & Kitschelt, *B* 23, 882) Neutral in reaction Its salts are acid in reaction May be diazotised Yields adipic acid on oxidation

(β) Naphthylamine *ac* tetrahydride

$C_6H_5 \begin{smallmatrix} \text{CH}_2\text{CHNH}_2 \\ \text{CH}_2\text{CH}_2 \end{smallmatrix}$ (162°) at 36 mm, (249.5 cor) at 710 mm SG 1.031 Formed by reducing (β) naphthylamine with sodium and isoamyl alcohol (Bamberger & Muller, *B* 20, 2916, *B* 21, 850, 1115, Bamberger & Kitschelt, *B* 23, 877) Purified by dissolving in ligroun and ppg by CO_2 The carbonate is then dissolved in acetic acid, which leaves a brown oil Colourless liquid, smelling like piperidine, sl sol cold, m sol hot water, v sol alcohol and ether Powerful base with alkaline reaction, displacing ammonia from its salts Absorbs CO_2 from the air Possesses no reducing power Carbon disulphide at 0° forms tetrahydronaphthyl amine

nium tetrahydronaphthyl thio carbamate $C_{10}H_{11}NH_2$, 8 OS $NHCO_2H$, [142°] Phenyl cyanate forms $PhNH.CO.NH.C_{10}H_{11}$ [166 5°] Does not yield colouring matters with diazo compounds Powerful midratic poison $KMnO_4$ oxidises it to o-carboxy phenyl-propionic acid and phthalic acid Benzoz aldehyde forms $C_{10}H_7CH.NCO_2H$, [52°] $HBr.Aq$ at 160° forms naphthalene

Salts— $B'H_2CO_3$, and $B'H_2CO_3$, are white crystalline pps, giving off CO_2 in the air— $B'HCl$ [237°] Plates, v e sol water and alcohol— $B'H_2PtCl_6$ — $B'HAuCl_4$ — $B'HClHgCl_2$ (?) [241°] Prisms, sol hot water— $B'H_2SO_4$ — $B'H_2CrO_4$ — $B'HNO_3$, [212°] White satiny tables, sl sol cold water, v sol conc HNO_3 , At 220° it dissociates with explosive violence— $B'HNO_3$, [c 160°] Needles (from water), v e sol water, insol ether Not decomposed by boiling water— $B'HOAc$ [156°] Thick mono clinic crystals, v sol alcohol, sl sol ether

Acetyl derivative $C_{10}H_{11}NHAc$ [107°] Prisms, v e sol chloroform and benzene, v sol hot water, m sol ether, insol petroleum ether Not attacked by bromine in the cold

Benzoyl derivative $C_{10}H_{11}NHBz$ [151°] Needles, v sl sol water, v sol benzene

Di-(a)-naphthylamine $C_{20}H_{19}N$ e ($C_{10}H_9$) $_2NH$ [111° uncor] (L), [113°] (G a V) (318°) at 15 mm

Formation—1 By heating (a)-naphthylamine hydrochloride with (a) naphthylamine (Girard a Vogt, *Bl* [2] 19, 68)—2 A product in the preparation of methyl (a)-naphthylamine from (a)-naphthylamine and $MeCl$ (Landshoff, *B* 11, 638)—3 By heating (a)-naphthol with four times its weight of ammoniacal $ZnCl_2$ to 260°, the yield being about 60 p c (Benz, *B* 16, 15)—4 By heating a mixture of (a)-naphthylamine, (a)-naphthol, and $CaCl_2$ to 260°, the yield being 22 p c (B)—5 In small quantities by heating (a)-naphthylamine with $CaCl_2$ or $ZnCl_2$

Properties—Dimetric leaflets (from alcohol), m sol alcohol, v e sol ether $FeCl_3$ gives a green pp in its alcoholic solution

Picric acid compound
 $NH(C_{10}H_7)_2.2C_6H_3(NO_2)_3OH$ [169°] Small glistening black needles

Acetyl derivative $NAc(C_{10}H_7)_2$ [217°]

Nitrosamine $(C_{10}H_7)_2NNO$ [262°] Formed by adding rather more than the theoretical quantity of powdered $NaNO_2$ to a solution of di-naphthylamine in $HOAc$ (L, Wacker, *A* 243, 800) Yellow crystalline powder, v sl sol alcohol and $HOAc$ On adding alcoholic HCl to its ethereal solution it is changed to the isomeric nitroso dinaphthyl amine $C_{10}H_7(NO)NH.C_{10}H_7$, or $C_{10}H_7.NH(C_{10}H_7)O$ [169°] (Fischer a Hepp, *B* 20, 1248)

(aβ) Di-naphthyl amine $NH(C_{10}H_7)_2$ [111°]

Formed by heating a mixture of (β)-naphthol and (a)-naphthylamine with $CaCl_2$ or $ZnCl_2$

Preparation—A mixture of (β)-naphthol (100 pts), (a)-naphthylamine (100 pts), and $CaCl_2$ (200 pts), is heated for 8 hours to 280°, the yield is 70 pts (Benz, *B* 16, 17). Long colourless prisms Soluble in benzene, alcohol, and ether, sparingly soluble in petroleum-ether

Picric acid compound
 $NH(C_{10}H_7)_2.2C_6H_3(NO_2)_3OH$ [178°], small dark brown needles

Acetyl derivative $NAc(C_{10}H_7)_2$, [125°].
Di-(β)-naphthyl amine $NH(C_{10}H_7)_2$, [171°]. (471°) (Ris, *B* 20, 2619)

Formation—1 Together with (β) naphthylamine by heating (β) naphthol with ammonia zinc chloride at 200° (Merz a Weith, *B* 13, 1300)—2 In small quantity by heating (β)-naphthol with ammoniacal $CaCl_2$ —3 By heating (β)-naphthylamine with $CaCl_2$ or $ZnCl_2$

Preparation—1 By heating a mixture of (β)-naphthylamine (100 pts), (β) naphthol (100 pts), and damp $CaCl_2$ (200 pts) for 8 hours at 275°, the yield being good (130 pts) (Benz, *B* 16, 9)—2 By passing HCl into fused (β)-naphthylamine at 180°, the yield being quantitative (Klopsch, *B* 18, 1585)

Properties—White silvery leaflets, sol benzene and $HOAc$, sl sol hot alcohol Its solutions exhibit blue fluorescence

Reactions—1 Conc $HCl.Aq$ at 150° has no action, but at 200° it forms (β) naphthol and (β) naphthylamine (Ris, *B* 19, 2016)—2 Ammoniacal $ZnCl_2$ and NH_4Cl at 370° forms (β)-naphthylamine $(C_{10}H_7)_2NH + NH_3 = 2C_{10}H_7NH_2$ 3 Bromine in $HOAc$ forms a tetra bromo derivative $C_{20}H_{17}Br_4N$ [246°] Bromine and $AlBr_3$ form $C_{20}H_{17}Br_4N$ [over 300°] (Ris, *B* 20, 2621)—4 S_2Cl_2 , acting on its benzene solution at 35°, forms two isomeric imido di-naphthyl disulphides $NH(C_{10}H_7)_2S_2$, one crystallising from benzene in plates [205°], and the other in needles [220°] (Kym, *B* 21, 2807)—5 Phosgene acts in the cold, forming $(C_{10}H_7)_2NCOCl$ [173°] (Kuhn a Landau, *B* 23, 511, cf Kym, *B* 23, 427). This chloro formyl derivative is v sl sol cold alcohol, v sol benzene, and reconverted into di-(β) naphthylamine by alcoholic potash

Salts— $B'HCl$, white crystalline pp— $B'2C_6H_3(NO_2)_3OH$ [165°] Brown needles

Acetyl derivative $(C_{10}H_7)_2NAc$ [115°] Small colourless needles

Benzoyl derivative $(C_{10}H_7)_2Nbz$ [157°] Large needles (Klopsch, *B* 18, 1585)

Nitrosamine $(C_{10}H_7)_2NNO$ [140°] Needles, m sol alcohol and ether, v sol benzene (Ris, *B* 20, 2622)

References—BROMO-, BROMO NITRO, CHLORO, IODO, IODO NITRO, and NITRO NAPHTHYLAMINE

(a) -NAPHTHYLAMINE (a) -SULPHONIC ACID $C_{10}H_7NSO_3$, e $C_{10}H_7(NH_2)(SO_3H)$ [1 4] *Naphthionic acid* *S* 022 at 15°

Formation—1 By boiling nitro naphthalene (1 pt) with alcohol (5 pts) and aqueous ammonium sulphite (5 pts of SG 1.24), and maintaining an alkaline reaction by frequent addition of ammonium carbonate (Piria, *A Ch* [3] 31, 217, *A* 78, 81)—2 By heating the acid sulphate of (a)-naphthylamine (Neville a Winther, *C J* 37, 632, Witt, *B* 19, 55)—3 By heating (a) naphthylamine (1 pt with H_2SO_4 (4 pts) at 120° for an hour (Schaal a Schmidt, *B* 7, 1868, Erdmann, *A* 247, 313)—4 By heating (a) naphthylamine with $KHSO_4$ at 230° (Eischoff, *B* 23, 1913)—5 By reducing the corresponding nitro-naphthalene sulphonic acid by ammonium sulphide (Cleve, *B* 28, 961)

Properties—Small needles (containing $\frac{1}{2}eq$)

NAPHTHYLAMINE SULPHONIC ACID.

(from hot water), blackened by heat without melting. \forall sol alcohol, almost insol water. Not affected by boiling aqueous acids or alkalis. Dilute solutions of its salts exhibit violet fluorescence. The azo-colouring matters (e.g. Congo red), formed by its combination with diazo-compounds give on reduction *o*-naphthylene-diamine sulphonic acid, proving that they are *ortho*-azo-compounds, and hence that the HSO_3 group occupies the *para*-position to the NH_2 (Witt, *B* 19, 1719).

Reactions—1 Split up by water at 160° into naphthylamine and H_2SO_4 (N a W)—2 *Chromic acid mixture* forms a brown resinous product—3 *Benzonic aldehyde*, acting on its sodium salt forms $\text{C}_6\text{H}_5\text{OH}$, $\text{C}_6\text{H}_5\text{SO}_3\text{Na}$, crystallising in yellow plates, \forall sol cold water (Erdmann, *A* 247, 325)—4 *Succinic acid* heated with its K salt at 170° forms $\text{C}_6\text{H}_5\text{C}_2\text{O}_2\text{N}$, $\text{C}_6\text{H}_5\text{SO}_3\text{K}$, crystallising from water or dilute alcohol in small needles (containing 2aq) (Pellizzari, *A* 248, 157)—5 *Phthalic anhydride* heated with the K salt at 160° forms $\text{C}_6\text{H}_5\text{C}_2\text{O}_2\text{N}$, $\text{C}_6\text{H}_5\text{SO}_3\text{K}$, crystallising from hot water in small needles (containing 3aq) (P)—6 On displacing NH_2 by Cl, and distilling the resulting chloro naphthalene sulphonic acid with PCl_5 , there is formed (1,4)-di-chloro naphthalene.

Salts— KA' small micaceous laminae, \forall sol water and alcohol, sl sol KOH aq — NaA' 4aq monoclinic prisms— BaA' 8aq— CaA' 8aq, monoclinic crystals, \forall sol water, almost insol alcohol— MgA' 8aq monoclinic prisms— MgA' 10aq— PbA' 2aq, needles, sl sol water— ZnA' 2aq— CuA' 2aq— AgA' aq crystalline grains— $\text{AgA}'\text{N}_2\text{H}_5$ 2aq.

Amide $\text{C}_{10}\text{H}_7(\text{NH}_2)\text{SO}_2\text{NH}_2$ [206°] Formed by reducing the amide of (1,4) nitro naphthalene sulphonic acid with HI in HOAc and P (Cleve, *B* 23, 361). Needles (from alcohol)— B'HCl colourless needles, \forall sl sol water.

Acetyl derivative of the amide $\text{C}_{10}\text{H}_7(\text{NHAc})\text{SO}_2\text{NH}_2$ [241°] Small needles (a) Naphthylamine (α') sulphonic acid $\text{C}_{10}\text{H}_7(\text{NH}_2)(\text{SO}_3\text{H})$ [14°] S 1064 at 15° .

Formation—1 By reducing (a) nitro naphthalene sulphonic acid by ammonium sulphide (Laurent, Cleve, *Bl* [2] 24, 511)—2 Together with naphthionic acid by adding (a)-naphthylamine hydrochloride (1 pt) at 0° to (2 pts of) fuming H_2SO_4 (containing 25 p.c. additional SO_3) (Witt, *B* 19, 578, Mauzelius, *B* 20, 3401). 3 Together with a small quantity of the (1,4) acid from the acetyl derivative of (a) naphthylamine and fuming H_2SO_4 (Erdmann, *A* 247, 315, G Schultz, *B* 20, 3161, cf Lange, *B* 20, 2940).

Properties—Needles. Solutions of its salts exhibit green fluorescence, and reduce AgNO_3 on warming. Auric chloride, FeCl_3 , and CuSO_4 colour its solution red, and suffer reduction. By boiling its diazo-compound with HCl a chloro naphthalene sulphonic acid is obtained, the K salt of which on distillation with PCl_5 yields (γ)-di-chloro naphthalene [107°] (M). Benzoic aldehyde, acting on its sodium salt, forms $\text{C}_6\text{H}_5\text{CH}_2\text{N}$, $\text{C}_6\text{H}_5\text{SO}_3\text{Na}$, crystallising in pearly plates, decomposed by boiling water (Erdmann, *A* 247, 826).

Salts— KA' aq needles or prisms, \forall e sol. water— NaA' aq plates (O) or needles (M).—

NaA' 5aq plates (W)— BaA' 8aq (W)— BaA' 6aq (C, M)— CaA' 6aq (W) pearly plates, \forall e sol water— CaA' 9aq (C, M) tables, \forall sol hot water— MgA' 8aq plates, \forall sol water— PbA' 4aq 'nodules'— ZnA' 9aq needles, m sol cold water— AgA' .

Amide [260°] Tables. Yields an acetyl derivative [232°] and a diacetyl derivative [200°] (Ekborn, *B* 23, 1119)— B'HCl — $\text{B'H}_2\text{SO}_4$.

(a) Naphthylamine *para*-sulphonic acid $\text{C}_{10}\text{H}_7(\text{NH}_2)(\text{SO}_3\text{H})$ [11°] Naphthylamine sulphonic acid S 42 at 100° , 0207 at 21° . Formed, together with the (1,4') acid from naphthalene (a) sulphonic acid by nitration and reduction (Mensching, Schillkopf's *Aniline Co*, G P 40,571, G Schultz, *B* 20, 3162). Formed also, together with the (1,4') acid, from (a)-nitro-naphthalene by sulphonation and reduction (Cleve, *B* 20, 1535). White needles (from water) or tufts of needles (from HOAc). Its sodium salt is less soluble than that of the preceding isomeride. FeCl_3 colours its cold aqueous solution violet. The diazo-compound treated with PCl_5 gives $\text{C}_{10}\text{H}_7\text{ClSO}_3$, crystallising in yellow needles [175°]— NaA' S 267 at 100° , 113 at 24° — KA' Plates S 149 at 100° , 356 at 19° (Erdmann, *A* 247, 306).

Anhydride $\text{C}_{10}\text{H}_6\langle\text{SO}_2\text{NH}\rangle$ [167°] Small crystals, \forall sl sol water (Cleve).

(a) Naphthylamine (δ) sulphonic acid $\text{C}_{10}\text{H}_7(\text{NH}_2)(\text{SO}_3\text{H})$ [$12'$ or $3'$] S 2 at 15° , 7 at 100° . Obtained by heating (a) naphthylamine (1 pt) with conc H_2SO_4 (5 pts) at 125° – 130° for 8 hours until the naphthionic acid at first formed has disappeared (Hirsch, *B* 21, 2370). Plates, sl sol water and alcohol, insol ether and benzene. The K salt is crystalline, and sl sol cold water. Gives, on treatment with HNO_3 and HNO_2 , di-nitro naphthol sulphonic acid isomeride with naphthol yellow S.

(a) Naphthylamine (β' -sulphonic acid $\text{C}_{10}\text{H}_7(\text{NH}_2)(\text{SO}_3\text{H})$ [13°] Formed, together with its (γ) and (δ) isomerides, from naphthalene (β)-sulphonic acid by successive nitration and reduction (Cleve, *Bl* [2] 26, 447, *B* 20, 1535). Slender needles (containing 2aq) or anhydrous tables, sl sol cold, \forall sol hot, water— NaA' 4aq thin tables— KA' aq needles— MgA' 10aq rhombohedra— BaA' aq flat needles, m sol water— CaA' 7aq rhombohedra.

Anhydride $\text{C}_{10}\text{H}_6\langle\text{NH}\text{SO}_2\rangle$ [180°] Formed by treating the chloride of (β') nitro naphthalene (β) sulphonic acid with HI and HOAc . Yellow needles, \forall sl sol HOAc .

(a) Naphthylamine (γ) sulphonic acid $\text{C}_{10}\text{H}_7(\text{NH}_2)(\text{SO}_3\text{H})$ [13°] Formed, with isomerides, from naphthalene (β) sulphonic acid by nitration and reduction (Cleve, *B* 19, 2181, 21, 3271). Small needles, sl sol water. Gives rise to a di-chloro-naphthalene [61°].

Salts— NaA' scales, \forall sol water— BaA' aq thin plates, \forall sol water— PbA' aq prisms— AgA' aq minute needles.

Amide $\text{C}_{10}\text{H}_7(\text{NH}_2)\text{SO}_2\text{NH}_2$ aq. [181°]. Needles— B'HCl prisms, sl sol cold water. With potassium cyanate it forms $\text{NH}_2\text{CO NH C}_6\text{H}_5\text{SO}_3\text{NH CO NH}_2$ [273°].

Acetyl derivative of the amide
 $C_{10}H_7(NHAc)SO_2NH_2$ [221°].

Anhydride $C_{10}H_7\langle\begin{smallmatrix} NH \\ SO_2 \end{smallmatrix}\rangle$ [124°] Formed from (γ)-nitro naphthalene (β) sulphonic chloride, HOAc, and HI (Cleve, *B* 20, 1536) Lemon-yellow needles, v sl sol HOAc and alcohol.

(α)-Naphthylamine (θ)-sulphonic acid
 $C_{10}H_7(NH_2)(SO_3H)$ [1 2°] Formed, with isomerides, from naphthalene (β) sulphonic acid by nitration and reduction (Cleve, *Bl* [2] 29, 415, *B* 21, 3264) Crystals (containing aq) Gives rise to di chloro-naphthalene [61°] Its alcoholic solution yields with nitrous acid a deep violet dye $SO_3H.C_{10}H_7.N_2.C_{10}H_7(NH_2)SO_3H.2\frac{1}{2}aq$

Salts —NaA' 3aq thin needles, v sol water —CaA' 2aq powder, v sol water, turning red in air —BaA' flat needles —ZnA' 4aq needles
Amide $C_{10}H_7(NH_2)SO_2NH_2$ [181°] Needles. B'HCl aq —B'HI aq Yields the urea derivative $NH_2.CO.NH.C_{10}H_7.SO_2.NH.CO.NH_2$ [225°]

Acetyl derivative of the amide
 $C_{10}H_7(NHAc)SO_2NH_2$ [213°]

Anhydride $C_{10}H_7\langle\begin{smallmatrix} NH \\ SO_2 \end{smallmatrix}\rangle$ [173°] Formed from (θ) nitro naphthalene (β) sulphonic chloride, HOAc, and HI (Cleve, *B* 20, 1536) Yellow needles, sol boiling Ac_2O

(β) Naphthylamine 'α'-sulphonic acid
 $C_{10}H_7(NH_2)(SO_3H)$ [2 1°] 'Badische acid' S 599 (Forsling)

Formation —1 By heating (β) naphthylamine (1 pt) with conc H_2SO_4 (3 pts) at 100°–105° for 5 or 6 hours there is obtained a mixture of the 'α', 'β', (γ), and (δ) sulphonic acids of (β) naphthylamine in the proportion of about 50 p.c. of the 'α' acid, 40 p.c. of the 'β' acid, 5 p.c. of the (γ) acid, and 5 p.c. of the (δ) acid (Green, *C* J 55, 35, cf Badische Anilinfabrik, *G* P 14,612, 20,760, Dahl, *G* P 29,084, 32,271, 32,276) The same mixture heated at 120° gives a greater quantity of the 'β' acid, and less of the 'α' acid Fuming sulphuric acid (20 p.c. SO_3 extra) at 70°–80° gives 30 p.c. of the 'α' and 70 p.c. of the (γ)-acid (β) Naphthylamine sulphate, shaken with conc H_2SO_4 for three days in the cold, yields a similar mixture (Dahl) but on heating (β) naphthylamine with conc H_2SO_4 for an hour at 160° the 'β' and (δ) acid are formed in about equal quantities (Bayer a Duisberg, *B* 20, 1426, *G* Schultz, *B* 20, 1358) —2 By heating the (β) naphthol 'α' sulphonic acid (of Bayer) with ammonia in a closed vessel (Pätzinger a Duisberg, *B* 22, 396, cf Landshoff, *B* 16, 1931)

Properties —Needles or broad tables, v sl sol water, insol alcohol Solutions of its salts exhibit blue fluorescence

Reactions —1 Yields naphthalene (α) sulphonic acid when its amido-group is removed by the diazo-reaction (P a D, Nietzki a Zübelen, *B* 22, 453) —2 Yields by Sandmeyer's method a bromo-naphthalene sulphonic acid which can be converted into di bromo naphthalene [75°] (Forsling, *B* 22, 619) —3 Yields by Skraup's method (β)-naphthaquinoline sulphonic acid, which may be oxidised to $\langle\begin{smallmatrix} CO \\ C,NH \end{smallmatrix}\rangle.C_{10}H_7.SO_3H$, showing that the naphthylamine sulphonic acid is heteronuclear (Immerheiser, *B* 23, 402, 412, cf Armstrong a Wynne, *C* J. Proc 4, 108, 5,

49)—4 Heating with H_2SO_4 (8 pts) at 160° for $1\frac{1}{2}$ hours converts it into a mixture of the 'β' and (δ) isomerides (Weinberg, *B* 20, 3353) —5. Reacts with diazobenzene sulphonic acid, forming a yellow diazo amido- and not an azo-compound (Witt, *B* 21, 3483) —6 On conversion into the corresponding chloro naphthalene sulphonic acid and treatment of this with PCl_5 there is formed di-chloro naphthalene [61 5°]

Salts (Forsling, *B* 20, 2099) —NaA' plates, v. sol water, insol alcohol —KA' $\frac{3}{2}$ aq tables, v sol water — NH_4A' large prisms, v sol water —BaA' 4aq columns, v e sol water *S* 4 35 in the cold —CaA' 6aq tables. *S* 9 09 in the cold —MgA', $3\frac{1}{2}$ aq nodules, v e. sol water —ZnA' 6aq columns —PbA', needles —CuA', crystalline powder —AgA'

(β) Naphthylamine 'β' sulphonic acid
 $C_{10}H_7(NH_2)(SO_3H)$ [2 3°] 'Brönner's acid'

Formation —1 By heating the corresponding (β) naphthol sulphonic acid with ammonia under pressure (Farbfabrik vormals Brönner, *G* P 22,547), or by passing a current of ammonia over the potassium (β) naphthol sulphonate at 200°–250° (Landshoff, *B* 16, 1931, Green, *B* 22, 723) —2 Together with the 'α', (γ), and (δ) isomerides by heating (β) naphthylamine (1 pt) with conc H_2SO_4 (3 pts) at 105° (Bayer a Duisberg, *B* 20, 1426, Green, *C* J 55, 35) —3 By heating (β) naphthylamine with $KHSO_4$ at 230°, the yield being 60 p.c. of the theoretical (Bischoff, *B* 23, 1914) 4 The chief product obtained by heating (β)-naphthylamine acid sulphate at 200°

Properties —Plates or flat needles, sl sol warm water (Schultz, *B* 20, 3158) According to Forsling the laminar crystals contain aq Its solution exhibits blue fluorescence Yields, by conversion into diazo naphthalene sulphonic acid and application of Sandmeyer's reaction a chloro naphthalene sulphonic acid whence PCl_5 forms $C_{10}H_7Cl_2$ [136°]

Salts (Forsling, *B* 20, 76) — NH_4A' aq large thin plates, v sl sol water, forming a solution exhibiting violet fluorescence (Green) —NaA' 2aq flat needles *S* 2 5 at 15° —KA' aq long needles *S* 2 4 —BaA' 6aq needles *S* 22 at 15° —CaA' 6aq laminæ *S* 4 4 in the cold —MgA' 6aq laminæ or broad needles —PbA' 2aq v sl sol water —ZnA' 4aq —CuA' 4aq sl sol water —AgA' aq powder

(β) Naphthylamine (γ) sulphonic acid
 $C_{10}H_7(NH_2)(SO_3H)$ [2 4°] *Dahl's acid* *S* 077 in the cold (F). Formed by sulphonating (β) naphthylamine, and is the chief product when the sulphonation is carried out at 15° to 20° (Dahl, *G* P 32,276, Forsling, *B* 20, 2099, Claus, *J* pr [2] 89, 315) Small plates or needles, insol alcohol Its solutions show blue fluorescence May be converted into di chloro naphthalene [48°] Conc H_2SO_4 (3 pts) at 160° converts it into a mixture of the 'β' and (δ) isomerides (Weinberg, *B* 20, 3353)

Salts — NH_4A' exceedingly soluble tables KA' aq rhombohedra, v e sol water —NaA' 5aq tables, v e sol water —BaA' 2 $\frac{1}{2}$ aq nodules, v e sol water —CaA' 11aq needles *S* 9 09 in the cold —AgA' 8aq crystalline

(β)-Naphthylamine (δ) sulphonic acid
 $C_{10}H_7(NH_2)(SO_3H)$ [2 2°] (β)-Naphthylamine sulphonic acid *F. Bayer's acid* *S* 2857.

Formation.—1. Together with the 'β' isomeride, by heating (β)-naphthylamine with H_2SO_4 at 160°–170° (Bayer a Duisberg, *B* 20, 1426, 3158, Schultz, *B* 20, 1858, 8161) It is also formed when the sulphonation is conducted at temperatures between 105° and 160°, the quantity increasing with the temperature (Green, *C J* 55, 86) —2 By heating the corresponding (β)-naphthol sulphonic acid with ammonia at 200° (Weinberg, *B* 20, 2908, Erdmann, *B* 21, 637), or by heating naphthalene 'α' disulphonic acid with NaOH at 250°, and afterwards with NH_4Cl (Weinberg, *B* 20, 2906, 3353) —3 By heating the 'α' or (γ) isomeride with H_2SO_4 at 160° (B a D)

Properties.—Long silky needles (containing aq), m sol hot water nearly insol cold water On boiling with water the needles change to an almost insoluble crystalline powder With tetra-azo diphenyl it gives a yellowish red colouring matter (β-purpurn) Yields by the diazo reaction the (β) naphthol sulphonic acid of Weinberg, which by fusion with NaOH is converted into dioxynaphthalene [129°] Gives rise to (β) di chlorio naphthalene [114°]

Salts.— KA' needles, v sol water — NaA' 4aq white needles (from water) or plates (from hot 90 p c alcohol), v e sol hot water, S 14 in cold water, v sol hot spirit (90 p c) (difference from 'α' isomeride) — $\text{NH}_4\text{A}'$ small plates, m sol water — BaA' 4½aq plates, sl sol cold water (difference from (γ) isomeride) — MgA' aq white needles (B a D) — MgA' 5aq (W) — CaA' 6aq plates, with blue fluorescence S 38 at 15°

(α) Naphthylamine v sulphonic acid

$\text{C}_{10}\text{H}_7\text{NHSO}_3\text{H}$ *Thionaphthamic acid* Formed, together with the (1, 4) acid, by the action of ammonium sulphite on (α) nitro naphthalene (Piria, *A* 78, 54) The free acid, liberated from its salts, splits up at once into naphthylamine and H_2SO_4 — KA' pearly plates, v sol water, sl sol KOHAq — $\text{NH}_4\text{A}'$ plates, sl water and alcohol — BaA' 3aq plates — PbA' (OAc)

(α) Naphthylamine 'α' disulphonic acid

$\text{C}_{10}\text{H}_7\text{NS}_2\text{O}_6$, i.e. $\text{C}_{10}\text{H}_7(\text{NH}_2)(\text{SO}_3\text{H})_2$ [1 3 3'] Formed by reducing (α) nitro naphthalene 'α' disulphonic acid by ammonium sulphide (Alén, *Bn* 2, 407) Crystals, v e sol water and alcohol, insol ether and benzene Yields (α)-naphthylamine on treatment with sodium amalgam — $\text{NH}_4\text{HA}'$ 2aq (?) slender needles — KHA' 3aq needles, m sol hot water — CaA' 5aq — BaA' 4aq tables, m sol water — PbA' 4aq (?)

(α) Naphthylamine 'β' disulphonic acid

$\text{C}_{10}\text{H}_7(\text{NH}_2)(\text{SO}_3\text{H})_2$ [1 3 2'] Obtained by reducing (α) nitro naphthalene 'β' disulphonic acid (Alén) Small needles (from water), v sol water, sl sol alcohol Yields (α) naphthylamine on treatment with sodium amalgam — $\text{NH}_4\text{HA}'$ needles, m sol hot water — KHA' needles — CaA' 2aq — BaA' aq minute needles, sl sol water — PbA'

(α) Naphthylamine (δ)-disulphonic acid

$\text{C}_{10}\text{H}_7(\text{NH}_2)(\text{SO}_3\text{H})_2$ [1 1' 4] *Naphthylamine δ-sulphonic acid S* (Sohollkopf's *G P* 40,571) Formed by sulphonating (1, 1') naphthylamine sulphonic acid — NaA' 2aq long needles (Bernthsen, *B* 23, 8090)

(α) Naphthylamine (ε)-disulphonic acid

$\text{C}_{10}\text{H}_7(\text{NH}_2)(\text{SO}_3\text{H})_2$ [1 3 1'] Formed by heating naphthalene with conc H_2SO_4 at 90° and fuming

H_2SO_4 at 100°–120°, then nitrating, and reducing the product (*G P* 45,776, 46,953, Bernthsen, *B* 22, 8328) Formed also from naphthalene (αδ) disulphonic acid (corresponding to $\text{C}_{10}\text{H}_7\text{Cl}_2$ [48°]) by nitration and reduction (Ewer a Pick, *Monat scient* 1889, 604, cf Armstrong a Wynne, *C N* 54, 255) Colourless scales (containing 3aq), v e sol warm water — NaHA' 2aq needles or thin prisms, sl sol cold water — NaA' 6aq needles or prisms, v e sol water — BaA' 3aq — BaA' 4aq flat needles, v sol hot, m sol cold, water — $\text{Ba}(\text{HA}')_2$ 5aq minute needles, sl sol cold water

(α) Naphthylamine disulphonic acid

$\text{C}_{10}\text{H}_7(\text{NH}_2)(\text{SO}_3\text{H})_2$ [1 4 2'] S 7 at 20°, 5 at 100° Formed, together with two isomeric acids, by sulphonating (α) naphthylamine (1 pt) with H_2SO_4 (containing 25 p c SO_3 extra) (5 pts) at 120° The acids are separated by treatment of the lime salts with dilute alcohol, which dissolves the salts of the two isomeric acids (Dahl & Co, *G P* 41,957, Armstrong a Wynne, *C J Proc* 6, 125) Formed, together with a smaller quantity of the following acid, by treating (α)-naphthylamine 'α'-sulphonic acid (1 pt) with H_2SO_4 containing SO_3 (1½ pts) at 30° Groups of needles, insol 85 p c alcohol Its solutions and those of its salts exhibit blue fluorescence

Salts.— KA' 3aq — NaA' 3aq v sol water CaA' aq v sl sol water

(α) Naphthylamine disulphonic acid

$\text{C}_{10}\text{H}_7(\text{NH}_2)(\text{SO}_3\text{H})_2$ [1 4 3'] S 17 at 20° Formed in the preparation of the preceding, from which it may be separated by extracting the mixed calcium salts with dilute alcohol Alcohol of 90 p c extracts the salt of a third isomeride, subsequent treatment with alcohol of 85 p c extracts the present acid Needles, insol alcohol, v sol boiling alcohol of 85 p c The calcium salt is v sol water, insol alcohol of 90 p c The K and Na salts are v sol water Solutions of the acid and its salts exhibit blue fluorescence

(β) Naphthylamine 'α' disulphonic acid

$\text{C}_{10}\text{H}_7(\text{NH}_2)(\text{SO}_3\text{H})_2$ [2 3 3'] (β) *Naphthylamine R-disulphonic acid* Formed by heating the corresponding (β) naphthol R disulphonic acid with ammonia Readily yields dyes with diazo salts

(β) Naphthylamine (γ) disulphonic acid

$\text{C}_{10}\text{H}_7(\text{NH}_2)(\text{SO}_3\text{H})_2$ [2 1' 3'] (β) *Naphthylamine G disulphonic acid* Formed by heating the corresponding (β) naphthol G disulphonic acid with ammonia Formed also by heating (β) naphthylamine sulphate (10 kilos) with H_2SO_4 (30 kilos containing 25 p c SO_3 extra) at 110°–140° (Gans a Co, *G P* 35,019) v sol water, m sol alcohol Does not react with diazo-compounds (G Schultz, *B* 21, 3487) The salts are v sol water

(β) Naphthylamine disulphonic acid

$\text{C}_{10}\text{H}_7(\text{NH}_2)(\text{SO}_3\text{H})_2$ [2 1 3'] Formed, together with the (2, 1', 3') acid, from (2, 3')-naphthylamine sulphonic acid and H_2SO_4 (with 20 p c SO_3) at 20° (Armstrong a Wynne, *C J Proc* 6, 130) Needles Yields $\text{C}_{10}\text{H}_7\text{Cl}_2$ [92°]

(β) Naphthylamine disulphonic acid

$\text{C}_{10}\text{H}_7(\text{NH}_2)(\text{SO}_3\text{H})_2$ [2 3' 2'] Formed by heating (Brønner's) (β)-naphthylamine 'β'-sulphonic acid at 160° to remove water of crystallisation, adding fuming H_2SO_4 (4 pts) and heating at 110° (Forsling, *B* 21, 3495) Possibly identical with the preceding acid. White needles, v. e.

sol water, sl. sol alcohol Its dilute aqueous solution fluoresces blue It gives rise to a tri-chloro-naphthalene [91°]

Salts— K_2A'' 2aq large needles, v sol water— KHA'' aq needles, sl sol cold water— $NaHA''$ 2aq— Na_2A'' long needles— $(NH_4)_2A''$ aq monoclinic crystals— NH_4HA'' — CaA'' plates

(β)-Naphthylamine disulphonic acid $C_{10}H_7(NH_2)(SO_3H)_2$ [2 1 4'] Formed, together with a greater quantity of the (2, 2', 4')-isomeride from (2, 4') naphthylamine sulphonic acid and H_2SO_4 (with 20 p.c. SO_3) at 20° (A & W) Yields $C_{10}H_7Cl_2$ [78°]

(β)-Naphthylamine disulphonic acid $C_{10}H_7(NH_2)(SO_3H)_2$ [2 2' 4'] Formed as above. Yields $C_{10}H_7Cl_2$ [80°]

(β)-Naphthylamine disulphonic acid $C_{10}H_7(NH_2)(SO_3H)_2$ [2 3 2'] Formed from the corresponding naphthol disulphonic acid

Di-NAPHTHYL-ANTHYLENE $C_{22}H_{12}$ \pm $C_{10}H_6$ C ||| (?) [270°] Formed by distilling di- $C_{10}H_6$ C

naphthyl-acetylene, or by distilling (β) di naphthyl-tri-chloro-ethane (1 pt) with ZnO (15 pts) (Grabowski, B 11, 302) Large leaflets (by sublimation)— $C_{22}H_{12}C_6H_4(NO_2)_2OH$ crystals (from chloroform)

NAPHTHYL ARSENIOSULFONIC ACID Described as *Naphthalene arsenic acid*, vol 1 p 322

NAPHTHYL-BENZAMIDINE $C_{17}H_{13}N$ \pm $C_{10}H_7C(NH)NHC_6H_5$ [141°] Formed by heating (α) naphthylamine hydrochloride with benzonitrile at 200° (Bornthsen a Trompeter, B 11, 1757) Tables (from alcohol), may be sublimed—B HCl white prisms— $B_2H_3PtCl_2$ yellow needles— BH_3CO_2 prisms, sl sol water—Chromate yellow pp

(α) NAPHTHYL-BENZYL AMINE $C_{17}H_{15}N$ \pm $C_{10}H_7NHCH_2Ph$ [67°] Formed from (α) naphthylamine and benzyl chloride (Froté a Tommasi, B [2] 20, 67)

(β)-Naphthyl-benzyl-amine $C_{10}H_7NHCH_2Ph$ [68°] Formed by reducing $C_{10}H_7NCHPh$ (Köhler, A 241, 358) Prisms

Nitrosamine $C_{10}H_7N(NO)CH_2Ph$ [112°] Yellow needles, sol alcohol and ether

(α)-NAPHTHYL-BENZYLIDENE-AMINE $C_{10}H_7NCHPh$ [78°] Formed from (α) naphthylamine and benzoic aldehyde or hydrobenzamide (Lachovitch, M 9, 695, cf Papasogli, A 171, 138) Yellow needles (from alcohol)

(β)-Naphthyl-benzylidene-amine [101°] Resembles the preceding in preparation and properties (Claisen, A 237, 261)

NAPHTHYL BENZYL KETONE v BENZYL NAPHTHYL KETONE

NAPHTHYL BENZYL OXIDE v BENZYL NAPHTHYL OXIDE

NAPHTHYL BROMO-METHYL KETONE $C_{12}H_9BrO$ \pm $C_{10}H_7COCH_2Br$ Obtained by adding bromine (9.5 g) to a solution of naphthyl methyl ketone (10 g) in CS_2 (Fampel a Schmidt, B 19, 2898) Pungent oil

(α)-NAPHTHYL-CARBAMATE $C_{11}H_9NO_2$ \pm $C_{10}H_7OCO NH_2$ [158°] Formed from (α) naphthol and $HCO NH_2$ (Gattermann, A 244, 48). Needles (from alcohol)

(β)-Naphthyl carbamate [187°] From (β) naphthol and $OICONH_2$ (G). Long needles, almost insol. water, v. sol alcohol and ether.

(α)-NAPHTHYL-CARBAMIC ACID *Ethyl ether* $C_{11}H_{11}NO_2$ \pm $C_{10}H_7NHCO_2Et$ [79°] Formed from (α) naphthylamine and $ClCO_2Et$ (Hofmann, B 3, 657) Needles, insol water

Isopropyl ether $C_{10}H_7NHCO_2Pr$ [78°] From (α) naphthylamine and $ClCO_2Pr$ (Spica, G 17, 168) Groups of needles, sl sol water

(β) Naphthyl-carbamic acid *Ethyl ether* $C_{10}H_7NHCO_2Et$ [73°] Formed from (β) naphthylamine and $ClCO_2Et$ (Cosiner, B 14, 60) Needles, insol hot water, v sol alcohol

Isopropyl ether $C_{10}H_7NHCO_2Pr$ [70°] Needles, sol alcohol and ether (S)

Di-(β) naphthyl carbamate *Methyl ether* $(C_{10}H_7)_2NCO_2Me$ [114°] Formed from di-(β) naphthylamine and $ClCO_2Me$ at 155° (Ris, B 20, 2620) Needles (from alcohol and ether), v sol cold alcohol and ether

(α)-NAPHTHYL-CARBAMINE $C_{11}H_{11}N$ \pm $C_{10}H_7NO$ Formed from (α) naphthylamine, chloroform, and alcoholic KOH (Liebermann, B 16, 1640) Solid, v sol alcohol

(β) Naphthyl-carbamine $C_{10}H_7NO$ [54°] Formed from (β) naphthylamine, chloroform, and alcoholic potash (Liebermann, B 16, 1640) Needles, sol alcohol, ether, and benzene

(α)-NAPHTHYL *semi* CARBAZIDE $C_{11}H_{11}N_2O$ \pm $C_{10}H_7NHNHCO NH_2$ [231°] Formed by heating (α) naphthylamine hydrochloride with urea at 140° (Pinner, B 21, 1219) Thin plates (from boiling isomyl alcohol), insol water and ether, sl sol dilute alkalis

(β)-Naphthyl-*semi*-carbazide [225°] (P), [221°] (H) Formed like its isomeride, and also by mixing equivalent quantities of (β) naphthylamine hydrochloride and potassium cyanate in aqueous solution (Pinner, B 21, 1223, Hillinghaus, B 22, 2657, Hauff, A 253, 28) Silky plates, sl sol hot water, v sol hot alcohol Reduces Fehling's solution With $HClAq$ at 140° it yields a naphthazine

DINAPHTHYL-CARBAZOLE, so called, is described as *Imido dinaphthyl*

(α) NAPHTHYL CARBINOL $C_{11}H_{13}O$ \pm $C_{10}H_7CH_2OH$ *Naphthobenzyl alcohol* [60°] (301° cor) at 715 mm Formed by warming (α) naphthyl carbinylamine hydrochloride with aqueous $NaNO_2$ (Bamberger a Lodter, B 21, 258) Long needles, v e sol ether and alcohol, v sl sol cold water Yields (α) naphthoic aldehyde on oxidation with chromic acid mixture

(β) Naphthyl-carbinol $C_{10}H_7CH_2OH$ [80 5°] Resembles the preceding in mode of preparation and properties (Bamberger, B 20, 1118)

Tri naphthyl-carbinol $C_{30}H_{21}O$ \pm $(C_{10}H_7)_3COH$ Formed from naphthalene, $C(NO_2)Cl_3$, and $AlCl_3$, the product being boiled with water (Elbs, B 16, 1275) Crystalline powder (from acetone), melting below 278°, v sol benzene, sl sol ether, almost insol alcohol

(α) NAPHTHYL-CARBINYLAMINE $C_{11}H_{11}N$ \pm $C_{10}H_7CH_2NH_2$ *Menaphthylamine Naphthobenzylamine* (292°) Formed, together with *s*-di-naphthyl ethane, by reducing the amide of thionaphthoic acid in alcoholic solution with zinc and $HClAq$ (Hofmann, B 1, 101, Bamberger a Lodter, B 21, 256) Caustic liquid, absorbing CO_2 from the air Reduced in alcoholic solution by Na to the tetrahydride— $B'HCl$ long needles, sl. sol. water— $B'H_2PtCl_6$ crystalline pp— $B'HNO_3$ prisms [148 5°].

Tetrahydride $C_{10}H_7CH_2NH_2$ (270°) at 722 mm. Obtained by reducing the nitrile of (a)-naphthoic acid $C_{10}H_7CN$ in alcoholic solution by sodium (Bamberger & Lodter, *B* 20, 1707) — $BHCl$, white needles, v sol hot water — $B'H.PtCl$, yellow needles, sl sol cold water — $B'C_6H_5(NO_2)OH$ needles, v sol hot water

(β)-Naphthyl-carbonylamine $C_{10}H_7CH_2NH_2$ [60°] Formed from the amide of thio-(β) naphthoic acid $C_{10}H_7CSNH_2$ by treating its alcoholic solution with zinc and HCl at 35° (Bamberger & Lodter, *B* 21, 1117) Prisms, sl sol cold water, v sol alcohol. It is a powerful base, ppg the hydroxides from solutions of salts of copper, zinc, and lead

Salts — $B'HCl$ [260°–270°]. Prisms, v sol alcohol and water, insol ether — $B'H.PtCl$, yellow needles — $B'C_6H_5(OH)(NO_2)$, golden-yellow needles, v sol hot water

Tetrahydride $C_{10}H_7CH_2NH_2$ (270° cor) at 729 mm Formed by adding sodium to a hot alcoholic solution of (β) naphthothionitrile (Bamberger & Boekmann, *B* 20, 1711) Yields an acetyl derivative [65°] (Bamberger & Helwig, *B* 22, 1916) Carbon disulphide reacts forming $C_{10}H_7CH_2NHCS SHNH_2$, $CH_2C_{10}H_7$, [128°], which on boiling with alcohol gives rise to $CS(NHCH_2C_{10}H_7)$, [143°] — $BHCl$ [229°] Needles, v sol water and alcohol With potassium cyanate it gives $CO(NHCH_2C_{10}H_7)$, [226°] and $CO(NH_2)(NHCH_2C_{10}H_7)$, [135°] — $B'H.PtCl$, Needles — $B'H_2CO$, white needles — $B'H_2SO$, Prisms, v sol water — $B'C_6H_5(NO_2)OH$ Yellow prisms, sl sol water

DI NAPHTHYL-TRI CHLORO-ETHANE ν TRI-CHLORO DI NAPHTHYL-ETHANE

(a) NAPHTHYL CYANAMIDE $C_{10}H_7NHCON$ [188°] Formed by heating a solution of oxy-(a) naphthyl thio urea $C_{10}H_7NHCSNH_2$ (Tiemann, *B* 22, 1940)

Di-(a) naphthyl-cyanamide $C_{10}H_7N_2$, s $C_{10}H_7N_2CN$ $C_{10}H_7$, Di-(a) naphthyl carbimide Carbo-di (a) naphthyl imide [94°] Obtained by adding HgO to a boiling solution of di (a) naphthyl thio-urea in dry benzene, the yield being c 30 p c of the theoretical (Huhn, *B* 19, 2405) Large prisms v sol benzene, sl sol cold ether and petroleum ether By heating with dilute alcohol it is converted into di (a) naphthyl-urea H_2S passed into the boiling solution in dry benzene converts it into di-(a) naphthyl thio urea Heated with CS_2 at 200° it yields (a) naphthyl thiocarbimide.

Di-(β)-naphthyl cyanamide $H_7C_{10}N_2CN$ $C_{10}H_7$, Di-(β) naphthyl-carbimide. [146°] Obtained by adding HgO to a boiling solution of di-(β) naphthyl thio urea in dry benzene, the yield being 25 p c of the theoretical (Huhn, *B* 19, 2408) White granular crystals. V sol hot benzene, sl sol ether and petroleum ether By boiling with dilute alcohol it is converted into di (a) naphthyl urea If H_2S is passed into its solution in dry boiling cumene, di-(β)-naphthyl-thio-urea is regenerated With CS_2 at 200° it yields (β)-naphthyl-thio carbimide

(a) - NAPHTHYL - CYANATE $C_{10}H_7NCO$ (270°) Formed in small quantity by heating di-naphthyl-urea with P_2O_5 , and in larger quantity by the like treatment of naphthyl carbamis ether (Hall, *Pr* 9, 366, Hofmann, *Pr* 19, 108; *O* R 47, 425). Pungent liquid. With oxy-azo-

benzene it forms $C_6H_5N_2C_6H_5COONHC_6H_5$, [149°] (Goldschmidt & Rosell, *B* 23, 492)

NAPHTHYL CYANIDE ν NITRILE OF NAPHTHOIC ACID

NAPHTHYL CYANURIC ACID ν Cyanuric acid in the article CYANIC ACIDS

TRI-NAPHTHYL-CYANURATES

($C_{10}H_7$), C_3N_3O , Formed from cyanuric chloride and sodium-naphthyl dissolved in naphthol (Otto, *B* 20, 2239) The (a)- compound decomposes between 160° and 225°, the (β)- compound begins to decompose at 220° Both are powders, sl sol hot water and alcohol, m sol benzene

NAPHTHYLENE-ACETAMIDINE ν NAPHTHYLENE ETHENYL-AMIDINE

ν NAPHTHYLENE DIAMINE $C_{10}H_7N_2$, s $C_{10}H_7(NH_2)_2$ [12] Di amido-naphthalene Amido-naphthylamine Mol w 158 [95°]

Formation — 1 By reduction of p sulpho benzene azo (β)-naphthylamine with tin and HCl (Griess, *B* 15, 2193, Witt, *B* 21, 3482) — 2 By reduction of benzene azo (β) naphthylamine, of (β) naphthalene azo (β) naphthylamine, of the dioxim of (β) naphthoquinone, of (2,1) nitro-(a) naphthylamine, or of (1,2) nitro-(β) naphthylamine (Lawson, *B* 18, 800, 2423, Leuckart, *B* 19, 174, Lellmann & Remy, *B* 19, 803, Bamberger & Schieffelin, *B* 22, 1376)

Properties — Silvery trimetric plates (from hot water), sl sol water, sol alcohol and ether Its ethereal solution quickly turns brown. $FeCl_3$ colours its solution green

Reactions — 1 Phenanthraquinone yields naphthophenanthrazine which forms yellowish white crystals giving a scarlet solution in H_2SO_4 2 Phenyl cyanate in benzene solution reacts forming $C_{10}H_7NHCO NH C_6H_5NH_2$ [335°] and ($C_6H_5NHCO NH$) $C_{10}H_7$ — 3 Phenyl thiocarbimide unites forming ($C_6H_5NHCS NH$) $C_{10}H_7$, [355°–360°] — 4 On heating with excess of oil of mustard in alcoholic solution it yields, in like manner, silky needles of $C_{10}H_7(NHCS NHCH_2C_{10}H_7)$, which decompose at 170°–200° into naphthylene thio urea and di allyl thio urea (Lellmann, *B* 19, 808) — 5 An alcoholic solution of benzil reacts forming di phenyl naphthoquinoxaline $C_{10}H_7 \begin{smallmatrix} < N & CC_6H_5 \\ & N & CO_6H_5 \end{smallmatrix}$, [148°] (Leuckart, *B* 19, 174)

6 o Aldehyde-benzene acid $CHO C_6H_4 CO_2H$ forms $C_{10}H_7 \begin{smallmatrix} < N & NH \\ & N & \end{smallmatrix} C_6H_4 CO_2H$, which decomposes at 280° (Bistreyzky, *B* 23, 1044)

Salts — $B'H_2Cl$, [90°] Prisms or plates v sol water — $B'H_2SO$, white plates, sl sol water — Picrate nearly insoluble powder

Acetyl derivative $C_{10}H_7(NHAc)_2$, [234°] White needles

Propionyl derivative $C_{10}H_7(NHC_2H_5O)_2$, [192°] Formed from the base and propionic anhydride Prisms (from alcohol), insol ether

Benzoyl derivative $C_{10}H_7(NHBz)_2$, [291°] Plates, sl sol alcohol and $HOAc$ (Hinsberg, *A* 254, 256).

ar. Tetrahydride $C_{10}H_7N_2$, s $C_{10}H_7$

$CH_2CH_2C(O(NH_2)C(NH_2))$, [84°], (220°) at $CH_2CH_2COH=CH$

81 mm Formed, together with a smaller quantity of the alicyclic isomeride, by reducing naphthylene-o-diamine with sodium (Bamberger & Schieffelin, *B* 22, 1377). Needles, v sol alcohol, ether, and hot water. Reduces $AgNO_3$. Gives

a red colour with cold aqueous FeCl_3 , KMnO_4 oxidises it to adipic acid

Salts— $\text{B}^+\text{H}_2\text{Cl}_2$ [c 260°] Tables, v sol water— B^+HNO_3 [201°] Plates, m sol water

Acetyl derivative of the tetrahydride
 $\text{C}_{10}\text{H}_8(\text{NHAc})_2$ [245°] Needles, v e. sol alcohol, sl sol ether and cold water

ac-Tetrahydride
 $\text{CH CH C OH}(\text{NH}_2) \text{CH}(\text{NH}_2)$
 $\text{CH CH C CH}_2\text{—CH}_2$ Formed as above
 (B & S) Its hydrochloride and platinochloride crystallise in needles

Naphthylene-p-diamine $\text{C}_{10}\text{H}_8(\text{NH}_2)_2$ [1 4] [120°] (G), [118°] (B & S)

Formation—1 By reduction of naphthalene-azo-(a)-naphthylamine by tin and HCl (Perkin, *C J* 18, 178, *A* 137, 359, Friedlander, *B* 22, 587).—2 By reduction of p-sulpho-benzene azo-(a)-naphthylamine with tin and HCl (Griess, *B* 16, 2192)—3 By reduction of (a) nitro-(a) naphthylamine (Liebermann, *A* 183, 238)—4 By boiling benzene-azo-(a) naphthylamine with zinc dust and water (Bamberger & Schieffelin, *B* 22, 1881)

Properties—Colourless prisms or needles, sl. sol water, sol alcohol and ether Yields (a)-naphthoquinone on oxidation with FeCl_3

Salts— $\text{B}^+\text{H}_2\text{Cl}_2$ white soluble four sided plates, nearly insol HClAq — $\text{B}^+\text{H}_2\text{SO}_4$ needles

Mono-acetyl derivative
 $\text{C}_{10}\text{H}_8(\text{NH}_2)(\text{NHAc})$ Formed by reducing the acetyl derivative of (a)-nitro-naphthylamine with tin and HCl (Liebermann)— B^+HCl long needles— $\text{B}^+\text{H}_2\text{Cr}_2\text{O}_7$,— $\text{B}^+\text{C}_6\text{H}_5(\text{NO}_2)_2\text{OH}$ yellow needles

Di-acetyl derivative $\text{C}_{10}\text{H}_8(\text{NHAc})_2$ [205°] Formed from the base or its mono-acetyl derivative and Ac_2O (Kleemann, *B* 19, 334, B & S) Needles, sl sol alcohol, nearly insol water and ether

Mono-benzoyl derivative
 $\text{C}_{10}\text{H}_8(\text{NH}_2)(\text{NHBz})$ [186°] Formed by reducing $\text{C}_{10}\text{H}_8(\text{NO}_2)(\text{NHBz})$ (Ebell, *A* 208, 326)—Needles— B^+HCl — B^+HNO_3 — $\text{B}^+\text{H}_2\text{SO}_4$

ar-Tetrahydride $\text{CH}_2\text{CH}_2\text{C}(\text{NH}_2)\text{CH}(\text{NH}_2)\text{CH}_2\text{CH}_2\text{C}(\text{NH}_2)\text{CH}(\text{NH}_2)\text{CH}_2$
 The sole product of the reduction of p naphthylene-diamine by sodium (Bamberger & Schieffelin, *B* 22, 1882) Needles, resinified on exposure to air Yields adipic acid on oxidation with KMnO_4 — $\text{B}^+\text{H}_2\text{Cl}_2$ crystalline powder

Acetyl derivative of the tetrahydride
 $\text{C}_{10}\text{H}_8(\text{NHAc})_2$ [285°] Needles, v sl sol cold water, m sol boiling alcohol Reduces ammoniacal AgNO_3 , FeCl_3 colours a solution of its hydrochloride green, changing to brown

Peri-naphthylene-diamine $\text{C}_{10}\text{H}_8(\text{NH}_2)_2$ [1 1] [67°]

Formation—1 By reduction of peri di nitro-naphthalene with iodide of phosphorus and water (De Aguiar, *B* 3, 27, 7, 307, Beilstein & Kuhlberg, *A* 169, 90, Ladenburg, *B* 9, 1651)—2 By reducing di-nitro-(a)-naphthoic acid [265°] with tin and HCl (Ekstrand, *B* 20, 1353, *J pr* [2] 38, 263)—3 By the action of ammonia on peri di oxy-naphthalene at 150°–300° (Erdmann, *A* 247, 368)

Properties—Needles (from dilute alcohol), m sol water Gives a reddish-brown colour and pp with FeCl_3 , NaNO_2 added to a solution of the sulphate ppts red needles of the azimide.

By the diazo-reaction it yields di chloro-naphthalene [84°] Benzoic aldehyde forms $\text{C}_{10}\text{H}_7\text{—}\langle\text{N}=\text{CPh}\rangle\text{—N}(\text{CH}_2\text{Ph})$ (Hinsberg, *B* 22, 861)

Phenanthraquinone does not yield an azine Oxalic ether at 100° yields $\text{C}_{10}\text{H}_8\text{N}_2\text{O}_2$, crystallising from chloroform in red needles, carbonised at 195° (Aguar)

Salts— $\text{B}^+\text{H}_2\text{Cl}_2$ [c 280°] Small plates— $\text{B}^+\text{H}_2\text{SO}_4$ — $\text{B}^+\text{H}_2\text{CO}_3$

(1,4')-Naphthylene diamine $\text{C}_{10}\text{H}_8(\text{NH}_2)_2$ [1 4'] [190°] Formed by reducing the corresponding di nitro naphthalene in alcoholic solution with tin and HCl (Erdmann, *A* 247, 361, of Zinnin, *A* 52, 362, 85, 329, Hollemann, *Z* [2] 1, 555, De Aguiar, *B* 3, 33, 7, 307) Formed also by heating the corresponding di oxy-naphthalene with ammonia at 150°–180°, and finally at 250°–300° (E) Thin white needles, which may be sublimed, sl sol cold water, v sol alcohol FeCl_3 colours its solution bluish-violet Converted by the diazo reaction into di chloro-naphthalene [107°]— $\text{B}^+\text{H}_2\text{Cl}_2$ — $\text{B}^+\text{H}_2\text{I}_2$ — $\text{B}^+\text{H}_2\text{SO}_4$ — $\text{B}^+\text{H}_2\text{C}_2\text{O}_4$

Tetrahydride $\text{CH CH—C CH}(\text{NH}_2) \text{CH}_2$
 $\text{CH C}(\text{NH}_2) \text{C CH}_2\text{—CH}_2$ [77°] (264°) at 60 mm Formed by the action of sodium on a solution of the base in isoamyl alcohol (Bamberger & Hoskyns-Abraham, *B* 22, 944) Prisms (from ether) or needles (from ligroin), sl sol hot water, v sol alcohol FeCl_3 gives a deep reddish-brown colour in its hot solution, but no colour in the cold Boiling $\text{K}_2\text{Cr}_2\text{O}_7$ gives a claret-colour Gives off NH_3 when heated This base can be separated into dextro- and laevo-rotatory varieties by crystallisation of the bitartrate, for on adding a crystal of dextro rotatory conine tartrate, the laevo-rotatory tetrahydro naphthylene diamine tartrate crystallises out, while the mother liquor deposits the dextro-rotatory compound after long standing (Bamberger, *B* 23, 291) The rotatory powers of the two hydrochlorides, $[\alpha]_D$ are –7° 30' and +8° 9' respectively

Reactions
 1 Nitrous acid at 0° forms a diazo-compound which when boiled with water yields $\text{CH CH—C CH}(\text{NH}_2) \text{CH}_2$, the di acetyl derivative of which crystallises in needles [152°] (Bamberger & Bammann, *B* 22, 960)—2 Diazoisation and reduction by SnCl_4 and HCl yields

$\text{CH CH—C CH}(\text{NH}_2) \text{CH}_2$
 the hydrazine $\text{CH C}(\text{NH}_2) \text{C—CH}_2\text{—CH}_2$

3 A dilute ethereal solution of CS_2 forms $\text{C}_{10}\text{H}_8(\text{NH}_2)_2 \text{NH}_2\text{S CS NH C}_{10}\text{H}_7\text{NH}_2$ [145°], the alicyclic amidogen entering into reaction By treatment with PbO this substance is converted into the urea $\text{CO}(\text{NH C}_{10}\text{H}_7\text{NH}_2)_2$ Boiling with alcohol yields $\text{CS}(\text{NH C}_{10}\text{H}_7\text{NH}_2)_2$ [155°]—4 A warm alcoholic solution of CS_2 forms

$\text{C}_{10}\text{H}_7\text{—}\langle\text{NH CS NH}\rangle\text{—C}_{10}\text{H}_7$ [175°], both amidogens taking part in the reaction

Salts $\text{B}^+\text{H}_2\text{Cl}_2$ trimetric prisms, $a b c = 574 \cdot 1 \cdot 906$, v sol water, v. sl. sol alcohol— $\text{B}^+\text{H}_2\text{PtCl}_4$ aq prisms— $\text{B}^+\text{H}_2\text{PtCl}_4$ crystalline solid— $\text{B}^+\text{H}_2\text{SO}_4$ 2aq triclinic prisms *Acetyl derivative* $\text{C}_{10}\text{H}_8(\text{NHAc})_2$ [262° cor] Prisms (from alcohol), sol ether, sl sol boiling water

(2,2') Naphthylene-diamine $\text{C}_{10}\text{H}_8(\text{NH}_2)_2$ [2 2'] [159°] (B. & S.); [161°] (L). Formed by heat-

ing the corresponding di oxy naphthalene [186°] of Ebert and Merz with ammoniacal CaCl_2 at 265° (Lange, *B* 21, Ref 839, Bamberger a Schieffelin, *B* 22, 1884). Plates (from water), v sol boiling water, alcohol, and ether

(2,3'-Naphthylene - diamine $\text{C}_{10}\text{H}_8(\text{NH}_2)_2$ [218°] Formed from di-oxy naphthalene [218°], by heating with ammonia and NH_4Cl at 200°-250° (Lange, *B* 21, Ref 839) Its salts are more soluble than those of the (2,2')-isomeride

m-Naphthylene diamine $\text{C}_{10}\text{H}_8(\text{NH}_2)_2$ [13] Obtained by reducing di nitro naphthalene [144°] with tin and HCl (Urban, *B* 20, 978) A solution of its hydrochloride is coloured yellow by nitrous acid — $\text{B}^{\text{H}}\text{Cl}_2$ v. e sol water, m sol alcohol, insol ether

Acetyl derivative $\text{C}_{10}\text{H}_8(\text{NHAc})_2$ [156°]. Prisms, v sol benzene

Dinaphthylene-amine $\text{C}_{20}\text{H}_{16}\text{N}_2$ s.s.

$\text{C}_{10}\text{H}_7\text{NH} > \text{NH} (?)$ [159° cor] Formed by heating (ββ) dioxy dinaphthyl with ammoniacal zinc chloride (Walder, *B* 15, 2173) White trimetric plates or needles v sol ether, benzene, or acetone, insol dilute acids

Picric acid compound

$\text{C}_{20}\text{H}_{16}\text{N}_2 \cdot 2(\text{C}_6\text{H}_3(\text{NO}_2)_3\text{OH})$ [219° cor] blue-black needles

Acetyl derivative $\text{C}_{20}\text{H}_{16}\text{N}_2\text{Ac}$ [144° uncor], fine white needles, v sol ether, less sol alcohol

Tri naphthylene diamine $\text{C}_{30}\text{H}_{24}\text{N}_6$ s.s. ($\text{C}_{10}\text{H}_7\text{N}_2$)₃ (?) Formed by heating a mixture of naphthylamine, naphthylamine hydrochloride, and nitro naphthalene in molecular proportions for 3 hours at 190°-220° (Salzmänn a Wichelhaus, *B* 9, 1107) Amorphous blue black powder (containing aq), insol water and ether, forming a red solution in warm benzene Begins to decompose at 180° — $\text{B}^{\text{H}}\text{Cl}$ amorphous violet powder

References — Bromo-, Di-chloro-, and ETHENYL-NAPHTHYLENE DIAMINE

o NAPHTHYLENE DIAMINE - 'α'- SULPHONIC ACID

$\text{C}_{10}\text{H}_7\text{N}_2\text{SO}_3$ s.s. $\text{C}_6\text{H}_4 \begin{matrix} \text{C}(\text{NH}_2) & \text{C}(\text{NH}_2) \\ & | \\ & \text{C}(\text{SO}_3\text{H})\text{CH} \end{matrix}$ Formed,

together with benzidine, by reduction of Congo-red Not isolated (Witt, *B* 19, 1719)

o-Naphthylene-diamine 'β'-sulphonic acid $\text{CH CH C}(\text{NH}_2) \text{C}(\text{NH}_2)$ Obtained by

reducing 'gold-brown,' an azo dye obtained from Brönner's (β)-naphthylamine sulphonic acid, and diazobenzene (Witt, *B* 21, 3484) Yellowish-white crystals (from alcohol or ether), v sl sol water Quickly turns brown in air K_2FeCy_4 turns its solution first brown, then yellow FeCl_3 colours its aqueous solution dirty-green Phenanthraquinone bisulphate yields, in presence of NaOAc and HOAc , naphthophenanthrazine sulphonic acid, which dissolves in H_2SO_4 with reddish-violet colour, and which is converted by potash fusion into a eurhodol, forming in H_2SO_4 a solution coloured a pure ultramarine, turned cherry-red by water Naphthylene-diamine 'α'-sulphonic acid acts in the same way, but the compound formed by phenanthraquinone dissolves in H_2SO_4 with bluish-violet colour, and the eurhodol with indigo blue colour, the sul-

phate being ppd as a crimson crystalline powder

o Naphthylene-diamine (γ)-sulphonic acid $\text{CH CH} \text{---} \text{C}(\text{NH}_2) \text{C}(\text{NH}_2)$ Formed by reduction of azo dyes prepared from (β)-naphthylamine (γ) sulphonic acid (of Dahl) (Witt, *B* 21, 8486) Plates, sl sol water (more soluble than the 'β'-isomeride) FeCl_3 colours its solution emerald green The corresponding azines are rendered violet by H_2SO_4 , becoming orange on dilution The eurhodol gives a dark violet solution in H_2SO_4 , becoming cherry-red on dilution, the eurhodol sulphate being deposited in dirty-red flakes

o Naphthylene-diamine (δ) sulphonic acid $\text{SO}_3\text{H C} \text{---} \text{CH C}(\text{NH}_2) \text{C}(\text{NH}_2)$ Obtained by

reducing azo-dyes prepared from (β)-naphthylamine (δ) sulphonic acid (W) Grey powder, more soluble in water than the 'β'-isomeride Occurs also in a gelatinous (?) hydrated condition, v e sol water It resembles the 'β'-isomeride in its reactions with FeCl_3 , with K_2FeCy_4 , and with phenanthraquinone

o Naphthylene-diamine 'α'-disulphonic acid $\text{C}_{10}\text{H}_7(\text{NH}_2)_2(\text{SO}_3\text{H})_2$ [121' or 4'3'] Obtained by reducing benzene azo (β) naphthylamine 'α'-disulphonic acid formed from (β) naphthylamine R disulphonic acid (Witt, *B* 21, 3437) — NaHA'' sandy crystalline powder, v sol water, forming a solution with green fluorescence FeCl_3 gives a green colour Yields lemon yellow sodium naphthophenanthrazine disulphonate, which forms a bluish magenta solution in H_2SO_4 The eurhodol yields a deep greenish blue solution in H_2SO_4 , becoming claret red on dilution

Naphthylene-diamine disulphonic acid $\text{C}_{10}\text{H}_7(\text{NH}_2)_2(\text{SO}_3\text{H})_2$ Formed by reducing di-nitro naphthalene disulphonic acid (Alén, *Bn* 3, 1025) — KHA'' 3aq needles, m sol hot water — $\text{Ba}(\text{HA}'')_2$ 6aq

NAPHTHYLENE-BENZAMIDINE o BENZ ENYL NAPHTHYLENE AMIDINE

NAPHTHYLENE ETHENYL-AMIDINE $\text{C}_{12}\text{H}_{11}\text{N}_2$ s.s. [23] $\text{C}_{10}\text{H}_7 \text{---} \text{N} \begin{matrix} \text{NH} \\ > \end{matrix} \text{CCH}_3$ [168°].

Formed by the action of cold conc alcoholic HCl upon (β) naphthyl ethyl nitrosamine (Fischer a Hepp, *B* 20, 1248) Nodules (from water) Crystallises from methyl alcohol in prisms (containing MeOH) [75°] Sl sol hot water — $\text{B}^{\text{H}}\text{Cl}$ 3aq colourless needles, sl sol water, m sol alcohol — $\text{B}^{\text{H}}\text{PtCl}_4$ 3aq — $\text{B}^{\text{H}}\text{SO}_4$ — $\text{B}^{\text{H}}\text{C}_6\text{H}_4(\text{NO}_2)_3\text{OH}$ needles

Naphthylene ethenyl-amine

[12] $\text{C}_{10}\text{H}_7 \text{---} \text{N} \begin{matrix} \text{N} \\ > \end{matrix} \text{CMe}$ Formed from the acetyl derivative of nitro (β) naphthylamine by reduction with tin and HCl (Liebmann a Jacobson, *A* 211, 67) Formed also by the action of cold alcoholic HCl on (β)-naphthyl-ethyl-amine (Fischer a Hepp, *B* 20, 2472) — $\text{B}^{\text{H}}\text{Cl}$ 2aq needles, v sol water

NAPHTHYLENE-ETHYL-DIAMINE

$\text{C}_{10}\text{H}_7(\text{NH}_2)(\text{NHEt})$ [14] Formed by reducing nitroso-(α) naphthyl ethyl amine with SnCl_4 (Kock, *A* 243, 312) The free base is unstable. It yields (α)-naphthoquinone on oxidation — $\text{B}^{\text{H}}\text{Cl}_2$ [152°] Plates — $\text{B}^{\text{H}}\text{C}_6\text{H}_4(\text{NO}_2)_3\text{OH}$ [180°]. Needles, sl. sol. water and alcohol

DI NAPHTHYLENE-GLYCOL, so called

$C_{20}H_{14}O_2$, \pm $C_{10}H_7C(OH)(?)$ Formed by the action of chloroform and aqueous NaOH on (β)-naphthol (Rousseau, *A Ch* [5] 28, 151). Small crystals, insol water and alkalis, ν sol alcohol, HOAc, and chloroform, ν sol ether

Reactions—1 *Chromic acid mixture* yields crystalline $C_{20}H_{14}O$ [188°]—2 Red hot soda lime forms dinaphthyl [187°]—3 Fuming hydrochloric acid (15 pts) at 160° forms $C_{20}H_{12}Cl_2O$ 3aq crystallising in red needles—4 With fuming hydrobromic acid it forms the corresponding $C_{20}H_{12}Br_2O$ 3aq crystallising in lustrous green plates, whence alcoholic ammonia produces $C_{20}H_{12}(OH)(NH_2)$ crystallising from benzene in needles and forming the crystalline salts $B'H_4Cl_2$, $B'H_4PtCl_6$, and $B'H_4Br_2$. The compound $C_{20}H_{12}Br_2O$ 3aq is converted by hot HOAc into $C_{20}H_{12}BrOHOAc$ crystallising in lustrous green tables and giving off HOAc at 100°—5 *Bromine* in CS_2 forms orange plates of $C_{20}H_{12}Br_2O$ —6 *HIaq* (S G 17) forms, on boiling, crystals of $C_{20}H_{12}I_2O$ —7 Dilute nitric acid (S G 12) forms $C_{20}H_{12}(OH)(NO_2)$, which separates from HOAc as a red crystalline mass $C_{20}H_{12}(OH)(NO_2)HOAc$. Boiling dilute nitric acid forms red needles of $C_{20}H_{12}(NO_2)_2$ [190°]—8 H_2SO_4 (5 pts) at 100° forms $C_{20}H_{12}(OH)(SO_3H)H_2SO_4$ 3aq crystallising in red needles with golden lustre, and separating from HOAc as $C_{20}H_{12}(OH)(SO_3H)HOAc$

Di-acetyl derivative $C_{22}H_{12}(OAc)_2$ [192 5°] Needles, sl sol alcohol, ν sol C_6H_6

Anhydride $C_{20}H_{10}O$ [198 5°] Formed by heating 'dinaphthylene-glycol' with PCl_5 . Formed also by the action of boiling alcohol on the compounds $C_{20}H_{12}Br_2O$ and $C_{20}H_{12}Cl_2O$ (*supra*). Yellow needles (from benzene), almost insol cold alcohol, ν sol boiling benzene. Yields on reduction a compound $C_{20}H_{14}O$

(a)-DI-NAPHTHYLENE KETONE OXIDE

$C_{20}H_{12}O_2$, \pm $C_{10}H_7C(=O)C(=O)C_{10}H_7$ [240°] Formed by boiling (a) naphthyl ethyl carbonate (Bender, *B* 13, 702). Yellow prisms, sl sol alcohol

(b)-Dinaphthylene-ketone oxide (?) $C_{20}H_{12}O_2$

\pm $C_{10}H_7C(=O)C(=O)C_{10}H_7$ [194°] Formed, together with (β)-naphthol, alcohol, and CO_2 , by long boiling of di (β) naphthyl-di ethyl ortho-carbonate ($C_{10}H_7O_2$), $C(OC_2H_5)_2$, its formation is explained by assuming the intermediate formation, by molecular change, of (β) naphthol-carboxylic ether $C_{10}H_7(OH)CO_2Et$, which by elimination of H_2O and di-ethyl carbonate would give dinaphthylene-ketone oxide. Thin colourless prisms (from benzene) ν sl sol alcohol (Bender, *B* 19, 2267)

NAPHTHYLENE MERCAPTAN $C_{10}H_7S$, \pm $C_{10}H_7(SH)_2$ [181°] (G.), [174°] (E) (210° at 15 mm). Prepared by reducing the chloride of naphthalene 'a'-disulphonic acid with zinc-dust and H_2SO_4 , and extracting the product with ether (Grosjean, *B* 23, 2370; Ebert, *B* 24, 145). Pearly leaflets (from alcohol), ν sl sol cold alcohol and ether. Its alcoholic solution gives a yellow ppt with lead acetate. Its alkaline solution is rapidly oxidised by air

Acetyl derivative. [110°]. Crystals.

Benzoyl derivative. [158°].

NAPHTHYLENE-DI-METHYL-DIAMINE

$C_{10}H_7(NH_2)(NMe_2)$ [14] Formed by reducing nitroso naphthyl di methyl diamine or benzene azo-dimethylnaphthylamine (Friedlander, *B* 21, 3124). Liquid, ν sol hot water

Acetyl derivative $C_{10}H_7(NHAc)(NMe_2)$ [195°]

NAPHTHYLENE (a)-NAPHTHYL-DIAMINE

$C_{20}H_{14}N_2$, \pm [14] $C_{10}H_7(NH_2)(NHC_{10}H_7)$. Formed by reducing nitroso di (a) naphthylamine with $SnCl_4$ and HCl (Wacker, *A* 243, 303). Minute crystals (from benzene), ν sol alcohol

NAPHTHYLENE-NAPHTHYL-BENZAM

$INDINE$ $C_{27}H_{18}N_2$, \pm $C_6H_5C \begin{smallmatrix} \diagup N C_{10}H_7 \\ \diagdown N C_{10}H_7 \end{smallmatrix}$ [163°]

Formed by reducing benzoyl-nitro di (β) naphthyl amine with tin and HCl (Ris, *B* 20, 2626). Slender needles (containing C_6H_5) [114°], ν sol alcohol and ether

NAPHTHYLENE DI-NAPHTHYL-SULPH-

IDEOXIDE $C_{30}H_{20}SO$, \pm $C_{10}H_7O C_{10}H_7S C_{10}H_7$ [111°] Formed in small quantity as a by-product in the preparation of (a) and (β)-naphtho nitrile by distilling a mixture of (a) and (β)-potassium naphthalene sulphonate with potassium ferrocyanide (Ekstrand, *B* 17, 2601, *J pr* [2] 38, 140). Long needles. By $K_2Cr_2O_7$ and acetic acid it is oxidised to a compound [162°]. By heating with dilute HNO_3 at 130°-140° it yields a body $C_{20}H_{12}N_2SO$, which crystallises from hot acetic acid in small yellow prisms [231° uncor], nearly insoluble in alcohol and CS_2 . Br and I in CS_2 yield $C_{20}H_{12}Br_2SO$ [182°]

(a)-DINAPHTHYLENE OXIDE $C_{20}H_{12}O$, \pm $C_{10}H_7C(=O)C(=O)C_{10}H_7$ [182°] Formed by distilling (a)-naphthol (1 pt) with lead oxide (3 pts), the yield being 7 pc (Knecht a Unzeitig, *B* 13, 1724, *A* 209, 134), and, together with naphthalene, by heating (a) naphthol at 350°-400° (Merz a Weith, *B* 14, 195) or distilling it with an equivalent quantity of lime (Niederhäusern, *B* 15, 1121). Colourless needles, insol water, sl sol alcohol, ν sol ether

Picric acid compound

$C_{20}H_{12}O \cdot 2C_{10}H_7(NO_2)_2OH$ [173°] Red needles (β) Dinaphthylene oxide $C_{20}H_{12}O$ [155°] (K a U), [157°] (W), [161°] (M a W) ν D 9 05 (obs). Formed by distilling (β) naphthol (1 pt) with PbO (8 pts) (K a U). Formed also by passing a current of air into boiling (β) naphthol (Merz a Weith, *B* 14, 200) and by heating di-oxy (β) dinaphthyl (1 pt) with $ZnCl_2$ (4 pts) for 6 hours at 270° (Walder, *B* 15, 2171). Silvery plates, insol water, sl sol alcohol, ν sol ether

Picric acid compound

$C_{20}H_{12}O \cdot 2C_{10}H_7(NO_2)_2OH$ Red needles, ν sol hot benzene

References—DI-BROMO- and DI-CHLORO- DI-NAPHTHYLENE OXIDE

(a) DINAPHTHYLENE OXIDE SULPHONIC ACID $C_{20}H_{12}(SO_3H)_2O$ Prepared by sulphonating (a)-dinaphthylene oxide—A⁺Ba₂ 2aq needles, sparingly soluble in water with a beautiful blue fluorescence (Knecht a Unzeitig, *B* 13, 1725)

(b)-DINAPHTHYLENE OXIDE-TETRA-SULPHONIC ACID $C_{20}H_4(SO_3H)_4O$ Prepared by sulphonating (β) dinaphthylene-oxide.—A⁺Ba₂ 2aq; tables (K a U.)

DINAPHTHYLENE PHENYL-AMINE ν

PHENYL DINAPHTHYLENE AMINE

NAPHTHYLENE DISULPHOCYANIDE

$C_{10}H_7(SCN)_2$ [78°] Formed from $C_{10}H_7S_2Pb$, alcohol, and cyanogen chloride (Ebert & Kleiner, *B* 24, 146) Needles

 $(\alpha\beta)$ NAPHTHYLENE TOLAZINE $C_{11}H_{11}N_2$

ν $C_6H_5(CH_2)_2 \begin{smallmatrix} \diagup N \\ | \\ \diagdown N \end{smallmatrix} C_{10}H_7$ [141°] Formed by mixing acetic acid solutions, cooled to 0°, of (8) naphthoquinone and tolylene-*o* diamine (Hinsberg, *B* 18, 1229) Distils without decomposition at a high temperature Yellowish crystals ν sol alcohol, acetic acid, and benzene, insol water Dissolves in strong HCl with a brownish red colour

DI NAPHTHYLENE DI THIO DI-UREA

Octohydrate $CS \begin{smallmatrix} \diagup NH \\ | \\ \diagdown NH \end{smallmatrix} C_{10}H_7 \begin{smallmatrix} \diagup NH \\ | \\ \diagdown NH \end{smallmatrix} CS$ [175°] Formed by boiling the tetrahydride of (1,4')-naphthylene diamine with CS_2 and alcohol (Bamberger & Bammann, *B* 22, 951) Crystalline powder, ν sol alcohol

NAPHTHYLENE-UREA $C_{11}H_9N_2O$ ν

$C_{10}H_7 \begin{smallmatrix} \diagup NH \\ | \\ \diagdown NH \end{smallmatrix} CO$ [ν 380°] Formed from naphthylene diamine and $COCl_2$ in toluene at 100° (Hartmann, *B* 23, 1048)

($\alpha\alpha$) DI-NAPHTHYL-ETHANE $C_{22}H_{18}$ ν $(C_{10}H_7)_2C_2H_4$ [160°] Formed, together with (a) naphthyl carbonylamine, by reducing the amide of thio (a) naphthoic acid in alcoholic solution with zinc dust and HClAq (Bamberger, *B* 21, 54) Hexagonal plates, ν sol chloroform and benzene, m sol ether, sl sol alcohol The alcoholic solution exhibits greenish blue fluorescence¹

($\beta\beta$) Di naphthyl-ethane $C_{10}H_7CH_2CH_2C_{10}H_7$ [253°] Formed, in like manner, from thio (8)-naphthoic amide (B) Plates, ν sol hot chloroform and benzene, sl sol ether and alcohol Its solutions fluoresce bluish violet

References — TRI CHLORO and TRI CHLORO-

TETRA NITRO DI NAPHTHYL ETHANE

NAPHTHYL ETHER ν DI NAPHTHYL OXIDE

(a) NAPHTHYL-ETHYL-AMINE $C_{12}H_{11}N$ ν $C_{10}H_7NHET$ Ethyl naphthylamine (303° ν V) at 723 mm (Bamberger & Helwig, *B* 22, 1312) Formed by cohobating naphthylamine with EtBr (Limpriecht, *A* 99, 117, Schiff, *A* 101, 90) Obtained also by reducing $C_{10}H_7NHCSCH_3$ with zinc dust and HClAq (Bernthsen & Trompeter, *B* 11, 1756) Colourless crystals, becoming dichroic (steel blue and brown red) in light Forms a nitrosamine, which, in contact with alcoholic HCl, changes to the isomeric nitroso derivative $C_{10}H_7 \begin{smallmatrix} \diagup N \\ | \\ \diagdown N \end{smallmatrix} O$ [133°] (Kock, *A* 243, 310) — $B'HCl$ [193°] — $B'_2H_2PtCl_4$, yellow prisms — $B'HB'r$ — $B'HI$ four sided prisms

(8) Naphthyl-ethyl-amine $C_{12}H_{11}NHET$ (305°) at 716 mm Oil (Henriques, *B* 17, 2663, Bamberger & Muller, *B* 22, 1297) Yields a red dye with diazotised sulphathio acid $FeCl_3$ gives no colour in the cold, a greenish-brown colour in warm solutions $K_2Cr_2O_7$ and H_2SO_4 give a brown colour and pp — $B'HCl$ [235°]. Plates, sl sol cold water

Nitrosamine $C_{10}H_7NHET(NO)$ [49°].

Crystals Converted by alcoholic hydrogen chloride at 5° into nitroso naphthyl-ethyl amine $C_{12}H_{11} \begin{smallmatrix} \diagup C(NO) \\ | \\ \diagdown CH \end{smallmatrix} NHET$ — $B'HCl$ [108°] Green prisms (from benzene) (Fischer & Hepp, *B* 20, 1248, 2471)

(a) Naphthyl-di-ethyl-amine $C_{12}H_{11}NEt_2$ (291°) S G 1005 Formed by heating (a)-naphthylamine (10 g) with EtBr (15 g) and alcohol or NaOHaq at 120° (B E Smith, *C J* 41, 180, Friedlander, *B* 21, 3129) Formed also by heating (a) naphthylamine hydrochloride with alcohol Oil, ν sol alcohol and ether Forms a nitroso-derivative $C_{12}H_{11}(NO)NEt_2$ [165°]. — $B'HCl$ silky plates, ν sol hot water. — $B'_2H_2PtCl_4$, golden yellow plates — Sulphate: thick prisms

Ethyl iodide $C_{12}H_{11}NEt_2I$ [100°] Cubes

Ethyl-bromide $C_{12}H_{11}NEt_2Br$ Tables (from water)

(8) Naphthyl-di-ethyl-amine $C_{12}H_{11}NEt_2$ (316° ν V) at 717 mm Oil (Bamberger & Williamson, *B* 22, 1760) Yields (8) naphthylamine on heating with lime — $B'HCl$ [175°] Tables or needles, ν e sol water — $B'_2H_2PtCl_4$ [95°]

Di (8) naphthyl-ethyl-amine $(C_{10}H_7)_2NEt$ [231°] From di (8) naphthylamine and EtI at 150° (Ris, *B* 20, 2619) Needles, m. sol cold alcohol, insol petroleum ether

(a) NAPHTHYL-ETHYL-AMINE TETRA

HYDRIDE $C_{12}H_{11}N$ ν $CH_2CH_2C(CNHET)CH$
 $CH_2CH_2C(CH)CH$ (287° ν V) at 717 mm Formed by adding sodium to a solution of naphthyl-ethyl-amine in isoamyl alcohol (Bamberger & Helwig, *B* 22, 1312) Colourless liquid, sl sol water, ν al sol NaOHaq, ν sol alcohol Reduces warm alcoholic $AgNO_3$ Diazobenzene sulphonic acid yields an orange dye $FeCl_3$ added to a solution of its hydrochloride gives a claret colour, changing to greenish yellow $K_2Cr_2O_7$ and H_2SO_4 give a dirty yellow pp in the cold, but in warm solutions a red colour, becoming greenish brown, further addition of $K_2Cr_2O_7$ ppts blue black flakes $KMnO_4$ oxidises it to adipic and oxalic acids — $B'HCl$ [118°] Prisms or needles, ν sol water — $B'_2H_2PtCl_4$ Plates, sl sol cold water

Nitrosamine $C_{10}H_{11}(NEtNO)$ Formed by adding $NaNO_2$ to a solution of the base in HClAq Yellow oil, exhibiting Liebermann's reaction Tin and HClAq reconvert it into $C_{10}H_{11}(NEtH)$ When dissolved in alcoholic HCl it slowly changes to the isomeric $C_{10}H_7 \begin{smallmatrix} \diagup C(NHET) \\ | \\ \diagdown C(NO) \end{smallmatrix} CH$ crystallising in golden needles [119°]

(α -(8) Naphthyl-ethyl-amine tetrahydride $CH_2CH_2C(CH)CH(CNHET)$ (291° ν) at 724 mm $CH_2CH_2C(CH)CH$ Formed, together with the alicyclic isomeride, by reducing (8) naphthyl ethyl amine (15 g) in isoamyl alcohol with sodium (24 g) (Bamberger & Müller, *B* 22, 1304) Colourless oil, volatile with steam, ν sol alcohol, ν al sol water, insol NaOHaq Smells like piperidine $FeCl_3$ colours a warm solution of its hydrochloride reddish-brown $KMnO_4$ oxidises it to adipic acid — $B'HCl$ [173° ν] Needles, ν sol water, ppd as plates by addition of HCl — $B'_2H_2PtCl_4$ Needles.

(α -(8) Naphthyl-ethyl-amine tetrahydride $C_{10}H_7 \begin{smallmatrix} \diagup CH \\ | \\ \diagdown CH \end{smallmatrix} NHET$ (267°) at 724 mm S G. $CH_2CH_2CH_2CH_2$

M 998 Formed as above (B & M) Colourless oil, sl sol water, very volatile with steam FeCl_3 , colours a warm solution of its hydrochloride reddish brown. Reacts with diazobenzene nitrate, forming $\text{C}_{10}\text{H}_{11}\text{N} \cdot \text{Et} \cdot \text{N}_2\text{C}_6\text{H}_4$, [58°] — B'HCl [223.5°] Prisms (from water) or needles (from CHCl_3). Ppd in plates by adding HCl to its aqueous solution — B'ONO_2 , [184°] Needles or plates, v sol hot water — B'ONO_2 , [180°] Needles — $\text{B'H}_2\text{PtCl}_6$, [204°] Orange yellow stellate crystals — $\text{B'C}_6\text{H}_5(\text{NO}_2)_2\text{OH}$, [183.5°] Needles, v sol water and alcohol.

Acetyl derivative $\text{C}_{10}\text{H}_{11}\text{N} \cdot \text{EtAc}$ (328° uncor.) at 718 mm Oil

Nitrosamine $\text{C}_{10}\text{H}_{11}\text{N} \cdot \text{Et}(\text{NO})$ Yellow oil, v sl sol cold water

Ar-(β) Naphthyl-di-ethyl-aminetetrahydride $\text{CH}_2\text{CH}_2\text{C}(\text{CH}_2\text{CH}_2\text{N} \cdot \text{Et})_2$ (298°) at 709 mm The $\text{CH}_2\text{CH}_2\text{C}(\text{CH}_2\text{CH}_2\text{N} \cdot \text{Et})_2$ chief product of the reduction of $\text{C}_{10}\text{H}_7\text{N} \cdot \text{Et}_2$ by sodium (Bamberger & Williamson, *B* 22, 1763) Liquid, sl sol water Yields a red dye with *p*-diazobenzene sulphonic acid KMnO_4 oxidises it to adipic acid — B'HCl granules, v e sol water

Ac-(β) Naphthyl-di-ethyl aminetetrahydride $\text{C}_6\text{H}_5\text{CH}_2\text{CH}_2\text{CH}(\text{CH}_2\text{CH}_2\text{N} \cdot \text{Et})_2$ Liquid, sl sol water, in-sol NaOH aq — $\text{B'H}_2\text{CO}_3$, white needles — B'HCl prisms, v sol water

(a) **NAPHTHYL ETHYL CARBONATE** $\text{C}_{10}\text{H}_7\text{O}_2$, v e $\text{C}_{10}\text{H}_7\text{OCO OEt}$ [31°] Obtained from (a)-naphthol and chloroformic ether (Bender, *B* 13, 702, 19, 2266) Tables, sol alcohol By heating to boiling for some time it splits up into (a) naphthol, a body $\text{C}_{21}\text{H}_{12}\text{O}_2$, [240°], which is probably a dinaphthylene-ketone-oxide $\text{C}_{10}\text{H}_6\text{C}(\text{O})_2\text{C}_{10}\text{H}_6$, alcohol, and CO_2

Di-(β) Naphthyl-di-ethyl-orthocarbonate $(\text{C}_{10}\text{H}_7\text{O})_2\text{C}(\text{OC}_2\text{H}_5)_2$ (298°–300°) Obtained by the action of chloroformic ether upon (β) naphthol (Bender) White amorphous mass, melting with the heat of the hand HCl at 250° decomposes it into (β) naphthol, EtCl , and CO_2 . By long boiling it is decomposed into (β) naphthol, a body $\text{C}_{21}\text{H}_{12}\text{O}_2$, which is probably a dinaphthylene ketone oxide, alcohol, and CO_2 .

(a) **NAPHTHYL-ETHYLENE** C_{10}H_6 , v e $\text{C}_{10}\text{H}_6\text{CH}(\text{CH}_2)_2$ Formed by the action of Na_2CO_3 on β bromo- α -naphthyl-propionic acid (Brandis, *B* 22, 2158) Oil, smelling like styrene Bromine in chloroform forms $\text{C}_{10}\text{H}_6\text{CHBrCH}_2\text{Br}$ [168°]

References — *Di-CHLORO-* and *Di-CHLORO-TETRA-NITRO-DI-NAPHTHYL ETHYLENE*

DI-(α)-NAPHTHYL-ETHYLENE-DIAMINE $\text{C}_{22}\text{H}_{20}\text{N}_2$, v e $\text{C}_2\text{H}_5(\text{NHC}_6\text{H}_4)_2$, [127°] Formed from naphthylamine and $\text{C}_2\text{H}_5\text{Br}_2$ (Reuter, *B* 8, 28) — $\text{B'H}_2\text{SO}_4$.

Di-(β) Naphthyl-ethylene-diamine $\text{C}_{22}\text{H}_{20}(\text{NHC}_6\text{H}_4)_2$, [153°] (M), [150°] (B) Formed, together with di-(β)-naphthyl-pyrazine tetrahydride $\text{C}_2\text{H}_4\text{N}(\text{C}_{10}\text{H}_7)_2\text{N}$, [228°] by the action of ethylene bromide on (β) naphthylamine in presence of sodium carbonate (Maschke, *O C* 1886, 824; Bischoff, *B* 23, 1885) Plates and needles, sl sol ether, in sol absolute alcohol.

DI-(α)-NAPHTHYL-ETHYLENE-DI-CARB-AMIC ETHER $\text{C}_{22}\text{H}_{22}\text{N}_2\text{O}$, v e $\text{C}_2\text{H}_5\text{N}(\text{C}_{10}\text{H}_7)_2\text{CO}_2\text{Et}$, [156°] Formed from $\text{C}_2\text{H}_5(\text{NHC}_6\text{H}_4)_2$ and ClCO_2Et (Reuter, *B* 8, 25) V sol alcohol

DI-(α)-NAPHTHYL-ETHYLENE DIOXIDE $\text{C}_{22}\text{H}_{18}\text{O}_2$, v e $\text{C}_2\text{H}_5(\text{OC}_6\text{H}_4)_2$, [126°] Formed from (a) naphthol, KOH , and $\text{C}_2\text{H}_5\text{Br}_2$ (Koelle, *B* 13, 1956) Plates

Di-(β) Naphthyl-ethylene dioxide $\text{C}_{22}\text{H}_{18}(\text{OC}_6\text{H}_4)_2$, [217°] Formed in like manner (K) Plates, sl sol benzene and HOAc , in sol water, alcohol, and ether

(β)-**NAPHTHYL-ETHYL-HYDRAZINE** $\text{C}_{10}\text{H}_7\text{N}_2$, v e $\text{C}_{10}\text{H}_7\text{N} \cdot \text{EtNH}_2$ Formed from (β) naphthyl hydrazine and EtI in EtOH (Hauff, *A* 253, 33) Yellow oil, v sol alcohol Reduces Fehling's solution and H_2O without forming a tetrazone — B'HCl plates

NAPHTHYL ETHYL OXIDE v *Ethyl ether* of *NAPHTHOL*

NAPHTHYL-ETHYL-NITROSAMINE v *Nitrosamine* of *NAPHTHYL-ETHYL-AMINE*

NAPHTHYL DI ETHYL PHOSPHINE $\text{C}_{10}\text{H}_7\text{P}$, v e $\text{C}_{10}\text{H}_7\text{PEt}_2$, (above 360°) Formed from $\text{C}_{10}\text{H}_7\text{PCl}_2$ and ZnEt_2 (Kelbe, *B* 11, 1501) Yellow oil

Ethyl iodide $\text{C}_{10}\text{H}_7\text{PEt}_2\text{I}$ [209°] Colourless leaflets

NAPHTHYL-GLYCOCOLL v *NAPHTHYL-AMIDO-ACETIC ACID*

DI-NAPHTHYL-GLYCOL v *DI-NAPHTHYLENE-GLYCOL*

NAPHTHYL-GLYCOLLIC ACID v *OXY-NAPHTHYL ACETIC ACID*

(a) **NAPHTHYL-GLYOXYLIC ACID** $\text{C}_{10}\text{H}_6\text{O}_3$, v e $\text{C}_{10}\text{H}_6\text{CO CO}_2\text{H}$ *Naphthoyl formic acid* [114°] Obtained by saponification of its nitrile (Boessneck, *B* 15, 3066, 16, 640), and by oxidation of (a) naphthyl methyl ketone by KMnO_4 (Claus & Feist, *B* 19, 3181) Needles or plates, in sol water, v sol alcohol and ether Gives a red colour on shaking with H_2SO_4 and benzene containing thiophene Yields (a)-naphthoic acid [160°] on oxidation — CaA_2aq v sl water — AgA' white pp, v sl sol water

Amide $\text{C}_{10}\text{H}_6\text{CO CONH}_2$, [151°] Long white needles (from alcohol)

Nitrile $\text{C}_{10}\text{H}_6\text{CO CN}$ *Naphthoyl cyanide* [101°] Formed by heating (a)-naphthoylchloride with HgCl_2 at 100° Needles

(β) **Naphthyl glyoxylic acid** $\text{C}_{10}\text{H}_6\text{CO CO}_2\text{H}$ [c 75°] Formed by gentle oxidation of (β)-naphthyl methyl ketone by dilute KMnO_4 (Claus & Tersteegen, *J pr* [2] 42, 518). Reduced by sodium amalgam to a oxy naphthyl-acetic acid $\text{C}_{10}\text{H}_6\text{CH}(\text{OH})\text{CO}_2\text{H}$

NAPHTHYL-GUANIDINE $\text{C}_{11}\text{H}_{11}\text{N}_3$, v e $\text{NH C}(\text{NH}_2)\text{NHC}_6\text{H}_4$ — $\text{B'H}_2\text{CO}_3$ Trimetric crystals, $a b c = 666 \cdot 1 \cdot 270$ — B'HCl Tri metric crystals (Haushofer, *J* 1882, 365)

Di (a) naphthyl-guanidine $\text{C}_{21}\text{H}_{21}\text{N}_3$, v e $\text{NH C}(\text{NHC}_6\text{H}_4)_2$ *Menaphthylamine* [c 200°] Formed by the action of gaseous cyanogen chloride on (a)-naphthylamine (Perkin, *C J* 9, 8, *A* 98, 238) Small white needles with bitter taste, nearly in sol water, sl sol alcohol and ether. Cyanogen passed into its ethereal solution forms $\text{C}_{22}\text{H}_{21}\text{N}_3$, a pale yellow crystalline mass, in sol. water, in sol alcohol and ether, decomposed by cold HCl aq in $\text{C}_{22}\text{H}_{21}\text{N}_3\text{O}_2$

crystallising in yellow scales [245°], and decomposed by acids into oxalic acid and di-naphthylguanidine—B'HCl amorphous, v sol alcohol and ether, sl sol water—B₂H₂PtCl₆.

Tri-(a) naphthylguanidine C₁₀H₇N₃ . 2e C₁₀H₇N C(NHC₁₀H₇)₂ [178°] Formed from (a) naphthylamine and MeS C(NC₁₀H₇)₂(NHC₁₀H₇) [174°] (Evers, B 21, 962) Flat needles, insol water, v sol alcohol

(β) NAPHTHYL-GUANIDO BENZOIC ACID C₁₀H₇N₂O₂ . 2e C₁₀H₇NH C(NH)NH C₆H₄CO₂H Formed by heating cyanocarbimidamido-benzoic acid with excess of (β) naphthylamine (Griess, B 16, 338) Small crystalline spherules, insol ether, v sl sol hot water and hot alcohol—HA'HCl sparingly soluble six sided plates.

(a) NAPHTHYL-HYDRAZINE C₁₀H₇N₂ . 2e C₁₀H₇NH NH₂ [116°] (203° at 20 mm) Formed by making a paste of (a) naphthylamine and HClAq, adding NaNO₂ in the cold, filtering, and reducing with SnCl₂, the yield being 66 p c (Fischer, A 232, 236) Plates, sl sol water, v sol other solvents—B'HCl plates—B₂H₂SO₄ plates

Reactions—1 Acetone forms C₁₀H₇N₂H CMe₂ [74°]—2 Pyruvic acid produces the acid C₁₀H₇N₂H CMe CO₂H [159°], which forms the ether EtA' [100°] (Schlieper, A 239, 231)—3 Di-bromo pyruvic acid forms the acid C₁₀H₇N₂H CH C(N₂HC₁₀H₇)₂CO₂H [196°] (Nast vogel, A 248, 89)—4 Aceto acetic ether forms oxy-naphthyl-methyl pyrazole C₁₀H₇N₂O [c 190°] (Knorr, B 17, 551)

Tetrahydride C₁₀H₇<CH CH>C(N₂H₇)OH Obtained by treating a solution of the tetrahydride of (a) naphthylamine hydrochloride (18 g) with an equivalent quantity of NaNO₂ and dropping the mixture into a solution of SnCl₂ (45 g) dissolved in HClAq at 0° (Bamberger a Bordt, B 22, 630) Prisms (from boiling ligroin), sl sol water Reduces Fehling's solution at 30°-40° K₂Cr₂O₇ sets free nitrogen in the cold—"B'HCl silvery plates, v sol water

(β) Naphthyl hydrazine C₁₀H₇N₂ . 2e C₁₀H₇NH NH₂ [124°] Formed in the same way as its (a) isomeride (Fischer, A 232, 242) Plates, m sol water, v sol hot alcohol Turns red in air Its solution in conc HOAc is ppd by water

Reactions—1 Acetone forms C₁₀H₇N₂H CMe₂ [65°] (Schlieper, A 236, 174)—2 Aldehyde yields C₁₀H₇N₂H CH CH₂ [128°]—3 Phenyl-acetic aldehyde forms a crystalline hydrazide decomposing at 100° (Ince, A 253, 40)—4 Acetophenone forms a hydrazide crystallising in needles [c 150°]—5 Acetyl-propionic ether forms C₁₀H₇N₂O₂ [130°], the acid corresponding to which at 175° forms the anhydride C₁₀H₇N<N CMe>CO CH₂ [119°] (Steché, A 242, 368)—6 Pyruvic acid in alcoholic solution yields C₁₀H₇N₂H CMe CO₂H [166°], which yields the ether EtA' [131°] (Schlieper, A 236, 176)—7 Di-bromo pyruvic acid yields yellow needles of C₁₀H₇N₂H CH CO C(OH)N₂HC₁₀H₇? insoluble in alkalis (Nastvogel, A 248, 86)—8 Aceto-acetic ether forms C₁₀H₇N₂O [190°]

Salts—"B'HCl needles or plates.—B₂H₂SO₄ plates, sl sol hot water—Nitrate very soluble needles.

(β) Naphthyl-thiocarbamate

C₁₀H₇N₂HS CS N₂H₇O₂H₇ [145°] Plates.

Acetyl derivative C₁₀H₇NH NHAc. [165°] (Hauff, A 253, 25), [167°] (Hillringhaus, B 22, 2657) Formed from (β) naphthyl-hydrazine and HOAc or Ac₂O Needles (from alcohol), m. sol hot water Reduces Fehling's solution

Benzoyl derivative C₁₀H₇NH NHBz [155°] Needles, insol water, v sol hot alcohol

Di-benzoyl derivative C₁₀H₇N₂HBz₂ [163°]

Di-(a)-naphthyl-hydrazine C₁₀H₇N₂H₂C₁₀H₇ Hydrazo naphthalene [275°]

Preparation—1 pt of azonaphthalene is finely suspended in a solution of 1½ pts of NaOH in 160-170 pts of alcohol, and the boiling mixture is treated with zinc dust till decoloured, it is then poured into water containing NH₄HS and the pp is dried and extracted with benzene, from which it crystallises on cooling

Properties—Colourless plates Sublimable v sol alcohol, ether, and benzene, insol water By warming with HCl it is converted into a mixture of two isomeric di-amido-dinaphthyls (Nietzki a. Goll, B 18, 3253)

(1,4)-NAPHTHYL-HYDRAZINE SULPHONIC ACID [14] C₁₀H₇(N₂H₇)SO₂H Formed by reducing the diazo-compound of naphthionic acid with stannous chloride (Erdmann, A 247, 333) Tufts of white needles, sl sol hot water, m sol hot HClAq—A'Na₄aq plates, sl sol cold water

(1,4') Naphthyl-hydrazine sulphonic acid [14'] C₁₀H₇(N₂H₇)SO₂H Formed by reducing the diazo compound of the (1,4') naphthylamine sulphonic acid with SnCl₂ Plates, m sol water, v sol HCl—A'Na₃aq needles, sl sol water

Peri-Naphthyl-hydrazine sulphonic acid [11'] C₁₀H₇(N₂H₇)SO₂H Formed, in like manner, from (1,1')-naphthylamine sulphonic acid Small plates, sl sol hot water—A'K needles, v sol hot water—A'Na plates, v sl sol water

(a) NAPHTHYL IMIDO-DIACETIC ACID C₁₀H₇N(CH₂CO₂H) [133°] Formed by the action of chloro-acetic acid and Na₂CO₃ on (a)-naphthyl amido acetic acid (Bischoff, B 23, 2004) Colourless crystals, v sol alcohol

(a) Naphthylamide C₁₀H₇N(CH₂CO₂H)(CH₂CONHC₁₀H₇) [199°] Crystals (from benzene alcohol)

Di-(a)-naphthylamide C₁₀H₇N(CH₂CONHC₁₀H₇)₂ [202°] Crystals, sl sol ligroin

(β)-Naphthyl-imido-diacetic acid C₁₀H₇NO [182°] Formed by heating chloro-acetic acid with (β)-naphthyl-amido-acetic acid and Na₂CO₃aq at 160° (B) Crystals, sl alcohol, sl sol ether, insol benzene The solutions have a bluish violet fluorescence

(a)-NAPHTHYL-β-IMIDO-BENZYL-MALONIC ETHER C₁₀H₇N₂ . 2e C₁₀H₇ C(NC₁₀H₇)₂CH(CO₂Et)₂ [145°] Formed by the action of α-chloro-benzylidene (a)-naphthylamine upon sodio-malonic ether Crystalline solid sl sol ether By dilute HCl at 120° it is split up into acetophenone and (a)-naphthylamine (Just, B 19, 987)

(β)-Naphthyl-β-imido-benzyl-malonic ether C₁₀H₇NO₂ . 2e C₁₀H₇ C(NC₁₀H₇)₂CH(CO₂Et)₂

[140°]. Formed by the action of ω -chlorobenzylidene-(β)-naphthylamine $C_{10}H_7N \cdot COCl \cdot C_6H_5$ upon sodio malonic ether. Crystalline solid. Sol in ether. By dilute HCl at 120° it is split up into acetophenone and (β)-naphthylamine (Just).

(β)-NAPHTHYL- β IMIDO-BUTYRIC ACID $CH_3C(ONC_6H_7)CH_2CO_2H$ [92°]. Needles, sol water. Formed, together with its naphthalide, by heating a mixture of acetoacetic ether and (β)-naphthylamine at 150°–180°. By boiling with HCl it is converted into (Py 13) oxy-methyl (β)-naphthoquinoline.

Naphthalide $C_{10}H_7NO$ [200°]. Needles, nearly insoluble in most solvents (Knorr, B 17, 543).

DI-NAPHTHYL-IMIDO-THIOCARBAMIC ETHERS $C_{10}H_7NH \cdot C(NC_6H_5)_2SR$ Di-naphthyl-alkyl ψ thioureas. Formed by heating di-naphthyl thio-ureas with alkyl iodides (Evers, B 21, 964).

Di-(a)-naphthyl-imido-thiocarbamic acid Methyl ether $C_{10}H_7NH \cdot C(ONC_6H_5)_2SMe$ [136°]. Plates, sol hot, v sl sol cold, alcohol. Gives off HSMe on heating, leaving di-(a)-naphthyl-cyanamide, which on boiling with dilute HCl aq yields di-(a)-naphthyl-urea. Boiling alcoholic potash also converts it into di-(a)-naphthyl-urea and HSMe. Dilute H_2SO_4 at 160° forms (a)-naphthylamine and $C_{10}H_7NH \cdot CO \cdot SMe$ —B'HI [174°]—B' $_2H_4PtCl_6$ [202°]. Yellow powder.

Ethyl ether EtA' [98°]. Prisms, sl sol hot alcohol—B'HI [157°]—B' $_2H_4PtCl_6$. Propyl ether PrA' [95°]. Plates—B'HI [97°]—B' $_2H_4PtCl_6$.

Ethylene derivative $C_{10}H_7N_2S$ v. $C_{10}H_7N \cdot C \left\langle \begin{smallmatrix} N(C_6H_5)_2 \\ S \cdot CH_2CH_2 \end{smallmatrix} \right\rangle$ [139°]. Formed by boiling di-(a)-naphthyl thio urea with ethylene bromide. Needles, v sol alcohol—B' $_2H_4PtCl_6$.

Di-(β)-naphthyl-imido thiocarbamic acid Methyl ether $C_{10}H_7NH \cdot C(ONC_6H_5)_2SMe$ [110°]. Needles, v. e sol warm alcohol and ether—B' $_2H_4PtCl_6$.

Ethyl ether EtA' [106°]—B' $_2H_4PtCl_6$ [155°].

Propyl ether PrA' [66°]—B' $_2H_4PtCl_6$ [120°].

Ethylene derivative $C_{10}H_7N \cdot C \left\langle \begin{smallmatrix} N(C_6H_5)_2 \\ S \cdot CH_2CH_2 \end{smallmatrix} \right\rangle$ [172°]. Plates (from alcohol)—B' $_2H_4PtCl_6$ [146°].

DINAPHTHYLYNE v DI-AMIDO-DINAPHTHYL ($\alpha\beta$) DI-NAPHTHYL-KETONE $C_{10}H_7O$ v. $(C_{10}H_7)_2OO$ Mol w 268 [136°]. S (alcohol) 13 at 14°. Formed by heating (a) naphthoic acid with naphthalene and P_2O_5 at 210° (Kolbantz a Merz, B 6, 544), by the action of a strip of zinc on a mixture of (a)-naphthoyl chloride and naphthalene (Grucarevic a Merz, B. 6, 1241), and by heating (β)-naphthoyl chloride with mercuric dinaphthyl at 175° (G a M). Pointed needles (from boiling alcohol). On distillation with soda lime it yields naphthalene and a mixture of (a) and (β) naphthoic acids.

($\beta\beta$) Di-naphthyl ketone $(C_{10}H_7)_2CO$ Obtained in two isomeric forms [125.5°] and [164°] by heating (β) naphthoic acid with naphthalene and P_2O_5 (K a M), or (β)-naphthoyl chloride with naphthalene and zinc (G a M). The two varieties may be separated by crystallisation

from ether chloroform. Both varieties yield naphthalene and (β) naphthoic acid on distillation with soda lime. The solubility of the variety melting at 164° in alcohol at 19° is less (08) than that of the variety melting at 125.5° (38). The latter variety may also be prepared by distilling calcium (β) naphthoate (Hausmann, B 9, 1515).

Di-naphthyl ketone $(C_{10}H_7)_2CO$ [140°]. Formed by distilling potassium naphthalene (β) sulphonate with KHC_2O_4 (Giuseppe, B 6, 546).

NAPHTHYL-MELAMINE v CYANTIC ACIDS

(a) NAPHTHYL MERCAPTAN $C_{10}H_7S$ v. $C_{10}H_7SH$ Thionaphthol Mol w 160 (285°) SG $\frac{1}{4}$ 1.1729, $\frac{2}{4}$ 1.1549. Formed by reducing naphthalene (a) sulphonic chloride with zinc dust and dilute H_2SO_4 (Scherer, A 132, 91, Krafft a Schönherr, B 22, 822). Formed also by saponifying its ethyl ether which may be formed by the action of potassium xanthate on (a) diazonaphthalene chloride (Leuckart, J pr [2] 41, 216). Colourless oil, with unpleasant smell, sl sol aqueous alkalis, v sol alcohol and ether. Oxidised in alcoholic solution by the air to di (a) naphthyl disulphide [91°]. Yields di naphthyl sulphide [107°] on heating— $Hg(SC_6H_5)_2$ —Pb(SC_6H_5) $_2$ yellow pp. Ethyl ether $C_{10}H_7SEt$ (187.5°) at 15 mm SG $\frac{1}{4}$ 1.1198, $\frac{2}{4}$ 1.0797. Formed from $C_{10}H_7SH$ by heating with EtI, alcohol, and KOH at 120°–150°.

Acetyl derivative $C_{10}H_7SAc$ (188° at 15 mm) SG $\frac{1}{4}$ 1.1519. Benzoyl derivative $C_{10}H_7SBz$ [117°] (262° at 15 mm).

(β) Naphthyl mercaptan $C_{10}H_7SH$ [75°] (Billeter, B 8, 463, L), [81°] (K a S) (286°). Formed by reducing naphthalene (β) sulphonic chloride, and also by heating (β) diazo-naphthalene chloride with a solution of EtOCS $_2$ K, saponifying the oily product, and boiling with zinc dust and HCl aq (Maikopar, Z 1869, 711, Leuckart, J pr [2] 41, 220). Small plates (from ether), sl sol water. Not volatile with steam. Yields the corresponding disulphide on oxidation—Pb(SC_6H_5) $_2$ orange powder.

Ethyl ether $C_{10}H_7SEt$ [16°] (170.5° at 15 mm).

Acetyl derivative $C_{10}H_7SAc$ [53.5°] (191° at 15 mm). Formed by heating the mercaptan with AcCl at 75°.

Benzoyl derivative $C_{10}H_7SBz$ [108°] (267° at 15 mm).

DI (a)-NAPHTHYL METHANE $C_{10}H_8$ v. $(C_{10}H_7)_2CH_2$ [109°] (above 360°) S (alcohol) 8 in the cold, 6.6 at 78°. Formed by the action of H_2SO_4 on a cooled mixture of naphthalene (5 pts), methylal (1 pt) and chloroform (20 pts) (Grabowski, B 7, 1805). Short prisms (from alcohol), v sol ether and chloroform. Not affected by chromic acid mixture. Picric acid compound $C_{10}H_7 \cdot 2C_6H_4(NO_2)_3OH$ [148°]. Reddish yellow prisms (from chloroform).

Di (β)-naphthyl-methane [92°]. Prepared by reducing di (β) naphthyl-ketone with P and HIAq (Richter, B. 13, 1728). Slender white needles, v sol alcohol and benzene. Yields $C_{10}H_7Br$ [164°] and $C_{10}H_7Br$ [150°–160°].

NAPHTHYL-METHYL-ALCOHOL v. NAPHTHYL CARBINOL.

(a) NAPHTHYL METHYL-AMINE

$C_{10}H_7NHMe$ *Methyl (a)-naphthylamine* (298° uncor) Formed, together with dinaphthylamine, by passing $MeCl$ into melted (a) naphthylamine (Landshoff, *B* 11, 688) Dark red oil Its alcoholic solution gives a violet pp with $FeCl_3$ — $B'_2H_3PtCl_6$, 2aq

Acetyl derivative $C_{10}H_7NMeAc$ [91°] (*L*), [95°] (Norton & Livermore, *B* 20, 2272) Small white prisms, sl sol water, v sol alcohol and ether Dilute nitric acid (10 p c) forms $C_{10}H_7(NO_2)NMeNO_2$, [157 5°]

Benzoyl derivative $C_{10}H_7NMeBz$ [121°] Formed by heating (a) naphthyl-di-methylamine with $BzCl$ at 180° (Hess, *B* 18, 687) Crystals

(a)-Naphthyl-di-methyl-amine $C_{10}H_7NMe_2$ ($C_{10}H_7NMe_2$, (267°) (*L*), (274 5° v at 711 mm) (Bamberger & Helwig, *B* 22, 1815) SG 22 1.0423 Formed by heating (a) naphthylamine (1 mol) with MeI (2 mols) and $MeOH$ (Landshoff, *B* 11, 643, *J pr* [2] 17, 386, Monnet, Reverdin, a Nolting, *B* 12, 2305) Prepared by heating (a)-naphthylamine hydrochloride with $MeOH$ for 8 hours at 170° (Hantzsch, *B* 13, 1348, Friedlander, *B* 21, 3124) Oil

Reactions—1 Forms a nitroso-compound which decomposes in an acid aqueous solution into (1, 4) nitroso naphthol and dimethylamine 2 Nitric acid forms two nitro derivatives [88°] and [128°]—3 By condensation with benzoic aldehyde in presence of $ZnCl_2$ at 110° it yields $C_{10}H_7CH(C_6H_5NMe_2)$, [188°]—4 $C_{10}H_7NMe_2CHO$ (1.4) yields, in like manner, $NMe_2C_6H_4CH(C_{10}H_7NMe_2)$, [179°]—5 $COCl_2$, followed by Aq , yields $NMe_2C_6H_4CO_2H$, [164°] Platinochloride $B'_2H_3PtCl_6$, yellow needles

Methylo-iodide $B'MeI$ Yellowish green flat needles, decomposed at 164° uncor Not affected by $NaOH$, but Ag_2O yields a strongly alkaline hydroxide—($B'MeCl$), $PtCl_6$

ar Tetrahydride $C_8H_7 \begin{smallmatrix} < NMe, CH \\ < CH CH \end{smallmatrix}$

(262°) at 721 mm Formed by reducing the base, dissolved in isoamyl alcohol, with sodium (*B a H*) Colourless oil Yields a colouring matter with *p* diazobenzene sulphonic acid Reduces $AgNO_3$. Oxidised to adipic acid by $KMnO_4$. Yields $B'_2H_3PtCl_6$ and $B'MeI$ [164 5°]

(*B*)-Naphthyl-methyl-amine

Benzoyl derivative $C_{10}H_7NMeBz$ [169°], glistering plates Formed by heating di-methyl-(*B*)-naphthylamine with benzoyl chloride at 180° (Hess, *B* 18, 688)

(*B*)-Naphthyl-di-methyl-amine $C_{10}H_7NMe_2$, [46°] (305° cor) Formed by heating commercial trimethylamine with (*B*)-naphthol at 200° (Hantzsch, *B* 13, 2055), and by heating (*B*)-naphthylamine with MeI and $NaOHAq$ at 120° (Bamberger & Müller, *B* 22, 1806) Forms very soluble salts— $B'_2H_3PtCl_6$

Methylo-iodide $C_{10}H_7NMeI$. Tables, sl sol cold water With Ag_2O it yields a strongly alkaline hydroxide

ar-Tetrahydride $C_8H_7 \begin{smallmatrix} < CH CNMe_2 \\ < CH OH \end{smallmatrix}$

(287°) at 715 mm Formed by reducing $C_{10}H_7NMeI$ dissolved in isoamyl alcohol with sodium (Bamberger & Müller, *B* 22, 1806) Colourless oil. Reduces auric chloride and $AgNO_3$. $FeCl_3$ gives a turbidity and a yellow

colour $K_2Cr_2O_7$ and H_2SO_4 give a yellow pp and, on heating, a dirty-green colour $KMnO_4$ oxidises it to adipic acid— $B'HCl$ — $B'_2H_3PtCl_6$ — $B'HClHgCl_2$. [127 5°] Needles, v sl sol cold, v sol hot water Picrate needles

ac-Tetrahydride $C_8H_7 \begin{smallmatrix} < CH_2 CHNMe_2 \\ < CH_2 CH_2 \end{smallmatrix}$

(166 5° at 22 mm) Formed at the same time as the aromatic isomeride— $B'HCl$ needles, v sol water— $B'_2H_3PtCl_6$, orange needles, v sol water

Di (B) naphthyl methyl-amine $(C_{10}H_7)_2NMe$ [140°] Formed from $(C_{10}H_7)_2NH$ and MeI at 100° (Ris, *B* 20, 2619) Needles, m sol cold alcohol, insol ligroin

Isomeride of naphthyl methyl-amine v NAPHTHYL-CARBINYL AMINE

(a) NAPHTHYL METHYL KETONE $C_{10}H_7O$ $C_{10}H_7COCH_3$, [34°] (297°) Formed by the action of $AcCl$ in presence of $AlCl_3$ on naphthalene dissolved in ligroin (Pampela Schmidt, *B* 19, 2898, Claus a Feist, *B* 19, 3180, *J pr* [2] 42, 517) Crystals, insol water, v sol alcohol and ether Oxidised by $KMnO_4$ to (a) naphthyl glyoxylic acid Yellow ammonium sulphide at 220° forms $C_{10}H_7CMe \begin{smallmatrix} < O \\ < NH \end{smallmatrix}$ [154°] (Willgerodt, *B* 20, 2468)

Oxim $C_{10}H_7CMe(NOH)$ [101°] (*P a S*), [145°] (*C a F*)

Phenyl hydrazide $C_{10}H_7CMe(N_2HPH)$ [146°] (*P a S*), [173°] (*C a F*) Needles

Anilide $C_{10}H_7CMe(NPh)$ [130°] It will be seen that the melting points of the oxim and phenyl hydrazide of the ketone prepared by Claus are the same as those of the like derivatives of the (*B*) ketone

(*B*) Naphthyl methyl ketone $C_{10}H_7COCH_3$, [52°] (301°) Formed, together with the (a)-isomeride by the action of Ac_2O on naphthalene in presence of $AlCl_3$ (Roux, *A Ch* [6] 12, 289, Müller a Von Pechmann, *B* 22, 2561) Leaflets, nearly insol cold water Oxidised by dilute $KMnO_4$ to (*B*) naphthyl glyoxylic acid [*c* 75°], further oxidation forms (*B*) naphthoic acid Sodium amalgam reduces it to $C_{10}H_7CH(OH)CO_2H$ [176°]

Oxim $C_{10}H_7C(NOH)CH_3$, [145°]

Acetyl derivative of the oxim [134°]

Phenyl-hydrazide [171°]

NAPHTHYL METHYL OXIDE v *Methyl ether of NAPHTHOL*

(a) NAPHTHYL-DI-METHYL-PYRROLE

$C_{10}H_7NMe_2$ $C_{10}H_7N \begin{smallmatrix} < CMe CH \\ < CMe CH \end{smallmatrix}$ [123°] (312°)

Formed by heating its dicarboxylic acid at 250° (Knorr, *A* 236, 309) Insol water, v sol alcohol, ether, and chloroform

(*B*) Naphthyl-di-methyl-pyrrole $C_{10}H_7N$ [71°] (341°) Formed in like manner

(a)-NAPHTHYL-DI METHYL PYRROLE DICARBOXYLIC ACID $C_{10}H_7NO_4$, $C_{10}H_7N \begin{smallmatrix} < CO CO_2H \\ < CO CO_2H \end{smallmatrix}$ [244°] Formed by sa

ponifying its ether, which is obtained by the action of (a)-naphthylamine on diacetyl succinic ether (Knorr, *A* 236, 308) Needles— K_2A'' — BaA'' — $AgHA''$.

Ethyl ether EtA'' . [91°]

(*B*). Naphthyl-di-methyl pyrrole-di-carboxylic acid $C_{18}H_{11}N_2$ $\begin{matrix} \diagup \\ \text{OMe} \text{ C CO}_2\text{H} \\ \diagdown \end{matrix}$ $\begin{matrix} \diagdown \\ \text{OMe} \text{ C CO}_2\text{H} \\ \diagup \end{matrix}$ Its di ethyl ether

is obtained by mixing acetic acid solutions of di aceto succinic ether and (*B*)-naphthylamine (Knorr, *B* 18, 304) Sparingly soluble in most solvents Begins to decompose at 260° with evolution of CO_2 —BaA' —BaH.A'.

Di-ethyl ether A' Et. [124°], needles
DI-NAPHTHYL METHYL- ψ THIOUREA *v*
Methyl ether of DI NAPHTHYL-IMIDO THIOCARBAMIC ACID

DI-(*a*)-NAPHTHYL OXIDE ($C_{10}H_7$)₂O *Naphthyl ether* [110°] Formed by heating (*a*) naphthyl with ZnCl_2 or HCl (Merz & Weith, *B* 14, 195) Plates or tables, sol hot alcohol and ether May be distilled unchanged —Picrate $C_{28}H_{17}O_{12}N_2$ (NO_2)₂(OH) [115°] Red crystals

DI-(*B*)-naphthyl oxide ($C_{10}H_7$)₂O [105°] Obtained by boiling (*B*) naphthol with dilute (50 p c) H_2SO_4 (Graebe, *B* 13, 1849), or by heating it with ZnCl_2 (2 pts) at 190°, or with gaseous HCl (M & W) Formed also by distilling aluminium (*B*)-naphthol (Gladstone & Tribe, *C J* 41, 15) Pearly plates (from alcohol), sol ether Gives an orange colour with conc H_2SO_4 .

Picrate $C_{28}H_{17}O_{12}N_2$ (NO_2)₂(OH) [122°] Small orange prisms

DI-(*a*) NAPHTHYL-PARABANIC ACID

$C_{28}H_{17}N_2O_4$ \pm $\text{CO} < \begin{matrix} \text{N}(\text{C}_6\text{H}_5) \text{ CO} \\ \text{N}(\text{C}_6\text{H}_5) \text{ CO} \end{matrix} >$ [246°]

Formed by passing cyanogen gas into an alcoholic solution of methyl di (*a*)-naphthyl-imido-thiocarbamate and heating the crystals that separate with alcoholic HCl (Evers, *B* 21, 973) Needles, \forall sol hot alcohol, sl sol ether, insol water and dilute acids Decomposed by boiling alcoholic potash into CO_2 , oxalic acid, and (*a*) naphthylamine

(*a*)-NAPHTHYL PHENYL-AMIDO METHYL

KETONE $C_{18}H_{13}NO$ \pm $C_{10}H_7$, $\text{CO CH}_2\text{NHC}_6\text{H}_5$ [130°] Formed by the action of aniline on $C_{10}H_7$, $\text{CO CH}_2\text{Br}$ in alcoholic solution (Pampel & Schmidt, *B* 19, 2899) Red crystals

NAPHTHYL-PHENYL compounds ψ .

PHENYL-NAPHTHYL COMPOUNDS

(*a*) NAPHTHYL DI PHENYL METHYLENE-AMINE $C_{28}H_{17}N$ \pm $C_{10}H_7$, N CPh_2 Formed from (*a*)-naphthylamine and benzophenone chloride Ph_2CCl_2 (Pauly, *A* 187, 215) Golden plates (from ether), split up by acids into benzophenone and (*a*)-naphthylamine

TRI-(*a*)-NAPHTHYL PHOSPHATE

($C_{10}H_7$)₃PO [145°] Prepared by heating (*a*)-naphthol with POCl_3 , the yield being 65 p c of the theoretical (Schäffer, *A* 152, 289, Heim, *B* 16, 1769) Small glistening needles

Tri-(*B*)-naphthyl-phosphate ($C_{10}H_7$)₃PO [111°] Prepared by heating a mixture of (*B*)-naphthol and phosphorus oxy-chloride, the yield being 65 p c of the theoretical (Heim, *B* 16, 1768, cf Schäffer) Fine white needles Insol water, sl sol cold alcohol

NAPHTHYL PHOSPHOROUS ACID *v* NAPHTHALENE PHOSPHINIC ACID

(*B*)-NAPHTHYL-PROPIONIC ACID $C_{18}H_{13}O_2$ \pm $C_{10}H_7$, $\text{CH}_2\text{CH}_2\text{CO}_2\text{H}$ [148°]

Formed by reducing naphthyl-acrylic acid with sodium amalgam (Brandis, *B* 23, 2156). Needles (from alcohol), sol. boiling water.

(*a*) NAPHTHYL PROPYLENE ψ THIO UREA $\text{CH}_2\text{CH}_2\text{CH}_2\text{S} > \text{CNH } C_{10}H_7$, [184°] Formed by

heating *a* naphthyl allyl thio urea [145°] with HClAq at 100° (Frager, *B* 22, 3001) Tables, \forall sol chloroform, m sol ether, insol water — $\text{B}'\text{H}_2\text{PtCl}_6$ [206°] — $\text{B}'\text{C}_6\text{H}_5(\text{NO}_2)_3\text{OH}$ [192°]

NAPHTHYL PURPURIC ACID $C_{18}H_{11}N_3O_4$ The K salt is formed by the action of aqueous KCy on di nitro naphthol (Sommaruga, *B* 4, 94, *A* 157, 328) The free acid is unstable The K salt is golden brown with metallic lustre Potash fusion yields benzoic, phthalic, and hemimellitic acids The following salts were dried at 100° $\text{NH}_4\text{A}'$ — KA' — CaA' , — BaA' .

DI-(*a*)-NAPHTHYL PYRAZINE TETRAHY

DRIDE $C_{18}H_{11}N_4$ $< \begin{matrix} \text{CH}_2 \text{ CH} \\ \text{CH}_2 \text{ CH} \end{matrix} > \text{NC}_6\text{H}_5$, *Di-naphthyl-di-ethylene diamine* [265°] Formed from

ethylene bromide, (*a*) naphthylamine, and NaOAc (Bischoff, *B* 22, 1782) Prisms, \forall sl sol alcohol

Di (*B*) naphthyl-pyrazine tetrahydride

[228°] Formed by the action of C_6H_5Br , on (*B*) - naphthylamine in presence of Na_2CO_3 (Bischoff, *B* 23, 1984) Colourless crystals, insol water, alcohol, ether, and HClAq , sol hot HOAc

(*a*) NAPHTHYL-PYRIDINE HEXAHY

DRIDE $C_{18}H_{11}N_2$ (NO_2)₂ (185°–190° at 5–10 mm) Formed by heating piperidine (2 mols) with (*a*) bromo naphthalene (3 mols) for 10 hours at 255° (Lellmann & Buttner, *B* 23, 1383) Thick yellow oil, with faint fæcal odour, rapidly becoming brown \forall sol alcohol and ether A solution of its hydrochloride is pptd by HgCl_2 , by ZnCl_2 , and by chloride of gold — $\text{B}'\text{HCl}$ groups of needles, \forall sol water — $\text{B}'\text{H}_2\text{PtCl}_6$ 2aq (*B*)-Naphthyl pyridine hexahydride [58°]

Resembles the preceding in mode of preparation and properties Colourless prisms, becoming grey on keeping — $\text{B}'\text{HCl}$ — $\text{B}'\text{H}_2\text{PtCl}_6$ 6aq

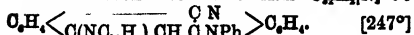
DINAPHTHYL-DIQUINONE, so-called,

$C_{28}H_{18}O_4$ \pm $C_{10}H_7$ $\begin{matrix} \text{OO CO OH} & \text{OH CO CO} \\ \diagdown & \diagup \\ \text{O} & \text{O} \end{matrix} > C_{10}H_7$

Obtained by oxidation of dinaphthyl-diquinhydrone $C_{28}H_{18}O_2$, a black powder formed by the action of dilute H_2SO_4 on (*B*) naphthoquinone (Stenhouse & Groves, *C J* 33, 415) Formed also by the oxidation of (*B*)-amido (*a*) naphthol by FeCl_3 or CrO_3 (Zincke & Rathgen, *B* 19, 2483) Small orange prisms, insol water, \forall sl sol other solvents Not affected by boiling HNO_3 or by H_2SO_4 .

Reactions —1 Yields on reduction 'dinaphthyl dihydroquinone' —2 Distillation with *vacuo* converts it into dinaphthyl-8 Alkalies form a green solution which, on exposure to air, changes to red from formation of $C_{28}H_{18}(\text{OH})_2O_4$ [245°–260°] (Korn, *B* 17, 3020) —4 KMnO_4 oxidises it to diphtallic acid $C_{10}H_6O_4$ —5 Aniline forms $C_{24}H_{15}N_2O_4$ crystallising in red plates [250°] and forming a hydrochloride $C_{24}H_{15}N_2O_4\text{HCl}$ crystallising in needles

Iso-dinaphthyl-diquinone $C_{28}H_{18}O_4$ [250°–260°] Formed by oxidising di-(*B*) naphthyl with CrO_3 in HOAc (Staub & Watson Smith, *C J* 47, 104) Amorphous yellow powder, sol. HOAc , sl. sol. most liquids. Turns brown at 215°.

NAPHTHYL-ROSINDULINE $C_{20}H_{11}N$, *tc*

Formed by heating benzene azo-di-(α)-naphthylamine with aniline and alcohol at 165° (Fischer & Hepp, *A* 256, 246) Black needles, forming a blue solution in conc H_2SO_4 . Conc $HClAq$ at 210° splits it up into (α)-naphthylamine and rosindone $C_{20}H_{11}N_2O$

(α) **NAPHTHYL SILICATE** $C_{10}H_7SiO_3$, *tc* ($C_{10}H_7$), SiO_3 (425° – 430° at 180 mm) Needles (Hertkorn, *B* 18, 1696)

(β)-Naphthyl silicate. (430° at 183 mm.) Crystalline (H)

NAPHTHYL-SULPHAMIC ACID ν **NAPHTHYLAMINE** ν **SULPHONIC ACID****NAPHTHYL SULPHATE.**

Naphthyl sulphuric acid $C_{10}H_7SO_3$, *tc* $C_{10}H_7O SO_3 OH$ Formed by the action of $Cl SO_3 OH$ on a cold solution of (β) naphthol in CS_2 (Armstrong, *B* 15, 204) It is also formed by dissolving (β) naphthol (1 pt) in cold H_2SO_4 (2 pts) (Nietzki, *B* 15, 305) —* NaA' Slender leaflets, ν e sol water Ppd from aqueous solution by $NaCl$ Split up by boiling $HClAq$ into (β) naphthol and $NaHSO_4$. Does not react with diazo salts —* KA' scales (from hot water) With PCl_5 it yields chloro naphthalene Bromine forms bromo naphthol [84°]

DI-(α) NAPHTHYL SULPHIDE $C_{20}H_{11}S$, *tc* ($C_{10}H_7$), S [110°] (290° at 15 mm)

Formation—1 By distilling a dry mixture of potassium naphthalene (α) sulphate and potassium sulphocyanide (Armstrong, *B* 7, 407) 2 By distilling the lead salt of (α) naphthyl mercaptan under diminished pressure (Krafft & Schönherr, *B* 22, 823) —3 By heating (α) naphthyl mercaptan as long as H_2S is evolved (Leuckart, *J pr* [2] 41, 217) —4 By heating ($C_{10}H_7S$), Pb with (α) bromo naphthalene at 235° for 4 hours (Krafft & Bougois, *B* 23, 3045)

Properties—Needles, ν sol benzene and $HOAc$, sl sol alcohol Chromic acid in $HOAc$ oxidises it to the sulphoxide [165°], and finally to the sulphone [187°] (Krafft, *B* 23, 2368)

($\alpha\beta$)-**Di-naphthyl sulphide** ($C_{20}H_{11}S$), S [61°] (290° at 15 mm) Formed by heating the lead salt of (β) naphthyl mercaptan with (α) bromo naphthalene at 230° (Krafft, *B* 23, 2368) Lustrous leaflets (from alcohol) Oxidised by CrO_3 in $HOAc$ to ($C_{20}H_9$), SO_2 [123°]

Di (β) naphthyl sulphide ($C_{20}H_{11}S$), S [151°] (296° at 15 mm) Formed by distilling the lead salt of (β) naphthyl mercaptan under 10 mm pressure (K & S) Plates, ν sl sol hot alcohol, ν sol CS_2 . Oxidised by CrO_3 in $HOAc$ to ($C_{20}H_9$), SO_2 [177°] (Krafft, *B* 23, 2366)

($\alpha\alpha$) **Di-naphthyl disulphide** ($C_{20}H_9$), S_2 , [91°] Formed by oxidising (α) naphthyl mercaptan by exposing its alkaline solution to the air (Schertel, *A* 182, 91, Leuckart, *J pr* [2] 41, 217) Monoclinic crystals, sl sol alcohol

Di-(β) naphthyl disulphide ($C_{20}H_9$), S_2 [137°] (L), [189°] (O) The chief product obtained by saponification of the oil which is formed by the action of potassium xanthate on (β) diazo-naphthalene chloride (Leuckart, *J pr* [2] 41, 221) Formed also by reducing naphthalene (β)-sulphonic chloride with $HIAq$ (Cleve, *B* 21, 1100). Colourless plates, insol. water, ν sol.

alcohol and ether Reduced by zinc and H_2SO_4 , to (β) naphthyl mercaptan.

(β)-**NAPHTHYL SULPHOCYANIDE** $C_{11}H_7NS$, *tc* $C_{10}H_7SON$ [35°] Formed by the action of cyanogen chloride on $Pb(SC_2H_5)_2$ (Billeter, *B* 8, 463) Decomposes on distillation KHS forms $KSCN$ and HSC_2H_5 . Conc $HClAq$ forms $C_{10}H_7SH$, NH_4 , and CO_2 . Sodium amalgam reduces it to sodium cyanide and ($C_{10}H_7$), S_2

($\alpha\alpha$) **DI-NAPHTHYL SULPHONE** ($C_{20}H_{11}S_2$), SO_2 [187°] Formed by oxidising ($\alpha\alpha$)-di-naphthyl sulphide by CrO_3 in $HOAc$ (Krafft, *B* 23, 2368, cf Leuckart, *J pr* [2] 41, 218)

($\alpha\beta$) **Di naphthyl sulphone** ($C_{20}H_{11}S_2$), SO_2 [123°] Formed, together with the ($\beta\beta$) isomeride by heating naphthalene (8 pts) with H_2SO_4 (8 pts) at 180° (Stenhouse & Groves, *B* 9, 682, Cleve, *B* 10, 1723, *Bl* [2] 25, 256, cf Berzelius, *A Ch* [2] 65, 290) Formed also by oxidising the corresponding di naphthyl sulphide (Krafft, *B* 23, 2369) Prisms (from CS_2), m sol boiling alcohol and ether

($\beta\beta$) **Di-naphthyl-sulphone** ($C_{20}H_{11}S_2$), SO_2 [177°] Formed as above, and also by dissolving ($\beta\beta$) di naphthyl sulphide (1 pt) in $HOAc$ (200 pts), warming, slowly adding a mixture of $K_2Cr_2O_7$ (3 pts) dilute (1 3) H_2SO_4 (20 pts) and $HOAc$ (50 pts), filtering, evaporating, and crystals from alcohol (Krafft, *B* 23, 2366) Long white needles When heated with PCl_5 it yields (β) chloro naphthalene and naphthalene (β) sulphonic chloride (Cleve, *Bl* [2] 25, 25)

DINAPHTHYL SULPHONIC ACID ν **DI****NAPHTHYL****($\alpha\alpha$) DI-NAPHTHYL SULPHOXIDE**

($C_{20}H_{11}S$), SO [164.5°] Formed by oxidation of ($\alpha\alpha$) di naphthyl sulphide with chromic acid mixture and $HOAc$ (Krafft, *B* 23, 2367) Formed also by oxidation of naphthylene di naphthyl sulphoxide $C_{20}H_{11}SO$ [111°] (Ekstrand, *B* 17, 2603) White crystals (from alcohol)

NAPHTHYL SULPHURIC ACID ν **NAPHTHYL****SULPHATE****NAPHTHYL SULPHYDRATE** ν **NAPHTHYL****MERCAPTAN****(α) NAPHTHYL-THIOCARBAMIC ACID**

$C_{10}H_7NH CO SH$ *Methylether MeA'* [122°] Formed by the action of dilute H_2SO_4 on methyl di naphthyl imido thiocarbamate [136°] (Evers, *B* 21, 970) Needles, ν sol dilute alcohol, insol water, alkalis, and dilute acids Alcoholic ammonia forms, on heating, (α) naphthyl-urea

Ethylene derivative

$CO < \begin{array}{c} N(C_{10}H_7) \\ S CH_2 CH_2 \end{array} > [102^\circ]$ Formed by the action of dilute $HClAq$ at 200° upon the substance $C_{10}H_7N CO < \begin{array}{c} N(C_{10}H_7) \\ S CH_2 CH_2 \end{array} >$ (Evers, *B* 21, 970) Needles

(β) **Naphthyl-thiocarbamic acid** *Ethylether* $C_{10}H_7NH CS OEt$ [97°] Formed by heating (β) naphthyl thiocarbamide with alcohol at 130° (Cosner, *B* 14, 58) Needles or plates, ν e sol chloroform, m sol alcohol and ether — $C_{10}H_7N Ag CS OEt$ Ppd by adding ammoniacal $AgNO_3$ to the alcoholic solution

(α)-Naphthyl-thiocarbamic acid

Methylether $C_{10}H_7NH CS SMe$ [185°], Formed by heating $C_{10}H_7NH CO SMe$ with CS_2 (E). Small needles, m sol, hot dilute alcohol.

Ethylene derivative $\text{OS} \left\langle \begin{smallmatrix} \text{N}(\text{C}_6\text{H}_5) \\ \text{S CH}_2\text{CH}_2 \end{smallmatrix} \right\rangle$

[199°] Formed by the action of CS_2 on $\text{CO} \left\langle \begin{smallmatrix} \text{N}(\text{C}_6\text{H}_5) \\ \text{S CH}_2\text{CH}_2 \end{smallmatrix} \right\rangle$ at 160° (Evers, *B* 21, 972). Pearly plates (from hot alcohol)

(*β*) Naphthyl-dithiocarbamic acid *Tetrahydride*. Tetrahydronaphthylamine salt $\text{C}_{10}\text{H}_{11}\text{NH}_2\text{S CS NHC}_6\text{H}_5$ [142°] Formed from the tetrahydride of (*β*) naphthylamine and an ethereal solution of CS_2 (Bamberger & Müller, *B* 21, 857) Needles

(*β*) NAPHTHYL-THIO-SEMI-CARBAZIDE $\text{C}_{10}\text{H}_7\text{NH NH CS NH}_2$ [204°] Formed by heating (*β*)-naphthyl hydrazine hydrochloride with ammonium sulphocyanide in alcoholic solution (Hauff, *A* 253, 80, Hüllinghaus, *B* 22, 2657) Crystals, insol water, sol cold alcohol

(*α*)-NAPHTHYL-THIOCARBIMIDE $\text{C}_{10}\text{H}_7\text{N OS Naphthyl mustard oil}$ [58°] Formed from di-naphthyl-thio urea by distilling with P_2O_5 (Hall, *P M* [4] 17, 804) or by heating it with HClAq (84 p.c.) at 150° (Manger, *B* 15, 1414) Formed also by heating di naphthyl-cyanamide with CS_2 at 200° (Huhn, *B* 19, 2406) Long white needles, sol alcohol, ether, and benzene With naphthylamine in alcoholic solution it combines, forming di naphthyl thio-urea Aniline forms phenyl naphthyl thio urea

(*β*) Naphthyl thiocarbimide $\text{C}_{10}\text{H}_7\text{NCS}$ [62°] Obtained in the same manner as its (*α*) isomeride (Cosner, *B* 14, 61, Huhn, *B* 19, 2407) Needles

(*β*)-NAPHTHYL-THIOCARBIZINE $\text{C}_{11}\text{H}_7\text{NS}$, *is* $\text{C}_{10}\text{H}_7\text{N} \left\langle \begin{smallmatrix} \text{NH} \\ \text{CS} \end{smallmatrix} \right\rangle$ [254°] Formed by heating (*β*)-naphthyl thio-semi-carbazide with dilute (20 p.c.) H_2SO_4 at 135° (Hauff, *A* 253, 31) Pearly plates, insol water, sol ether, v sol warm alcohol May be sublimed— B'HOI needles, v sol hot water— B'HPtCl ,— B'HNO ,

DI-(*α*)-NAPHTHYL-THIOHYDANTOIN

$\text{C}_{10}\text{H}_7\text{N}_2\text{SO}$ *is*. $\text{C}_{10}\text{H}_7\text{N} \left\langle \begin{smallmatrix} \text{N}(\text{O}_2\text{H}) \\ \text{S CH}_2 \end{smallmatrix} \right\rangle \text{CO}$ [176°]. Formed by the action of chloro-acetic acid on di-(*α*) naphthyl thio-urea (Evers, *B* 21, 974) Plates, insol water, v sol alcohol

Di (*β*)-naphthyl-thiohydantoin [174°] Resembles the (*α*) isomeride in mode of preparation and properties

NAPHTHYL-THIOSINAMINE *is* NAPHTHYL-ALLYL-THIO UREA (*q v*)

(*α*) NAPHTHYL-THIO-UREA $\text{C}_{11}\text{H}_9\text{N}_2\text{S}$ *is*. $\text{C}_{10}\text{H}_7\text{NH CS NH}_2$ [198°] Formed by heating naphthylamine hydrochloride with ammonium sulphocyanide (De Clermont & Wehrhn, *C B* 82, 512) Small trimetric prisms (from alcohol); sol water, ether, and cold alcohol

Acetyl derivative $\text{C}_{10}\text{H}_7\text{NH CS NHAc}$ [198°]. *S* (alcohol) 2.5 at 78° Formed from (*α*)-naphthylamine and acetyl sulphocyanide (Miquel, *Bl* [2] 28, 103) Slender needles

Benzoyl derivative $\text{C}_{10}\text{H}_7\text{NH CS NHBz}$. [178°] *S* (alcohol) 2 at 78° Formed from (*α*)-naphthylamine and benzoyl sulphocyanide (Miquel, *A Ch* [5] 11, 328) Lustrous yellow prisms (from alcohol), insol ether

(*β*) - Naphthyl- thio- urea $\text{C}_{10}\text{H}_7\text{NH OS NH}_2$. [180°] Formed by heating (*β*)-naphthylamine hydrochloride with potassium sulphocyanide (Cosner, *B* 14, 61). White trimetric plates.

Di-(*α*)-naphthyl-thio-urea $\text{C}_{21}\text{H}_{15}\text{N}_2\text{S}$ *is*. $\text{CS}(\text{NHC}_6\text{H}_5)_2$ [207° cor] (*E*), [197°] Brieger, *B* 12, 1860, Huhn, *B* 19, 2405) Formed by heating (*α*) naphthylamine (100 g) with CS_2 (50 g) and alcohol (500 g) for 14 hours at 75° (Delbos, *A* 84, 371, Evers, *B* 21, 963) Formed also by passing H_2S into a boiling solution of di-naphthyl-cyanamide ($\text{C}_{10}\text{H}_7\text{N}_2\text{C}$) in dry benzene Needles, almost insol alcohol, ether, and benzene, sol nitrobenzene By adding HgO to its boiling solution in dry benzene it is converted into $\text{C}(\text{NC}_6\text{H}_5)_2$. Mel at 100° forms methyl di- naphthyl- imido- thio- carbamate $\text{C}_{10}\text{H}_7\text{N} \left\langle \begin{smallmatrix} \text{NHC}_6\text{H}_5 \\ \text{SMe} \end{smallmatrix} \right\rangle$ [186°] (*v* Di NAPHTHYL-IMIDO THIO CARBAMIC ETHERS

Octohydride $\text{CS}(\text{NHC}_6\text{H}_5)_2$ [170°] Formed by heating (*α*) naphthylamine tetrahydride with CS_2 and alcohol as long as H_2S is given off (Bamberger, *B* 21, 1795) Needles, v sol alcohol, m sol ether

Di (*β*) naphthyl-thio-urea $\text{CS}(\text{NHC}_6\text{H}_5)_2$ [193°] (*C*, *H*), [203° cor] (*E*) Prepared by heating (*β*) naphthylamine in alcoholic solution with CS_2 for 14 hours at 75° (Cosner, *B* 14, 61, Evers, *B* 21, 963) Formed also by passing H_2S into a boiling solution of di (*β*) naphthyl-cyanamide (carbo di (*β*) naphthyl imide) in dry cumene (Huhn, *B* 19, 2407) White plates (from hot nitrobenzene), v sol alcohol and ether With Mel it forms the compound $\text{C}_{10}\text{H}_7\text{NH} \left\langle \begin{smallmatrix} \text{N}(\text{C}_6\text{H}_5) \\ \text{CSMe} \end{smallmatrix} \right\rangle$ [110°], *v* Di NAPHTHYL-IMIDO THIO CARBAMIC ACID By adding HgO to its boiling solution in benzene it is converted into $\text{C}(\text{NC}_6\text{H}_5)_2$. When heated with alcoholic NH_3 at 100° it yields (*β*) naphthylamine and (*β*) naphthyl-thio urea (Gebhardt, *B* 17, 3045) With mercuric cyanide and ammonia it yields $\text{CN} \left\langle \begin{smallmatrix} \text{C}(\text{NC}_6\text{H}_5) \\ \text{NHC}_6\text{H}_5 \end{smallmatrix} \right\rangle$ [166°], which forms an acetyl derivative [141°] and a benzoyl derivative [188°] (Hefelmann, *C C* 1885, 884)

Octohydride $\text{CS}(\text{NHC}_6\text{H}_5)_2$ [166°] Formed by boiling with alcohol the product obtained by the action of CS_2 on the tetrahydride of (*β*)-naphthylamine (Bamberger & Müller, *B* 21, 858) White needles, v sol alcohol, v e sol ether and benzene

NAPHTHYL-TOLYL-AMINE *v* TOLYL NAPHTHYL AMINE

(*α*) NAPHTHYL UREA $\text{C}_{11}\text{H}_9\text{N}_2\text{O}$ *is* $\text{C}_{10}\text{H}_7\text{NH CO NH}_2$. Obtained by saturating a solution of (*α*)-naphthylamine in dry ether with cyanic acid gas, and crystallising from hot alcohol (Schiff, *A* 101, 90) Formed also, together with di naphthyl urea, by heating naphthylamine hydrochloride (3 pts) with urea (1 pt) at 150°–170° (Pagliani, *G* 9, 80) Flat needles, nearly insol water, m sol alcohol, v sol ether Decomposes at 250° without previous fusion

(*β*)-Naphthyl-urea $\text{C}_{10}\text{H}_7\text{NH CO NH}_2$ [c. 287°] Prepared by heating urea with (*β*)-naphthylamine hydrochloride (Cosner, *B* 14, 62) White needles, sol hot alcohol and hot water

Di-(*α*) naphthyl-urea $\text{CO}(\text{NHC}_6\text{H}_5)_2$ [270°].

Formation—1 By heating the acid oxalate of (*α*)-naphthylamine (Delbos, *A Ch* [4] 21, 68), di-naphthyl-oxamide being first formed (Zinn, *A* 108, 228)—2 By gradually heating (*α*)-naphthylamine (2 pts) with urea (1 pt) to 120° (Pagliani, *G* 9, 28)—3 By boiling di-(*α*) naphthyl-cyanamide with dilute alcohol (Huhn, *B* 19,

2405) —4 By heating (a)-naphthylamine with carbamic ether at 185° (Smolka, *M* 11, 200)

Properties —Plates or needles, insol water, al sol boiling alcohol. Yields naphthylamine and no NH_3 on decomposition by KOH

s-Di-(β)-naphthyl-urea $\text{CO}(\text{NHCO}_2\text{H})_2$ [293°] (H.), [286°] (E)

Formation —1 By the action of HgO on di (β) naphthyl thio-urea suspended in spirit (Huhn, *B* 19, 2406) —2 By boiling di (β) naphthyl cyanamide $\text{C}(\text{NC}_6\text{H}_4)_2$ with dilute alcohol (H.) —3 By boiling potassium di (β) naphthoyl-hydroxylamine with water (Ekstrand, *B* 20, 1360)

Properties —Slender needles, al sol alcohol, ether, benzene, and nitrobenzene

u Di-(β) naphthyl urea $\text{C}_{10}\text{H}_7\text{N CO NH}_2$ [193°] Formed by heating the chloro-formyl derivative of di (β) naphthylamine with ammonia for an hour at 140° (Kym, *B* 23, 428) Groups of long needles (from alcohol), al sol cold alcohol

Tetra (β) naphthyl-urea $(\text{N}(\text{C}_{10}\text{H}_7)_2)_2\text{CO}$ [288°] (K & L.), [295°] (K) Formed by heating di (β) naphthylamine with $(\text{C}_{10}\text{H}_7)_2\text{N COCl}$ at 200°–260° (Kuhn & Landau, *B* 23, 811, 2161, Kym, *B* 23, 1542) Prismatic needles, al sol alcohol and ether, v sol hot benzene

NAPHTHYL-URETHANE v NAPHTHYL-CARBAMIC ETHER

NARCEINE $\text{C}_{20}\text{H}_{24}\text{NO}_4$. [184°] (Blyth), [145° cor.] (Hesse, *A* 129, 251), [162°] (Claus Meixner, *J pr* [2] 37, 1, cf Dott, *Ph* [3] 20, 335) *S* 08 at 13° *S* 80 p alcohol 1 Occurs in opium (Pelletier, *A Ch* [2] 50, 262, Couerbe, *A Ch* [2] 59, 151)

Preparation —1 The aqueous extract of opium, from which morphine has been separated by Gregory's process, is mixed with ammonia, filtered, and pptd by lead acetate. The filtrate is freed from lead by H_2SO_4 , neutralised by ammonia, and evaporated. The narcaine is recrystallised from water (Anderson, *Tr E* 20, 347) —2 A solution of the opium bases in HCl aq is mixed with excess of NaOAc and allowed to stand for 24 hours. The filtrate evaporated to a small bulk on the water-bath deposits, after 24 hours, pure narcaine (Plugge, *Ar Ph* [8] 25, 343)

Properties —Silky needles (containing 2aq), v. sol hot water and alcohol, al sol cold water and chloroform, insol ether. Cannot be sublimed. It loses its water of crystallisation at 100°, and at 140° gives off another H_2O (Hesse, *B* 7, 105) It is insol conc KOH aq , al sol dilute caustic potash and ammonia. Ppd by NaHCO_3 from solutions of its salts. Inactive to light (Hesse, *A* 176, 198) Can be extracted both from acid and alkaline solutions by shaking with benzene or chloroform (Plugge, *Ph*, [8] 20, 401) Narcaine is a somniferous poison, 5 g being probably a fatal dose

Reactions —1 Zinc and HCl aq form a small quantity of an amorphous base $\text{C}_{20}\text{H}_{24}\text{NO}_4$, or $\text{C}_{20}\text{H}_{22}\text{NO}_4$ (Beckett & Wright, *C J* 28, 701) —2 Water at 150° carbonises it —3 Chromic acid mixture yields hemipic acid (10 p c), and methylamine (Beckett & Wright, *C J* 29, 467) —4 Ferric chloride forms hemipic, but no opianic acid. Hemipic acid is also formed, though in smaller quantity, by oxidation by KMnO_4 , or by

MnO_2 and H_2SO_4 . Alkaline KMnO_4 forms narcic acid (*v infra*) —5 Conc HNO_3 yields oxalic acid —6 Boiling caustic potash gives off ammonia and NMe_3 , and forms a sparingly soluble acid $\text{C}_{20}\text{H}_{22}\text{NO}_4$ [210°] —7 Potash-fusion yields protocatechuic acid

Tests —1 Weak sodium solution colours solid narcaine dark blue, the colour not being removed by ammonia. The blue colour is destroyed by boiling water —2 Conc H_2SO_4 turns it brown, and then dissolves it, forming a yellow solution. Narcaine is not coloured by diluted sulphuric acid, but on heating over a water bath a violet red colour appears, which ultimately becomes cherry red. If the red liquid be cooled and a trace of HNO_3 or KNO_3 be added, bluish-violet stripes appear (Plugge, *Ar Ph* [3] 25, 425) —3 Erdmann's solution gives a brown colour, turning reddish-brown on heating —4 Chloride of iodine forms a greenish yellow pp., which dissolves on heating (Dittmar, *B* 18, 1612) —5 Potassium chromate gives no pp in cold saturated solutions of salts of narcaine, but in hot solutions there is formed a pp of narcaine chromate and free narcaine (Plugge, *Ar Ph* [3] 25, 793) —6 Narcaine is a feeble base, so that its salts may be titrated by standard alkali as if they contained no base (Plugge, *Ar Ph* [3] 25, 45) —7 When strongly heated it gives off vapours smelling like herring brine (Hesse) —8 Chlorine-water followed by ammonia gives a red colour

Salts — B'HCl (dried at 100°) concentric needles, sol water and alcohol *S* 36 — B'HCl 2aq (Petit, *Bl* [2] 18, 534) *S* 7 — B'HCl 5aq (Wright, *C J* 27, 109) — B'HCl (Petit) — B'HCl (P) — $\text{B'H}_2\text{PtCl}_6$ (dried at 100°) crystalline powder [195°] Golden needles (Claus) — $\text{B'H}_2\text{PtCl}_6$ 2aq amorphous pp., changing to prisms (Hesse, *A* 129, 250) — $\text{B'H}_2\text{SO}_4$, 10aq crystals (Beckett & Wright, *C J* 27, 69) — $\text{B'H}_2\text{SO}_4$ — $\text{B'H}_2\text{SO}_4$ — $\text{B'H}_2\text{SO}_4$ —Mercuric-chloride, only at first, ultimately crystallising in concentric prisms (Hesse) [118°–122°] (Roser, *A* 247, 175) — $\text{B'H}_2\text{I}$, slender needles (Jørgensen, *B* 2, 460) — $\text{B'H}_2\text{I}$: needles (from alcohol)

Methylo-sodide B'MeI [173° uncor.] colourless needles (Claus & Ritzefeld, *B* 18, 1569)

Methylo-chloride B'MeCl [210° uncor.], easily soluble small white needles — (B'MeCl) PtCl_6 nearly insoluble yellow powder

Methylo-nitrate B'MeNO_3 [186° uncor.], small white soluble needles (C & R.)

Ethylo-sodide B'EtI granular crystals (Beckett & Wright, *C J* 28, 708) — B'EtI 2aq Yields alkaline B'EtOH which is readily split up into narcaine and alcohol

Ethylo-bromide B'EtBr [165° uncor.], fine white soluble needles — (B'EtBr) CdBr_2 : small colourless needles (C & R)

Ethylo-chloride B'EtCl [170° uncor.], small colourless needles — (B'EtCl) PtCl_6 glistering yellow crystals [170° uncor.] (C & R) — (B'EtCl) HgCl_2 aq [127°], white crystalline pp.

Ethylo-nitrate B'EtNO_3 : [155° uncor.], long colourless silky needles, sol water (C & R.) *Ethylo-oxalate* $\text{B'EtC}_2\text{O}_4$ [c 174° uncor.], glistering needles (containing 6aq)

Benzoylo-chloride $\text{B'CH}_2\text{Cl}$ [162° uncor.], fine white needles, v. sol. hot water and

alcohol, insol ether—(B'C,H,Cl),PtCl₄. [165° uncor]; yellow crystalline powder

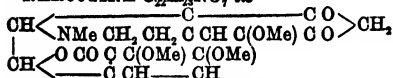
Methyl-narceine C₂₂H₃₂(OH)₂O₂N [175° uncor] Formed by adding KOH (2 pts) to a boiling solution of narceine methyl-iodide (1 pt) in 10 pts of water (Claus & Ritzfeld, *B* 18, 1578) Fine colourless needles *V* sol alcohol, *sl* sol water, nearly insol ether

Benzyl-narceine C₂₂H₃₂(O,H)₂O₂N [169° uncor] Formed by the action of aqueous KOH upon narceine benzyl-iodide (Claus & Ritzfeld, *B* 18, 1574) Long white needles *V* sol alcohol, nearly insol water and ether Alkaline reaction—(B'HCl),PtCl₄ 2aq [128° uncor], dark yellow crystalline pp

Narceic acid C₂₂H₃₂NO₂ [184°] Formed by oxidising narceine with KMnO₄ (Claus & Meixner, *J pr* [2] 87, 3) Prisms (containing 3aq), *v* sol alcohol, ether, chloroform, and hot water Decomposes at 180°–200° into CO₂, dimethylamine, and di-oxy-naphthalene dicarboxylic acid C₁₂H₈O₄ Not acted upon by FeCl₃, —Na₂A'''—Na₂HA''' 5aq [85°]—NaH₂A''' 4½aq—Ba₂A''' 5aq—Ag₂A'''

ψ-Narceine *v* NARCOTINE

NARCOTINE C₂₂H₃₂NO₂ *s*



(Roser, *A* 254, 357) Mol *w* 413 [155°] (Wynter Blyth, *C J* 33, 317, [176°] (Hesse) *S* 014 at 100° *S* (85 p c alcohol) 1 in the cold, 5 at 78° (Duflos, *B J* 12, 214) *S* (ether) 77 in the cold, 21 on boiling (Duflos), 6 at 16° (Hesse) *S* (isoamyl alcohol) 325 in the cold *S* (benzene) 461 in the cold (Kubly, *J* 1866, 823) *S* (chloroform) 38 (Pettenkofer) *S* (EtOAc) 17 (Henry) $[\alpha]_D = -130^\circ$ in ethereal solution (Bouchardat, *A Ch* [3] 9, 213), -185° in alcoholic solution, -207° in chloroform at 22° (Hesse, *A* 176, 192) Contained in opium to the amount of 5 to 8 p c, and was the first alkaloid extracted therefrom (Derosne, *A Ch* 45, 257, Robiquet, *A Ch* [2] 5, 275, Dumas & Pelletier, *A Ch* [2] 24, 188, 50, 269, Liebig, *A* 6, 35, Brandes, *A* 2, 274, Couerbe, *A Ch* [2] 59, 159, *A* 17, 174, Regnault, *A Ch* [2] 68, 137, Wöhler, *A* 50, 1, Blyth, *A* 50, 29, Wertheim, *A* 70, 71, 73, 208, Flückiger, *J* 1869, 797)

Preparation—1 The mother-liquor obtained in the preparation of morphine (*q v*) by Gregory's process is ppd by ammonia, and the pp recrystallised from boiling spirit (Anderson, *A* 86, 179)—2 Opium is extracted with dilute HCl and the bases ppd by KOH The bases are treated with oxalic acid to ppt papaverine, and the filtrate from acid papaverine oxalate ppd by ammonia The pp is recrystallised from alcohol (Hesse, *A Suppl* 8, 284)—3 Conc aqueous NaOAc added to a solution of the hydrochlorides of the opium bases throws down narcotine and papaverine The pp is redissolved in dilute HClAq and mixed with K₂FeCy₄ which ppts papaverine ferricyanide From the filtrate the narcotine is ppd by ammonia (Plugge, *Ar Ph* [3] 25, 348)

Properties—Trimetric prisms, or groups of needles Cannot be sublimed If cooled slowly after melting it forms slender radiating needles.

Insol cold water, *m* sol alcohol and ether. Lævogyrate in neutral solutions, dextrogyrate in acid solutions Insoluble in aqueous potash, nearly insol NH₄Aq Dissolves in boiling baryta-water, but ppd therefrom by NH₄Aq Its solutions do not give a blue colour with FeCl₃ Narcotic poison, 3 g killing a cat Not acted upon by Ac₂O

Reactions—1 Distillation with conc HIAq yields Mel (3 mols) (Matthiessen & Foster, *Tr* 1863, 345, *C J* 16, 342, *A Suppl* 6, 60)—2 Conc HClAq at 110° yields MeCl and, successively, di-methyl-nornarceine, methyl nornarceine, and nornarceine (Matthiessen & Foster, *C J* 21, 257, Matthiessen, *Pr* 17, 337, Matthiessen & Wright, *Pr* 17, 340) Distillation with equal volumes of H₂SO₄ and water gives MeHSO₄ and the three bodies last mentioned (Armstrong, *C J* 24, 56)—3 Boiling dilute KOHAq has no action, but potash-fusion at 200°–220° yields trimethylamine and other volatile amines (Hofmann, *A* 75, 367)—4 Dilute nitric acid at 50° forms meconin, opianic acid, hemipic acid and cotarnine—5 Distillation with HCl and platinum chloride also yields opianic acid and cotarnine The same products are formed by oxidation with H₂SO₄ and MnO₂ (Wöhler)—6 Water at 140° splits it up into meconin and hydrocotarnine (Beckett & Wright, *C J* 28, 583) Sodium-amalgam gives the same products—7 Iodine acting on an alcoholic solution of narcotine forms tarconine methyloperiodide and iodo tarconine methyloperiodide (Jørgensen, *J pr* [2] 2, 446, Roser, *A* 245, 317)

Tests—1 Conc H₂SO₄ forms a yellow liquid which, when heated becomes orange red and finally violet red (Husemann, *A* 128, 305)—2 H₂SO₄ containing a little HNO₃ gives a reddish-yellow colour—3 Does not liberate iodine from iodic acid—4 Does not reduce alkaline K₂FeCy₄ (Kieffer, *A* 103, 277)—5 Bromine gradually added to a boiling solution of narcotine in dilute HClAq forms a rose coloured liquid, the colour being destroyed by excess of bromine—6 Potassium mercuric iodide forms a yellowish white pp (Groves, *C J* 11, 97) Phosphomolybdic acid, picric acid, potassium sulphocyanide, and auric chloride also give pps in solutions of salts of narcotine—7 A solution of potassium chromate added to one of a salt of narcotine, ppts free narcotine (Plugge, *Ar Ph* [3] 25, 793), K₂Cr₂O₇ ppts narcotine bichromate—8 Salts of narcotine require, on titration, the same amount of alkali for neutralisation as if the base were absent (Plugge, *Ar Ph* [3] 25, 45)—9 Chloride of iodine gives a curdy yellowish pp (Dittmar, *B* 18, 1612)

Salts—B'HCl Needles, *v* sol water $[\alpha]_D = -47^\circ$ to -50° in a 2 p c solution containing excess of HCl $[\alpha]_D = -104.5^\circ$ in a 2 p c solution in 80 p c alcohol containing excess (1 mol) of HCl (Hesse, *A* 176, 192)—B'HClAq (Dott, *J* 1884, 1889)—B'HCl (Beckett & Wright, *C J* 29, 164)—B'HCl—B'HCl (B a. W.)—Mercuric chloride white pp (Hinterberger, *A* 82, 311)—B'H₂PtCl₄ yellow crystalline pp (Blyth) Decomposed by boiling water (De Coninck, *Bl* [2] 45, 181)—B'H₂PtCl₄ 2aq (Hesse)—B'H₂SO₄ 4aq (Dott)—B'H₂CrO₄—B'HI, shining laminae, converted by boiling alcohol into tarconine periodide C₂₂H₃₁NO₂HI,

Jørgensen, *B* 2, 460) — $B'HOAc$ aq needles (Dott) — Binoxalate v e sol water — Cyanurate $B'H_2O_2C_2N_3$ 1½aq, needles [c 175°], v sl sol water (Claus, *J pr* [2] 38, 229)

Methylo-chloride $B'MeCl$ Formed by heating narcotine with MeI and decomposing the viscid $B'MeI$ with $AgCl$ (Roser, *A* 247, 168) Groups of needles, v sol water and alcohol Yields ψ narceine (v *infra*) on distillation with aqueous $NaOH$ — $B'_2Me_2PtCl_2$

Ethylo iodide $B'EtI$ (How, *A* 92, 327)

Oil **Ethylo chloride** $B'EtCl$ Yields ψ methyl narceine [173°] on treatment with $NaOHAq$ — $B'_2Et_2PtCl_2$

Nornarcotine $C_{12}H_{17}NO$, Obtained by heating narcotine with fuming $HIAq$ (Matthiessen & Wright, *Pr* 17, 340) White amorphous mass, turning brown in air Almost insol water, insol alcohol and ether A concentrated solution of its hydrochloride is ppd by HCl and also by water The hydrochloride is tasteless, and its solution gives with KOH , Na_2CO_3 , and NH_3 , pps soluble in excess, and with platinum chloride a yellow pp, quickly turning brown

Methyl derivative $C_{12}H_{19}NO$, Formed by heating narcotine with $HClAq$ for some days White amorphous mass, almost insol water, insol alcohol and ether Its solution in conc $HClAq$ is ppd by water Astringent taste The platinumchloride is a yellow pp, slowly turning brown The ppd base is soluble in excess of NH_4Aq and Na_2CO_3Aq but sl sol $KOHAq$

Di-methyl derivative $C_{14}H_{21}NO$, Formed by heating narcotine with $HClAq$ for 2 hours or with diluted H_2SO_4 at 100° White amorphous mass, almost insol water, v sol alcohol, sl sol ether Its solution in $HClAq$ is ppd by water Its hydrochloride tastes bitter The platinumchloride is a yellow pp The ppd base is insol Na_2CO_3Aq , sl sol NH_4Aq , sol $KOHAq$

ψ -Narceine $C_{22}H_{27}NO_3$, [c 175°] Formed by boiling narcotine with MeI , decomposing the resulting methylo iodide by $AgCl$, adding an equivalent amount of $NaOH$, and distilling with steam (Roser, *A* 247, 169) Slender white needles (containing 3aq), v sol alcohol and hot water, sl sol cold water, insol ether Dissolves in aqueous KOH and NH_3 , but is reppd by CO_2 , Inactive to light Coloured blue by iodine Conc H_2SO_4 forms a brownish yellow solution, turned dirty violet on heating Chlorine water followed by ammonia gives a red colour According to Roser, this body is probably identical with narceine — $B'_2H_2SO_4$ 2aq tufts of needles — $B'HCl$ 3aq — $B'_2H_2PtCl_2$, [198°] Thin needles, insol water — Aurachloride [180°], yellowish red needles — Mercury double salt [123°]

ψ -Methyl-narceine $C_{14}H_{21}NO_3$, ψ -Homo-narceine [173°] Formed by passing steam through a mixture of narcotine ethylo chloride and aqueous $NaOH$ (Roser, *A* 247, 178) White needles (containing 3aq), v sol water and alcohol, insol ether Gives a blue colour with iodine The hydrochloride and sulphate are v sol water — $B'_2H_2PtCl_2$ 2aq small yellow needles

Teropiammon $C_{10}H_{15}NO_{12}$, A crystalline compound, insol. water, found among the pro-

ducts of the action of dilute HNO_3 on narcotine (Anderson) It forms a crimson solution in H_2SO_4 , and yields NH_3 and opianic acid on boiling with potash.

Cotarnine $C_{17}H_{19}NO_4$, s

$[314s] OH_2 \cdot \langle O \rangle \cdot C_6H(OMe)(CHO) OH, CH_2NMe_2$ (Roser, *A* 254, 354) [133°]

Formation — 1 By the oxidation of narcotine by MnO_2 and H_2SO_4 (Wöhler, *A* 50, 19, Matthiessen & Foster, *Pr* 11, 55, Beckett & Wright, *C J* 28, 575), by platinum chloride (Blyth, *Mem C J* 2, 168), by dilute nitric acid (Anderson, *C J* 5, 266, *A* 86, 196) or by $K_2Cr_2O_7$ (M & F) — 2 By the oxidation of hydrocotarnine (Beckett & Wright, *C J* 28, 580)

Properties — Colourless needles, sl sol water, v sol alcohol and ether Decomposed on fusion A solution of its hydrochloride is neither ppd by HCl nor by water Its salts taste bitter The freshly ppd base dissolves in excess of aqueous NH_3 and Na_2CO_3 , but is v sl. sol. $KOHAq$ Decomposes on fusion Cotarnine is not poisonous

Reactions — 1 Nitric acid forms a red solution and, on heating, oxidises it to apophyllenic and oxalic acids — 2 Aqueous HCl , HI , or H_2SO_4 at 140° yields $MeCl$, MeI , or $MeHSO_4$ and cotarnamic acid — 3 Zinc and dilute $HClAq$ yield hydrocotarnine — 4 Bromine-water forms $C_{17}H_{17}Br_2NO_4$ (Wright, *C J* 32, 533) Bromine water added to a solution of cotarnine hydrochloride forms $C_{17}H_{17}Br_2NO_4HBr$, a crystalline pp [190°–200°] (Von Gerichten, *B* 14, 311)

Salts — $B'HCl$ 2½aq long silky crystals (Blyth) — $B'_2H_2PtCl_2$ lemon-yellow crystalline pp, or dark red six sided prisms — $B'HClHgCl_2$ pale-yellow pp — B'_2H_2Br 2aq — $B'HI$ yellow needles, sl sol cold water and alcohol (Roser, *A* 249, 156) — $B'HI$, [142°] (Jørgensen, *J pr* [2] 2, 455)

Benzoyl derivative $C_{22}H_{21}BzNO_4$ ½aq [123°] Formed by shaking cotarnine with $BzCl$ and dilute $NaOHAq$ (Roser, *A* 254, 335) Long needles, insol water, v e sol hot alcohol

Oxim $C_{17}H_{19}NO_4(NOH)$ [168°] Prisms, insol water, m sol alcohol, sol alkalis — $B'HCl$ small yellow needles, v sol water, m sol alcohol — $B'_2H_2PtCl_2$ yellow crystalline pp decomposed by boiling water

Oxim of the benzoyl derivative $C_{24}H_{23}O_5(CHNOH) C_6H_5NMeBz$ [166°]

Formed by warming benzoyl cotarnine with alcoholic hydroxylamine hydrochloride Small pointed crystals, insol water and ether, v sol alcohol, sol $NaOHAq$

Methyl-cotarnine

Methylo-iodide $C_{17}H_{19}IO_4NMe_2I$ Formed by warming cotarnine with MeI (Roser) Yellow needles, sl sol cold water and alcohol Decomposed by boiling $NaOHAq$ into cotarnone and NMe_2

Methylo chloride $C_{17}H_{19}ClO_4NMe_2Cl$ ½aq. Formed from the iodide and $AgCl$ Crystals (from water) On warming with alcoholic hydroxylamine hydrochloride at 100° it yields $C_{17}H_{19}Cl_2N_2O_4$ 2aq or $C_6H_5O_2(CN) C_{17}H_{19}NMe_2Cl$ a crystalline body, v sol water, converted by hot $NaOHAq$ into NMe_2 and $C_6H_5O_2(CN) \cdot CH_2CH_2$

[16 The last body forms a dibromide [140°].

(C₁₁H₁₁O₂NMe₂Cl)₂PtCl₄
Bromo-cotarnine C₁₁H₁₁BrNO₂ [100°]

Formed by treating hydrocotarnine hydrobromide with bromine-water (Wright, *C J* 82, 525) Crystals (from ether), decomposing at 100° Reduced by zinc and HClAq to bromo-hydrocotarnine [78°] — B'Br — Crystals, v sol water Above 200° it melts, gives off gases, and forms a small quantity of tarconine hydrobromide, together with a blue substance C₁₁H₁₁N₂O₂HBr insoluble in alcohol

Cotarnone C₁₁H₁₁O₂ s s
CHO C₁₁H₁₁O₂ CH₂CH₂ [78°] Formed by warming methyl cotarnine methylo chloride with aqueous NaOH (Roser, *A* 249, 163) Plates (from alcohol), slightly volatile with steam Insol cold water, m sol alcohol, ether, and HOAc

Oxam C₁₁H₁₁O₂(CH NOH) CH CH₂ [182°]
Slender needles (from dilute alcohol).

Cotarn-lactone C₁₁H₁₁O₂ s s

CH₂—O—C₁₁H(OMe)—CH(CH₂OH)— [154°]

Formed by oxidising cotarnone with KMnO₄ (Roser, *A* 254, 341) Prisms, sl sol cold alcohol or water Forms with baryta the salt Ba(C₁₁H₁₁O₂)₂5aq

Acetyl derivative C₁₁H₁₁AcO₂ [174°]

Benzoyl derivative C₁₁H₁₁BzO₂ [184°]

Cotarnic acid C₁₁H₁₁O₂ s s

CH₂—O—C₁₁H(OMe)(CO₂H)₂[₂ 3 5 6] [178°]

Formed by oxidising cotarnone or cotarn lactone with aqueous KMnO₄ (Roser, *A* 249, 165, 254, 845) Plates Forms, on melting, the anhydride C₁₁H₁₁O₂ [162°] On heating with P and HIAq it yields gallic acid HClAq at 100° forms CH₂—O—C₁₁H(OMe)CO₂H [210°]

Bromine in HOAc produces the compound

CH₂—O—C₁₁HBr₂(OMe) [160°]

Salts —KHA''2½aq needles, v sol. water —BaA'': plates —Ag₂A'

Hydrocotarnine C₁₁H₁₁NO₂ s s

CH₂—O—C₁₁H(OMe)—CH₂CH₂· [50°] (H).

[55°] (B & W) Occurs in opium (Hesse, *A Suppl* 8, 326) Formed by reducing cotarnine with zinc and HClAq, and, together with meconin, by the action of zinc and HClAq on narcotine (Beckett & Wright, *C J* 28, 577) It is also formed in small quantity (2 to 5 p c) in the preparation of opianic acid by the oxidation of narcotine. Monoclinic prisms (from ether), v sol alcohol and benzene The crystals contain water of crystallisation (½aq) Conc H₂SO₄ forms a yellow solution, changing on heating to crimson and, finally, to a dirty violet colour Easily oxidised to cotarnine Not attacked by AcO Bromine added to a solution of its hydrobromide forms C₁₁H₁₁BrNO₂ [78° cor], which yields the crystalline salts C₁₁H₁₁BrNO₂HBr and (C₁₁H₁₁BrNO₂)₂H₂PtCl₄

Salts —B'HCl aq —B'₂H₂PtCl₄ —B'HI. S 2 at 18° —B'Br 1½aq

Ethyl-o-codide B'EtI White micaceous plates, sl. sol water Yields (B'Et)₂CO₂ 4aq, B'EtCl [100°], both crystalline, and (B'EtCl)₂PtCl₄

Acetyl-hydrocotarnine-acetic acid C₁₁H₁₁NO₂ [201°] Formed by boiling cotarnine (1 pt) with acetic anhydride (10 pts) for 1½ hours Small needles Sol alcohol and benzene, sl sol boiling water, insol cold water, and ether —A'Ag white pp —A'Ca very soluble small needles

Ethyl ether A'Et [118°], white feathery needles (Bowman, *B* 30, 2431).

Oxy-hydrocotarnine methylo-iodide

CH₂—O—C₁₁H(OMe)—CH(OH)NMe₂I

Methyl derivative

C₁₁H₁₁O₂ C₁₁H₁₁(OMe)NMe₂I [178°] Formed by the action of MeI in the cold on a solution of cotarnine in MeOH (Roser, *A* 254, 360) Yellow needles, v sol hot water and alcohol Ppd unchanged by adding NaOH to its warm aqueous solution Boiling with water and Ag₂O expels NHMe₂ —(C₁₁H₁₁NO₂Cl)₂PtCl₄ orange pp

Ethyl derivative C₁₁H₁₁O₂(OEt)NMe₂I [168°] Formed by the action of MeI on a cold solution of cotarnine in EtOH Plates, v e sol water and alcohol.

Isobutyl derivative C₁₁H₁₁O₂N(OEt)I [120°] Formed from cotarnine, MeI, and iso butyl alcohol Plates (from water)

Cotarnamic acid C₁₁H₁₁NO₂ Formed by heating cotarnine with aqueous HCl at 140° (Matthiessen & Foster, *A Suppl* 2, 379, Gerichten, *B* 14, 310) Yields apophyllenic acid on oxidation by dilute HNO₃ —C₁₁H₁₁NO₂HCl aq White needles, sl sol cold water On treatment with nitrous acid or on exposure to air its solution becomes green, with red fluorescence, and exhibits a spectrum resembling that of chlorophyll

Tarconine C₁₁H₁₁NO₂ Formed by heating bromo cotarnine hydrobromide at 200° (Wright, *C J* 82, 535) or tarconine methylo chloride with conc HClAq for 4 hours at 150° (Roser, *A* 245, 321) Ppd from solutions of its salts by KOH, but not by Na₂CO₃ Its hydrochloride heated at 200° forms a blue base C₁₁H₁₁N₂O₂, forming the blue salts B'₂H₂Br₂ and B'₂H₂SO₄ — Salts B'HCl 1½aq needles, v e sol water and alcohol —B'₂H₂PtCl₄ insol alcohol

Methylo-iodide B'MeI Formed by the action of H₂S on the periodide which is formed by boiling narcotine in alcoholic solution with iodine and some HCl Tufts of yellow needles (from water or alcohol) —B'MeI, [160°] Red dish brown needles (from alcohol or HOAc) —B'MeI, (Jørgensen) —B'MeBrI,

Methylo-chloride B'MeCl yellow needles —(B'MeCl)₂PtCl₄ yellow crystalline pp, sl sol water —B'HAuCl₄

Bromo-tarconine C₁₁H₁₁BrNO₂ s s

C₁₁H₁₁BrO C₁₁H₁₁N—CH₂—O [235°–238°] Formed by heating the hydrobromide of the compound C₁₁H₁₁BrNO₂ (v Cotarnine, Reaction 4) with water at 180° (Wright, Gerichten, *A* 210, 84) Orange red needles (containing 2aq), becoming crimson and anhydrous at 100° Sl sol cold water, insol. ether Water at 160° splits it up into cupronine and tarconine Conc HClAq at 120° forms narcic acid, tarconine, CO₂, and CO Chromic acid mixture yields bromoform and apophyllenic acid Bromine-water added to a solution of its hydrochloride yields cuprine, bromo apophyllenic acid, and dibromo apophyll-

lin. Distillation over soda-lime yields pyridine. —B'HCl 2aq crystals, sl sol cold water. —B'HBz 2aq —B'H₂PtCl₄ needles (from hot conc. HClAq)

Methylo-chloride C₁₁H₇BrNO₂MeCl slender needles, v e sol water, m sol alcohol —(B'MeCl)₄PtCl₄ —B'MeAuCl₄

Methylo-bromide C₁₁H₇BrNO₂MeBr Formed by the action of bromine water on an aqueous solution of tarconine methylo chloride, the perbromide first formed being decomposed by H₂S (Roser, A 245, 324) Long needles (containing aq), changing on standing to flat prisms —B'MeBr₂ [165°] Yellow rectangular plates (from alcohol or HOAc)

Methylo-iodide C₁₁H₇BrNO₂MeI [204°] Formed from bromo tarconine and MeI (Gerichten, A 210, 170) Yellow needles, sol water, insol ether Turns brown at 170° Boiling baryta water converts it into formic aldehyde and methyl bromo tarconic acid Moist Ag₂O forms the hydroxide crystallising in small red needles

Ethyl-iodide C₁₁H₇BrNO₂EtI [206°] Formed from ethyl iodide and bromo tarconine Yellow needles Yields (B'EtCl)₄PtCl₄

Iodo-tarconine C₁₁H₇INO₂ Formed by heating its methylo chloride at 180° (Roser, A 245, 319) Crystallises from water in yellowish red needles (containing aq), becoming dark red on drying —B'HCl 2aq silky yellow needles, v sol water, sl sol HClAq

Methylo-chloride C₁₁H₇INO₂MeCl aq Formed from the methylo-iodide and AgCl yellowish white needles (from alcohol), m sol water —(B'MeCl)₄PtCl₄ needles, sl sol hot water —B'MeAuCl₄ needles (from hot water)

Methylo iodide C₁₁H₇INO₂MeI Formed by the action of iodine on narcotine in alcoholic solution, the resulting periodides being decomposed by H₂S (Roser, A 245, 317) Yellow needles, changing on standing to prisms, almost insol cold water, sl sol hot alcohol —B'MeI₂ [171°] Needles, sl sol alcohol

Methyl tarconic acid C₁₁H₇NO₂ [244°] Formed from the aqueous solution obtained by the action of Ag₂O and water on tarconine methylo chloride by boiling alone or with baryta (Roser, A 245, 322, 254, 366) Not a true homologue of tarconic acid Thin yellow needles (containing 2aq), v sol hot water, alcohol, mineral acids, and KOHAq, insol ammonia —C₁₁H₇NO₂HCl aq white prisms or needles, turns yellow on drying at 100° —C₁₁H₇NO₂H₂SO₄ 3aq white prisms —(C₁₁H₇NO₂)₂H₂SO₄ 6aq yellowish crystals

Bromo - methyltarconic acid C₁₁H₇BrNO₂ [238°] Formed by boiling bromo tarconine methylo hydroxide with baryta water (Gerichten, A 210, 79, Roser, A 245, 326) The ppd Ba salt is decomposed by H₂SO₄ or HOAc Yellow prisms (containing 2aq), insol cold water and ether, sl sol hot alcohol Darkens at 215° Heated with conc HClAq it yields MeCl, HBr, and tarconic acid —CuA', —BaA', yellow pp —(HA'HCl)₄PtCl₄ needles, m sol dilute HCl

Ethyl - bromo - tarconic acid C₁₁H₇BrNO₂ [226°] Formed by the action of baryta on bromo-tarconine ethylo iodide (or ethylo hydroxide) (Gerichten, A 212, 182) Yellow needles (containing 2 aq), v sl sol cold water, v sol. alco-

hol, insol ether Its aqueous solution is neutral. Conc H₂SO₄ forms a yellow solution Conc HClAq converts it, on heating, into EtCl, HBr, and tarconic acid —HA'HCl yellow needles —(HA'HCl)₄PtCl₄ —CuA'.

Tarconic acid C₁₁H₇NO₂ Formed by heating bromo methyl tarconic acid with conc HClAq at 155° (Gerichten, A 212, 184) Slender yellow needles, turning brown in air Its alkaline solution is brown, becoming greenish blue on standing It reduces AgNO₃ in the cold —HA'HCl prisms, m sol hot water, insol cold alcohol FeCl₃ colours its solution red

Nartic acid C₂₀H₁₅N₂O₂ *Nartine* Formed by heating bromo tarconine or tarnine with conc HClAq at 130° (Gerichten, A 212, 194, 212, 170) Yellow needles, decomposing at 200° without melting Turns brown in air NaOH does not ppt it from acid solutions Its solution in NaOHAq turns greenish blue when exposed to air Reduces AgNO₃, KMnO₄ oxidises it to a pyridine carboxylic acid On distillation with soda lime it yields pyridine. —H₂A'H₂Cl₂ [above 273°] Yellow needles, m sol water —H₂A'HCl yellow needles, sl sol water

Tarnine C₁₁H₇NO₂ [above 290°] Formed by heating bromo-tarconine with water at 130° (Gerichten) Orange needles (containing 1½ aq) m sol hot water and dilute alcohol, insol ether. —B'HCl yellow needles, sol water, —B₂H PtCl₄ Decomposed by boiling water

Cupronine C₂₀H₁₅N₂O₂ Formed, together with tarnine, by heating bromo tarconine with water at 140° (Gerichten, B 14, 315, A 210, 190°) Black powder, insol hot water, alcohol, and ether Forms a brown solution in aqueous NaOH or Na₂CO₃ Conc H₂SO₄ forms a magenta-red solution, changing to violet on dilution Conc HClAq does the same —B'HCl coppery needles —B'HBz bluish green needles with coppery lustre, sl sol water, forming a blue solution

Cuprine C₁₁H₇NO₂ Formed by the action of bromine on a solution of bromo tarconine hydrobromide (Gerichten, A 210, 89) Minute blue needles with coppery lustre, sol water and alcohol, insol ether Weak base, the hydrochloride crystallising in concentric groups of needles, the platinochloride being a deep-blue flocculent pp

Apophyllic acid is the mono methyl ether of PYRIDINE DICARBOXYLIC ACID

Di bromo-apophyl. lin v vol i p 553

Oxy-narcotine C₂₀H₁₅NO₂ Crystals which remain undissolved in the process of purifying narcaine (Beckett & Wright, C J 29, 461) Small crystals (from alcohol), v sl sol water and alcohol, insol ether, nearly insol benzene and chloroform Ppd from solutions of its salts by NaOH and Na₂CO₃, but redissolved in excess Oxidised by FeCl₃ to hemipic acid and cotarnine —B'HCl 2aq crystals —B₂H₂PtCl₄

NARINGIN C₂₁H₁₉O₁₁ *Aurantium Hesperidins* [171°] S 35 in the cold [α]_D = -84.5 in aqueous solution, -87.6 in alcoholic solution (W), [α]_D = -64.6 (H) Occurs in the flowers and other parts of *Citrus decumana* The dry orange-blossoms contain about 2 p c It crystallises on cooling from the liquor left in the still after distilling over the oils with steam (Hoffmann, Ar Ph [3] 14, 189, Will, B 18, 1311.

K K

20, 295) White crystals (containing 4aq), nearly insol cold water, sol alcohol, insol ether. Has a very bitter taste. Split up by dilute H_2SO_4 (8 p c) at 95° quantitatively into naringenin and isodulcitol. It dissolves in alkalis with a yellowish-red colouration. Ferric salts produce a brownish-red colouration with dilute aqueous solutions. By sodium-amalgam it is converted into a colouring matter which dissolves in alcohol with a red colour and bluish fluorescence.

Naringenin $C_{15}H_{12}O_6$, probably

[4 1] $C_6H_4(OH)CH=CHCOOC_6H_4(OH)$, [1 8 5] [248 γ] Formed, together with isodulcitol, by heating naringin with dilute (2-3 p c) H_2SO_4 on the water bath. Glistening colourless, tasteless, and odourless crystals. V sol alcohol, ether, and benzene. Dissolves in alkaline hydrates forming yellow solutions, and is reprecipitated by CO_2 . Ferric salts give a brown-red colouration. By sodium-amalgam it is converted into a colouring matter which dissolves in alcohol with a red colour and bluish fluorescence. By boiling with concentrated aqueous NaOH it is decomposed into phloroglucin and *p* coumaric acid (Will, *B* 18, 1322, 20, 297).

Naringenic acid is *p* COUMARIC ACID

NARTIC ACID v NARCOTINE.

NATALOIN v ALOIN

NEODYMIUM The name given by Auer von Welsbach to that constituent of didymium which yields rose coloured salts (v DIDYMIUM, vol II p 883, cf METALS, RARE, this vol p 240)

NEOSSIDINE v PROTEIDS, Appendix C.

NEOSSINE v PROTEIDS, Appendix C

NERIODORIN A bitter yellow powder, extracted by alcohol from the bark and wood of *Nerium odorum*, an Indian plant prescribed for leprosy (Greenish, *Ph* [3] 11, 873). It is sol water, insol chloroform. Its solution is ppd by tannin and by ammoniacal lead acetate, and reduces Fehling's solution.

Neriodorin A bitter resin, accompanying the preceding body, sl sol water and alcohol, v sol $CHCl_3$.

NEURIDINE $C_8H_{11}N_2$ A product of the putrefaction of flesh, appearing on the second day and disappearing about the fourth day (Brieger, *B* 16, 1187, 1405; Bocklisch, *B* 18, 86; Gautier, *Bl* [2] 48, 12). It also occurs in fresh human brain (Brieger, *J Th* 1884, 92). Gelatinous mass with disgusting smell, v e sol water, insol alcohol and ether. Very poisonous (G.) Its solution is ppd by $HgCl_2$ and by lead acetate. Yields di- and tri-methylamine on boiling with NaOHaq— $B^+H_2O_2$ needles, v. sol water, insol alcohol and ether— $B^+H_2PtCl_6$.

NEURINE $C_8H_9NO_2$, v e

$CH_3(OH)CH_2NMe_2OH$ *Choline Di-methyl-oxy-ethyl-amine methyl-hydroxide Tri-methyl-oxyethyl ammonium hydroxide Sinealme*

Occurrence—In cotton seeds (Böhm, *J pr* [2] 80, 87), in putrefying flesh (Gautier, *Bl* [2] 48, 18), in germinating pumpkin sprouts (E Schulze, *H* 11, 365), in the seeds of *Trigonella Fenum graecum* (Jahns, *B* 18, 2518), in the seeds of vetch, *Vicia sativa* (Schulze, *B* 22, 1827), in hops and beer (Griess a. Harrow, *O J* 47, 298, *B* 18, 717), in areca nuts (Jahns, *B* 23, 2972), in the fly agaric (Harnack, *J* 1876, 808), in herring brine (Bocklisch, *B* 18, 1928),

in ergot of rye (Brieger, *H* 11, 184); and in beet root juice (Lippmann, *B* 20, 8201)

Formation—1 By boiling the bile of pigs or oxen with baryta (Strecker, *C R* 52, 1270, *A* 123, 353, Dybrowsky, *J pr* 100, 163, Hüfner, *J pr* [2] 19, 802)—2 By boiling ox-brain with baryta (Liebreich, *A* 184, 29, Baeyer, *A* 140, 306)—3 By extracting yolk of egg with ether and alcohol, evaporating the extract, and boiling the residue with baryta (Diaconoff, *J* 1867, 776, 1868, 780). In this preparation two similar bases occur, containing 34 and 32 p c Pt in their platinumchlorides (Hundeshagen, *J pr* [2] 28, 247)—4 Together with glycerophosphoric acid, oleic, and palmitic acids by the action of alkalis on lecithin (Liebreich, Gobley, *C R* 70, 1297, Bokay, *H* 1, 157, 12, 148)—5 Together with sinapic acid and barium sulphocyanide, by heating sinapine sulphocyanide with baryta water (Claus a. Keesé, *Z* [2] 4, 46)

Syntheses—1 By heating trimethylamine (5 g) with glycolic chlorhydrin (10 g) at 100° , the resulting chloride $HOCH_2CH_2NMe_2Cl$ being decomposed by moist Ag_2O (Wurtz, *C R* 65, 1015, 68, 1434, *A Suppl* 6, 116)—2 By the union of ethylene oxide with trimethylamine (Wurtz, *A Suppl* 6, 201)

Properties—Strongly alkaline syrup, decomposed on boiling in concentrated aqueous solution into glycol and trimethylamine. Conc HIAq and P at 140° forms $C_8H_{11}NMe_2I$. Weaker HIAq yields $C_8H_{11}NMe_2Cl$. Yields muscarine and betaine on oxidation. Not poisonous.

Salts— $C_8H_{11}(OH)NMe_2Cl$ Dimorphous usually as long needles, sometimes in thin trimetric plates— $(C_8H_{11}(OH)NMe_2Cl)_2PtCl_6$. Trimorphous orange prisms (from warm saturated solution), reddish brown trimetric tablets from a cold saturated solution, or regular octahedra from a solution containing 15 p c of alcohol. The first and third forms left in contact with their mother liquid change into the second form (Hundeshagen, *J pr* [2] 28, 245)— $(C_8H_{11}(OH)NMe_2Cl)AuCl$, yellow needles, sl sol cold water— $C_8H_{11}(OH)NMe_2I$ Formed from $C_8H_{11}(OH)NMe_2$ and MeI (Knorr, *B* 22, 1116)

Acetyl derivative of the chloride $C_8H_{11}(OAc)NMe_2Cl$ — $C_8H_{11}NO_2ClAuCl$, nodules

NEURO-KERATIN v PROTEIDS, Appendix C
NICKEL At w. 58.6 Mol w unknown, as element has not been gasified [α 1400° — 1420°] (Schertel, *N* 22, 543), [α 1450°] (Pictet, *C R* 88, 1317) SG 8.97 to 9.26 (Rammelsberg, *J* 2, 282, for other values v Clarke's *Table of Specific Gravities* [new ed.], 12) SH 14° to 97° 10916 (Regnault, *A Ch* [3] 63, 1) $V_r = V_0 (1 + 3 \times 0.0001279t)$ $t = 40^\circ$ (Fizeau, *C R* 68, 1125) EC at 0° (Hg at $0^\circ = 1$) 7.374 (Matthiessen a. Vogt, *P M* [4] 26, 242) SVS c 6.5 H_2O [Ni^2+ , O^2 , $3H^+$]=120,380 (*Th* 3, 307) For spectrum of Ni (ultra violet) v Liveing a. Dewar, *Pr* 43, 430

Occurrence—In some meteorites, from 8 to 8 p c associated with 2 to 1 p c of Co. In the sun's atmosphere (Cornu, *C R* 86, 983) Ni has been found, to the extent of 75 p c, in a Pt ore from the Ural (Terrell, *C R* 82, 1116). Chiefly as copper-nickel Ni_2As_2 , white nickel $NiAs_2$, and Mg-Ni silicate, garnierite. Ni antimonide, arsenate, oxide, sulphide, carbonate, &c., also occur, chiefly in combination with com-

pounds of Co, Fe, Sb, and Bi Ni compounds are always present in Co ores, and Co is almost invariably a constituent of Ni ores. Ni compounds have been known to the Chinese for centuries, Ni forms a constituent of Chinese armour. Copper nickel was known to the German miners in the Middle Ages, having in vain attempted to extract Cu from this mineral, they gave it the name of *kupfer nickel*, or false copper. In 1751 Cronstedt showed the mineral to contain a special metal, to which he gave the name of nickel. The metal was obtained in impure condition by Cronstedt, and examined by Bergmann, Richter, and others. Fleitmann prepared larger quantities of Ni, and applied it to coat iron and steel. Bottger, c 1840, introduced the practice of electro nickel plating, this application of Ni has become an important trade since c 1869.

Formation—*Copper nickel*, or *speiss* (a by-product in the production of smalt), is powdered and roasted (to remove As and oxidise Ni), again roasted with charcoal, dissolved in HNO_3Aq , saturated with H_2S , and the filtrate is pptd by Na_2CO_3 , the Ni is separated from the Fe and Co in the pp by different processes (*v Preparation*), the Ni is pptd by alkali as $\text{NiO} \cdot x\text{H}_2\text{O}$, the pp is dehydrated by heat, and reduced by C or in H. An ammoniacal solution of NiNH_4S sulphate is sometimes decomposed by electrolysis.

Preparation—A salt of Ni, tolerably free from Co, is obtained by such a method as the following from arsenical Ni sulphide, or *speiss*, *speiss* is a deposit formed in the pots in which roasted Co arsenide, mixed with *copper nickel*, is fused with K_2CO_3 and quartz, in the preparation of smalt. The roasted ore, or *speiss*, is fused with chalk and fluorspar, the slag being poured off, the metal is powdered and roasted for a long time in a reverberatory furnace till fumes of As_2O_3 cease to come off, the product is dissolved in HClAq , the solution is diluted, bleaching powder is added to oxidise the Fe salts, and milk of lime to ppt Fe_2O_3 with As oxide, the filtrate is acidified and pptd by H_2S , the filtrate from this pp is heated with bleaching powder to ppt Co oxide, and the Ni remaining in solution is pptd as oxide by milk of lime (Louyet, *J Ph* [3] 15, 204, for other methods v Wagner, *Berg und Hüttenm Zeit* 1870 134, Gilchrist, *B* 16, 264, Donath, *D P J* 236, 327, Wöhler, *P* 6, 227, Cloez, *J* 1857 619, v also DICTIONARY OF APPLIED CHEMISTRY).

Impure Ni may be purified by dissolving in HClAq with addition of HNO_3 , evaporating to dryness, dissolving in water, acidulating with HClAq , boiling with excess of NaHSO_4 (to reduce As_2O_3 to As_2O_5), saturating the warm liquid with H_2S (after SO_2 is all removed), filtering after 12 hours or so, evaporating the filtrate to dryness, dissolving the residue in water, filtering, treating the filtrate with Cl and then ppg Fe and Co by BaCO_3 , removing excess of Ba by dilute $\text{H}_2\text{SO}_4\text{Aq}$, filtering, and ppg Ni as carbonate by Na_2CO_3 (Cloez, *J* 1857 619). After ppg As, Cu, Sb, Pb, and Bi by H_2S , the filtrate may be much concentrated and Ni pptd. as oxalate, by addition of oxalic acid (Deville, *A Ch* [8] 46, 82). Winkler (*Fr* 6, 18) boils a solution of commercial NiCO_3 with NaClO till all Co is pptd, this occurs only when a large quantity of Ni is also thrown down; he filters, ppts. by H_2S , boils the filtrate, and

ppts NiCO_3 by addition of Na_2CO_3 . Co is the most difficult impurity to separate from Ni salts; the most effectual method is to dissolve in HNO_3Aq , concentrate, neutralise by KOH Aq , mix with KNO_3Aq , strongly acidify with acetic acid, and allow to stand for a few days, when the Co is completely pptd as Co K nitrite, and the filtrate contains the Ni (*cf* Zimmermann, *A* 232, 324).

Ni is obtained from NiCO_3 (or $\text{NiO} \cdot \text{H}_2\text{O}$ pptd from solutions by KOH Aq) by washing thoroughly, spreading out to dry, heating till CO_2 is all removed, and then reducing by H, at c 270 (Müller, *P* 136, 51), or by heating with C, the latter method is conducted by making the NiO into a paste with oil, placing this in a crucible lined with charcoal, and heating in a powerful air- or blast furnace.

NiCO_3 may be dissolved in HClAq , the solution evaporated to dryness, the residue of NiCl_2 thoroughly dehydrated, then sublimed in a porcelain tube in a stream of Cl, and finally reduced by heating in H (Winkler, *Fr* 6, 18).

NiCl_2O yields Ni when heated out of contact with air, this may be done by heating under a layer of pounded glass which does not contain any heavy metal.

Ni is also obtained by electrolysis solutions of its salts, using a weak current, the best solution to use seems to be an ammoniacal one of NiNH_4S sulphate (*v* Becquerel, *C R* 55, 18, *cf* Zimmermann, *A* 232, 324).

Properties—A very lustrous metal, white, with a slight greyish yellow tinge. Hard, easily polished, ductile, malleable, and very tenacious. By heating in a porcelain oven, crystals of Ni, apparently regular, have been obtained. Slightly magnetic, but less so than Fe or Co. Ni obtained by reduction of NiO by charcoal usually contains a little C, such C-containing Ni is softer and less malleable than the purer metal (Boussingault, *Chem Ind* 1878 130). As obtained by reducing NiO by H, or by heating NiCl_2O , Ni forms a greyish black, somewhat porous, pyrophoric powder. As obtained by reducing NiCl_2 in H, the metal forms a compact sponge, and also lustrous leaflets (Winkler, *Fr* 6, 18, Gard, *J* 1877 266). By electrolysis of Ni solutions by a very weak current, Ni is obtained as coherent, lustrous, white plates (Becquerel, *C R* 55, 18). After fusion of large quantities of Ni, the metal becomes porous and crystalline, and loses its ductility, this is probably due to absorption of gases, and may be prevented by adding $\frac{1}{2}$ p c of Mg (Fleitmann, *B* 12, 454).

Ni shows *passivity*, i.e. under certain conditions it is unacted on by HNO_3Aq . According to Saint-Etienne (*C R* 106, 1079), commercial sheet Ni is passive in ordinary HNO_3Aq , and passive Ni remains passive when heated to bright redness in H, whereas Fe loses its passivity (*cf* *Passivity of iron*, under Iron, this vol p 52).

Ni is a metallic element. The oxides are basic, or react as peroxides, NiO forms a series of corresponding salts, Ni_2O_3 probably forms salts by reacting with acids, but these salts are reduced very easily to salts of NiO . Ni is closely related in its chemical properties to Co, it is classed with Co and Fe, and also shows analogies with Mn, v Iron group of elements, this vol. p. 65. No Ni salts are known correspond-

ing with the cobalto and cobalti-cyanides, M_2CoCy_6 and M_2CoCy_8 , nor are nickel-ammonio-bases known analogous with the cobaltamines, although some Ni compounds, *eg* the haloid compounds, combine with NH_3 . The ultra-violet spectra of Ni and Co do not exhibit much analogy (*v. Luveing a Dewar, Fr 43, 430*).

The at w of Ni has been determined (1) by reducing NiO in H (Erdmann a Marchand, *A* 82, 76, Russell, *C J* [2] 1, 51), (2) by analysing Ni oxalate (Schneider, *P* 101, 387, 107, 605, 180, 303), by determining Cl in NiCl₂ (Dumas, *A Ch* [3] 55, 149), (3) by ppg NiSO₄Aq by BaCl₂Aq (Sommaruga, *W A B* 54 [2] 57), (4) by the reaction of AuCl₃Aq on Ni, and determining the Au ppd (Winkler, *Fr* 6, 22, *cf* Kruss a Schmidt, *B* 22, 11, but *v* also Winkler, *B* 22, 890), (5) by decomposing by heat strychnine Ni cyanide and bromine Ni cyanide (Lee, *C N* 24, 287), (6) by dissolving Ni in HClAq, and measuring H evolved (Russell, *C J* [2] 7, 294), (7) by determining SH of Ni, the result obtained shows that 58.6, and not a multiple of this number, is the at w, this result has been confirmed recently, (8) by determining V D of gaseous Ni(CO)₄ (Mond, Langer, a Quincke, *C J* 57, 749).

Alleged decomposition of nickel—Kruss a Schmidt (*B* 22, 11) found that by repeatedly treating NiS, prepared from the ordinary sulphate, with NH_4 sulphide, until the latter was no longer coloured brown, a bright yellow residue was obtained, from this residue they prepared various salts, which they regarded as compounds of a new metal, and finally they obtained the supposed new metal by electrolysis an aqueous solution of the chloride, and also by reducing the dichloride in H. Kruss a Schmidt (*B* 22, 2026) prepared what they considered to be pure Ni, and by fractionally ppg this as basic Ni NH_4 arsenite they obtained two different substances, one having an at w 56.58, and the other an at w 61.100, hence they concluded that their former results were confirmed, that is, that Ni is really a compound or a mixture of two elements. Considerable doubt was thrown on these results by the work of Winkler on the reaction between pure AuCl₃ and Ni (*B* 22, 890), and Fleitmann's experiments on large quantities of Ni also tend to show that Ni has not been separated into unlike parts (*Chem Zeitung*, 18, 757).

Reactions and Combinations—1 Pieces of Ni do not oxidise in ordinary air, heated in air Ni is superficially oxidised to NiO, Ni obtained by reduction of NiO at low temperatures is pyrophoric—2 Ni burns to NiO when heated in oxygen—3 By heating in chlorine, bromine, or iodine, NiCl₂, NiBr₂, or NiI₂ is produced—4 Heated with sulphur, NiS is formed—5 Ni combines with phosphorus when the elements are heated together—6 Combines with arsenic in several proportions (*v Nickel, arsenides of*, p 501)—7 Absorbs, and probably also combines with, carbon (*v Nickel, carbides of*, p 501)—8 Combines with carbon monoxide (*v Nickel carbon-oxide*, p 501)—9 By reducing NiO in presence of SiO₂, Ni is obtained, containing silicon (*v Nickel, silicides of*, p 503)—10 Steam is slowly decomposed by Ni at red heat, NiO being formed (Regnault, *A Ch.* [3] 62, 852).

11 Nitric acid forms Ni(NO₃)₂, but Ni is passive in conc HNO₃Aq—12 Sulphuric and hydrochloric acids react very slowly with Ni (*v Tassier, C R* 50, 106)—13 Aqueous sulphurous acid is decomposed by Ni at c 200°, with production of Ni₂S₃ (Geitner, *A* 139, 354)—14 Carbon dioxide is reduced to CO by heating with Ni to redness (Bell, *C N* 23, 358)—15 Heated in carbon monoxide to 850°–450°, C is deposited and CO₂ produced (*v supra*, No 8, also *Nickel carbon-oxide*, p 501)—16 Forms alloys with several metals (*v Nickel, alloys of, infra*)—17 Occludes about 165 times its volume of hydrogen (Raoult, *C R* 69, 826).

Detection and Estimation—Ni compounds give a bead with borax which is violet when hot and reddish brown when cold in the oxidising flame, and in the reducing flame becomes opaque and grey from reduced Ni. Brown black NiS is ppd by alkali sulphides, insol dilute cold HClAq, somewhat sol yellow NH_4 sulphide, H_2S produces no pp in acid solutions, moist NiS ppd from cold solutions, oxidises rather easily in the air. Very small traces of Ni may be detected, in the absence of most other metals, by the rose-red colour produced by adding K₂CS₂Aq (Braun, *J* 1868 376). To detect small quantities of Ni in presence of Co, Papasogli places a piece of Zn in a solution of the double cyanides of the two metals if Ni is present a red colour is produced (*J* 1879 1055), Donath a Mayrhofer (*Fr* 20, 879) add excess of NaOH Aq then I, boil, treat the pp with NH_4 Aq and NH_4 ClAq, and test the solution by NH_4 HS (*v* also Jorisson, *Fr* 21, 208).

Ni may be estimated as NiO after ppn as NiO xH_2O by addition of KOHAq to a hot solution. For separation of Ni from Co *v Cobalt*, vol II p 218. Classen recommends the ppn of Ni as oxalate, which on heating out of contact with air gives NiO (*Fr* 16, 471, 18, 189, 386). Ni may also be estimated by electrolysis a solution of Ni NH_4 oxalate in presence of excess of NH_4 oxalate (*v Classen*, for description of apparatus *v Dittmar's Exercises in Quantitative Chemical Analysis* [Glasgow, 1887], *cf* Merrick, *C N* 24, 100).

Technical Applications—Alloys of Ni with Cu, and with Cu and Zn, are used for coinage in some countries, and for other purposes. These alloys are nearly white and hard. Many articles of iron and steel are now covered with a deposit of Ni. This covering is only very slightly acted on by ordinary air. *Nickel plating* is usually accomplished by electrolysis an ammoniacal solution of Ni NH_4 sulphate, using Ni as one of the electrodes, and the substance to be plated as the other (*v D P J* 201, 145, 206, 288, 211, 74, 212, 160, 219, 469, *v* also Adams, *C R* 70, 123, 137, Bequerel, *ibid* 70, 124, 137, 181, *v* also *DICTIONARY OF APPLIED CHEMISTRY*).

Nickel, alloys of An alloy of Ni with aluminium, approximately Al₃Ni, is obtained by melting together 8 parts Al with 3 parts dry NiCl₂ and 20 parts mixed KCl and NaCl, and treating with dilute HClAq (Michel, *A* 115, 102). Alloys of Ni with copper, and with copper and zinc, are used for coinage and other purposes under the names of *German silver*, *packfong*, &c. Ramsay (*C J* 55, 582) found that an amalgam

of Ni with *mercury* could be obtained in small quantities by electrolysis dilute solutions of Ni salts in contact with Hg

Nickel, ammonio-compounds of Some Ni compounds, *eg* NiCl_2 and NiBr_2 , combine with NH_3 , *v* *Nickel bromide, Nickel chloride, &c*

Nickel, antimonates of, $\text{Ni}(\text{SbO}_3)_2 \cdot 6\text{H}_2\text{O}$ and $\text{Ni}(\text{SbO}_3)_2 \cdot 12\text{H}_2\text{O}$, *v* Heffter, *P* 86, 446

Nickel, antimonide of NiSb occurs native as *breithauptite*

Nickel, arsenates of, *v* vol 1 p 309 $\text{Ni}_2(\text{AsO}_4)_2 \cdot 8\text{H}_2\text{O}$ occurs native as *nickel bloom*

Nickel, arsenides of A brittle compound, Ni_2As , is formed by heating Ni with excess of As, also by very strongly heating Ni arsenates in a charcoal-lined crucible (For other arsenides *v* Descamps, *C R* 86, 1065) Various Ni ores are essentially compounds of Ni and As, Ni_2As , = *spess*, NiAs = *copper-nickel*, Ni_3As , = *white-nickel*

Nickel, arsenite of, *v* vol 1 p 306

Nickel, borate of, *v* vol 1 p 530

Nickel, bromide of, NiBr_2 This, the only compound of Ni with Br, is obtained by passing Br vapour over finely divided Ni heated to low redness NiBr_2 forms a brownish-yellow solid, it sublimes in yellow, glittering scales (Berthelot, *A Ch* [3] 44, 389, Rammelsberg, *P* 55, 243) Heated in air, or in steam, NiBr_2 forms NiO , it is completely decomposed by HNO_3 Aqueous alcohol and ether, deliquescent, soluble water to a green liquid

The *hydrate* $\text{NiBr}_2 \cdot 3\text{H}_2\text{O}$ is obtained, in green needles, by digesting Ni with BrAq , also by dissolving NiO or NiCO_3 in HBrAq , and evaporating (Rammelsberg, *P* 55, 243) This hydrate is dehydrated at c 200° H F [Ni, Br, Aq] = 71,820 (*Th* 3, 307)

A compound of *nickel bromide with ammonia*, $\text{NiBr}_2 \cdot 6\text{NH}_3$, is obtained as a violet powder, by passing NH_3 over powdered NiBr_2 , also, as a blue powder, by warming conc NiBr_2Aq with excess of NH_3Aq , and cooling NH_3 is given off when the compound is heated $\text{NiBr}_2 \cdot 6\text{NH}_3$ is soluble in a little water without decomposition, much water causes ppn of $\text{NiO} \cdot \text{H}_2\text{O}$ (Rammelsberg, *lc*)

Nickel, carbides of Commercial Ni always contains a little C For experiments on carbonisation of Ni *v* Gard, *Am S* [3] 14, 274, Boussingault, *C R* 86, 509, Péal, *A* 233, 160, Gautier, *A Hallopeau*, *C R* 108, 1111, Mond, Langer, *A Quincke*, *C J* 57, 749

Nickel carbon-oxide Ni(CO) , Mond, Langer, *A Quincke* (*C J* 57, 749) found that Ni decomposes CO at 850° – 450° , with separation of C and formation of CO_2 . When the product was allowed to cool in CO they noticed that the escaping gas caused a Bunsen flame to become very luminous, and when heated deposited Ni Following up this observation they found that when finely divided Ni, produced by reducing the oxide in H, is allowed to cool in a slow current of CO the gas is readily absorbed by the Ni when the temperature has fallen to c 100° , and that by replacing the CO by CO_2 , N, H, or air, a mixture of gases is obtained which deposits Ni when heated above 150° By analysing the mixture of gases thus obtained, and determining the Ni by passing the gases through a capillary tube at 180° , M, L, *A. Q* found that one volume of

the Ni compound present in the gases gave four volumes of CO. The analyses led to the formula $\text{Ni}_2\text{C}_2\text{O}_4$. By passing the mixed gases through a tube surrounded with salt and ice a colourless mobile liquid was obtained, which was proved, by estimations of Ni and C, to be $\text{Ni}_2\text{C}_2\text{O}_4$. The V.D. determined at 50° was found to be 86.9, $\text{Ni}_2\text{C}_2\text{O}_4$ requires 80.4 The new compound is called *nickel carbon-oxide* by its discoverers, it boils at 43° at 751 mm, solidifies at -25° to needle-shaped crystals, and has S.G. 1.8185 at 17° . The vapour is very poisonous The compound is sol. alcohol, benzene, and chloroform, it is not acted on by dilute acids or alkalis, nor by conc HClAq ; conc HNO_3 and *aqua regia* dissolve it readily. The vapour ppts Ag from AgCl in NH_4Aq , it is decomposed by Cl, giving NiCl_2 and COCl_2 . Br acts similarly, electric sparks produce Ni and CO

Nickel, chloride of, NiCl_2 H F. [Ni, Cl] = 74,530 (*Th* 3, 307), [NiCl_2 , Aq] = 19,170 S.G. 2.56 (Schiff, *A* 108, 21) Prepared by gently heating powdered Ni in a stream of dry Cl, and subliming in the Cl (H Rose, *P* 20, 156) Also by dissolving Ni or NiCO_3 in HClAq , or Ni in *aqua regia*, and evaporating to dryness Golden yellow scales Sublimes readily without melting Prepared in the wet way, NiCl_2 is deliquescent and easily soluble in water, sublimed NiCl_2 dissolves slowly in boiling water Heated in air Cl is evolved and NiO formed Heated in a stream of O, is entirely changed to Ni_2O_3 (Schulze, *J pr* [2] 21, 407) KOH Aq decomposes sublimed NiCl_2 only after prolonged boiling PH_3 forms Ni_2P_3 , and HCl , molten P forms Ni_3P_2 , and PCl_3 (H Rose, *P* 27, 117) For S.G. of conc NiCl_2Aq *v* Franz, *J pr* [2] 5, 274

The *hexa hydrate* $\text{NiCl}_2 \cdot 6\text{H}_2\text{O}$ (Laurent, *A Ch* [3] 60, 354) is obtained by cooling conc NiCl_2Aq Soluble in 1½–2 parts water, sol in alcohol Sabatier (*Bl* [3] 1, 88) describes a *dehydrate* $\text{NiCl}_2 \cdot 2\text{H}_2\text{O}$, obtained by placing the hexahydrate over H_2SO_4 *in vacuo* at 20° for three months Thomsen gives [Ni, Cl, $6\text{H}^\circ\text{O}$] = 94,860 (*Th* 3, 307)

The *oxychloride* $\text{NiCl}_2 \cdot 8\text{NiO} \cdot 18\text{H}_2\text{O}$ is obtained by adding a little NH_3Aq to NiCl_2Aq (Rasch, *C R* 69, 826)

Compounds with ammonia (1) $\text{NiCl}_2 \cdot 6\text{NH}_3$, is obtained by passing NH_3 over dry NiCl_2 (H. Rose, *P* 20, 155), also by adding alcohol to NiCl_2 in NH_3Aq (Erdmann, *Gm.-K* (6th edit) 3, 553, F. Rose, *Gm. K* (6th edit) 8, 561). Soluble without change in cold water, decomposed slowly by much cold water, quickly by hot water Slightly sol conc NH_3Aq , insol alcohol. (2) $\text{NiCl}_2 \cdot 9\text{NH}_3$, obtained by heating $\text{NiCl}_2 \cdot 6\text{NH}_3$ to 120° Gives NiCl_2 when heated *in vacuo*, and a little Ni when very strongly heated

Compounds with ammonium chloride (1) $\text{NiCl}_2 \cdot \text{NH}_4\text{Cl} \cdot 6\text{H}_2\text{O}$, green deliquescent crystals, obtained by evaporating a solution of the constituents in the ratio $\text{NiCl}_2 \cdot \text{NH}_4\text{Cl}$ (Hantz, *A* 66, 288) Jørgensen (*Gm.-K* (6th edit) 8, 561) obtained crystals containing c. 5 p.c. NiCl_2 by evaporating NiCl_2Aq with a large excess of NH_4Cl (2) $\text{NiCl}_2 \cdot 4\text{NH}_4\text{Cl} \cdot 7\text{H}_2\text{O}$, yellow, star-shaped, crystals, obtained by decomposing Ni-NH_4 sulphate solution by an equivalent of BaCl_2 , filtering and evaporating over H_2SO_4 (Adams *A Meyrick*, *J* 1871 308, *cf* Tuppitt, *A. Ch* [3] 78, 169)

NiCl_2 form *double salts* with CrCl_3 , CdCl_2 , and

AuCl_3 (v. Goddefroy, *B* 8, 9, von Hauer, *W A B* 20, 40; *ibid W A B* 17, 848)

Nickel, chlorates of, v. vol u p 156

Nickel, cyanide and double cyanides of, v. vol u p 848

Nickel, ferrieyanide of, v. vol u p 339

Nickel, ferrocyanides of, v. vol u p 336

Nickel, fluoride of, NiF_2 , SG 2 855 at 14° (Clarke, *Am S* [3] 13, 291) The hydrate $\text{NiF}_2 \cdot 3\text{H}_2\text{O}$ is obtained by dissolving NiO H_2O or NiCO_3 in HFAq , and evaporating (Berzelius, Clarke, *Am S* [3] 13, 291) Decomposed by much hot water to oxyfluoride $\text{Ni}_2\text{OF}_2 \cdot \text{H}_2\text{O}$ (Berzelius) Combines with AlF_3 , fluorides of the alkali metals (Wagner, *B* 19, 896), and with SiF_4 (v. Nickel, silicofluoride of, p 503), TiF_4 (v. TITANIUM FLUORIDE), and ZrF_4 (v. ZIRCONIUM FLUORIDE) With Mo_2O_7 forms the compound $\text{NiF}_2 \cdot \text{Mo}_2\text{O}_7 \cdot 6\text{H}_2\text{O}$ (Delafontaine, *J* 1867 236)

Nickel, hydroxides or hydrated oxides of, v. Nickel, oxides and hydrated oxides of, *infra*

Nickel, iodide of, NiI_2 [Ni , I , Aq]=41,400 (*Th* 3, 807) Prepared by heating Ni , reduced from NiO by H , with I , and subliming the NiI_2 from the residue of Ni and NiO (Erdmann, *J* pr 7, 249) Also by heating $\text{NiI}_2 \cdot 6\text{H}_2\text{O}$, which is obtained by dissolving NiO H_2O in HIAq , or by treating finely divided Ni with excess of I and water Iron black, lustrous, metal like, scales Deliquescent, soluble in water, forming a brown liquid, which becomes green on dilution Partly decomposed when strongly heated in air By digesting NiI_2 Aq with NiO H_2O , or by evaporating NiI_2 Aq , Erdmann (*lc*) obtained the oxyiodide $\text{NiI}_2 \cdot 9\text{NiO} \cdot 15\text{H}_2\text{O}$

Compounds with ammonia (1) $\text{NiI}_2 \cdot 4\text{NH}_3$, a yellow-white mass, by passing NH_3 over NiI_2 (Rammelsberg, *P* 48, 119) (2) $\text{NiI}_2 \cdot 6\text{NH}_3$, by adding excess of NH_3 Aq to conc NiI_2 Aq , warming, and cooling or adding alcohol (Erdmann, *lc*, Rammelsberg, *lc*)

Nickel, nitride of A compound of Ni with N is said to be formed by heating NiO to c 200° in NH_3 , it is decomposed at a higher temperature (Warren, *C N* 56, 155)

Nickel, oxides and hydrated oxides of Nickel forms three oxides, NiO , Ni_2O_3 , and Ni_3O_4 , a fourth oxide, Ni_4O_5 , probably exists, there are indications of the existence of oxides intermediate between Ni_2O_3 and Ni_3O_4 , and also of an oxide containing more O than Ni_3O_4 Hydrates of NiO and Ni_2O_3 , and perhaps of Ni_3O_4 , have been isolated The oxides of Ni are basic, but the only Ni salts which have been prepared with certainty correspond with NiO NiO is oxidised by heating to c 400°, but the product is reduced to NiO at c 600°

NICKEL MONOXIDE NiO (*Nickelous oxide Protoxide of nickel*) Occurs native as *bunsenite* Obtained by heating Ni or NiCl_2 in steam (Regnault, *A Ch* [3] 62, 352), by heating the hydrate or NiCO_3 in absence of air, by strongly heating $\text{Ni(NO}_3)_2$ (Russell, *C J* [2] 1, 58), NiSO_4 (Baubigny, *C R* 97, 951), or a mixture of NiSO_4 and K_2SO_4 (Debray, *C R* 52, 985), by reducing Ni_2O_3 by H at 190°–230° (Moissan, *A Ch* [5] 21, 288, cf Wright a Luff, *O J* 33, 1, also Müller, *P* 136, 59), or by NH_3 at c 180° (Vorster, *Dissertation*, Göttingen, 1861), at a moderate temperature NiO is obtained in green regular octahedra by heating Ni borate with CaO

in a porcelain oven, and treating the product with HClAq (Ebelmen, *C R* 33, 526)

NiO is a green powder, becoming deep yellow when heated (Moissan, *A Ch* [5] 21, 238, Zimmermann, *A* 232, 324) SG 5 6 (Playfair a Joule, *C S Mem* 3, 81), 6 66 (Rammelsberg, *J* 2, 282), 6 8 crystallised (Ebelmen, *C R* 33, 526) Heated to 350°–440°, NiO is oxidised to Ni_2O_3 , which is again deoxidised to NiO at c 600° (Moissan, *A Ch* [5] 21, 199) NiO is readily reduced to Ni , reduction by CO begins at c 120°, by H at c 220°, by C at c 450° (Wright a Luff, *C J* 33, 1), reduction by NH_3 begins at c 200° (Vorster, *Dissertation*, Göttingen, 1861) NiO is oxidised to Ni_2O_3 by ozonised O (Schönbein, *J pr* 93, 35) NiO , if not strongly heated, dissolves in NH_3 Aq , it decomposes NH_4 salts with evolution of NH_3 NiO dissolves in acids forming salts NiX_2 , $\text{X} = \text{NO}_3$, ClO_4 , $\frac{1}{2}\text{SO}_4$, $\frac{1}{3}\text{PO}_4$, &c

HYDRATE OF NICKEL MONOXIDE $4\text{NiO} \cdot 5\text{H}_2\text{O} = 4\text{Ni(OH)}_2 \cdot \text{H}_2\text{O}$ (*Nickelous hydrate, nickelous hydroxide*) This hydrate is obtained as a pale greenish pp, by adding alkali solution to solution of a Ni salt According to Teichmann (*A* 156, 17) the hydrate can be obtained free from acid only from $\text{Ni(NO}_3)_3$ Aq , T recommends to add NaOH Aq , free from carbonate, to cold $\text{Ni(NO}_3)_3$ Aq , to wash the excess of pp with cold water till the alkaline reaction disappears, then to wash with water containing a little NH_3 , and finally with boiling water, and to dry at 100° $4\text{NiO} \cdot 5\text{H}_2\text{O}$ is also obtained, as a green crystalline powder, by heating NiO or NiCO_3 in NH_3 Aq (*Gm K* (6th edit) 3, 536)

Nickelous hydrate is a pale green powder, slightly soluble in water (Fresenius) If not washed free from alkali it oxidises in presence of air and SO_2 Aq , but the product is reduced to NiO by excess of SO_2 Aq (Wicke, *Z* 1865 86) When strongly heated, H_2O is evolved and NiO remains $\text{H}_2\text{O Aq}$ forms a hydrate of Ni_2O_3 (Schönbein, *J pr* 93, 35), but, according to Bayley, $\text{H}_2\text{O Aq}$ is without action (*P M* [5] 7, 126) Oxidised by Cl and hypochlorites to $\text{Ni}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ Soluble in NH_3 Aq and solutions of NH_4 salts $4\text{NiO} \cdot 5\text{H}_2\text{O}$ reacts with acids as a strong base, forming salts NiX_2 , $\text{X} = \text{NO}_3$, $\frac{1}{2}\text{SO}_4$, $\frac{1}{3}\text{PO}_4$, &c Thomsen gives $[\text{NiO} \cdot \text{H}^2, \text{H}^2\text{SO}_4 \text{Aq}] = 26,110$, and $[\text{NiO} \cdot \text{H}^2, 2\text{HClAq}] = 22,580$ (*Th* 8, 307)

NICKEL SESQUIOXIDE Ni_2O_3 (*Nickelic oxide* Sometimes called *nickel peroxide*) A black powder, SG 4 84 at 16° (Herapath, *P M* 64, 321) Obtained by decomposing by heat at the lowest possible temperature $\text{Ni(NO}_3)_2$ (Berzelius, Vorster, *Dissertation*, Göttingen, 1861), or $\text{Ni(ClO}_4)_2$ (Wachter, *J pr* 30, 327), also by melting NiCl_2 with KClO_4 (Schulze, *J pr* [2] 21, 407), also by treating Ni salts in solution with KClO_4 Aq or KBrO_3 Aq (Schröder, *C C* 1890 931) Heated in air to c 600° Ni_2O_3 is reduced to NiO (Moissan, *A Ch* [5] 21, 199), reduction in H begins at c 190° (*M, lc*) Dissolves in H_2SO_4 Aq or HNO_3 Aq with evolution of O , in HClAq with evolution of Cl_2 , in each case salts of NiO are produced Soluble in NH_3 Aq with evolution of N (Müller, *P* 136, 59)

HYDRATES OF NICKEL SESQUIOXIDE.

(1) $\text{Ni}_2\text{O}_3 \cdot 2\text{H}_2\text{O}$; brownish crust, SG 2 744, obtained by electrolysis an alkaline solution of

Ni K tartrate (Wächter, *J pr* 80, 827) (2) $\text{Ni}_2\text{O}_3 \cdot 8\text{H}_2\text{O}$, by oxidising NiO or $4\text{NiO} \cdot 5\text{H}_2\text{O}$, suspended in water, by ClO or BrAq , also by adding alkali and NaClO to solution of a Ni salt (Wächter, *lc*) A black solid, which reacts with acids and NH_4Aq as Ni_2O_3 does, reduced to $4\text{NiO} \cdot 5\text{H}_2\text{O}$ by SO_2Aq (Wicke, *Z* 1865 86), also by $\text{Na}_2\text{SO}_3\text{Aq}$ (Schulze, *J* 1864 270) HF [$\text{Ni}_2\text{O}_3 \cdot 8\text{H}_2\text{O}$] = 120,380 (*Th* 3, 307) Carnelley & Walker (*C J* 53, 91) think that no definite stable hydrate of Ni_2O_3 exists

NICKEL O NICKELIC OXIDE Ni_2O_3 A grey, metal like, non magnetic solid, obtained by passing O over NiCl_2 at 350° – 440° (Baubigny, *C R* 87, 1082)

NICKEL O NICKELIC HYDRATE A black powder, having the composition $\text{Ni}_2\text{O}_3 \cdot \text{H}_2\text{O}$ (= $\text{Ni}_2\text{O}_3 \cdot 6\text{NiO} \cdot \text{H}_2\text{O}$), is said to be formed by heating NiCO , to 300° (H Rose, *P* 84, 571)

NICKEL SUBOXIDE An oxide, Ni_2O , is said to be produced by reducing NiO in H at 210° – 214° (Muller, *P* 186, 59), also by reducing NiO in CO at a low temperature (Bell, *C N* 23, 258, 267)

NICKEL P FROXIDE By the reaction of hypochlorites on $\text{Ni}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$ one or more oxides are produced containing more O than Ni_2O_3 Wicke gives the composition Ni_2O_7 (*Z* 1865 303), Bayley (*C N* 39, 81) gives the formula Ni_2O_8 , Carnot (*C R* 108, 610) says that Ni salt solutions give Ni_2O_8 when treated with hypochlorites or with Br and KOH Aq

Nickel, oxychloride of, *v* Nickel chloride, p 501

Nickel, oxyfluoride of, *v* Nickel fluoride, p 502

Nickel, oxyiodide of, *v* Nickel iodide, p 502

Nickel, phosphides of Ni and P combine when heated together Various compounds have been described (1) Ni_3P_2 , by heating Ni , bone ash, quartz sand, and C (*Gm K* (6th ed) 3, 542) (2) Ni_2P , by reducing $5\text{NiO} \cdot \text{P}_2\text{O}_5$ in H (Struve, *J* 1860 76) (3) Ni_3P_2 , by reducing $3\text{NiO} \cdot \text{P}_2\text{O}_5$ in H (H Rose, *P* 24 332), also by heating NiCl or NiS in PH_3 , or by passing PH_3 over heated Ni (Davy, Schrotter, *W A B* 2, 304)

Nickel, salts of Compounds obtained by replacing H of acids by Ni The Ni salts which have been studied all correspond with the oxide NiO , and belong to the form NiX , where $\text{X} = \text{ClO}_4$, NO_3 , $\frac{1}{2}\text{SO}_4$, $\frac{1}{2}\text{CO}_3$, $\frac{1}{2}\text{PO}_4$, &c The oxide Ni_2O_3 probably forms salts, but they are very easily reduced to salts of NiO The Ni salts are obtained by dissolving Ni , NiO , or NiCO in acids The salts of Ni are generally yellowish when dehydrated, and green when combined with water Some of the compounds of Ni combine with NH_3 The haloid compounds, the sulphate and nitrate of Ni , and some of the other salts, are soluble in water, the oxides, sulphides, phosphate, carbonate, and a few other salts, are insoluble in water Solutions of Ni salts in water are green, they redden litmus slightly Most Ni salts are decomposed by heating in air, NiCl_2 , NiBr_2 , and NiI_2 can be sublimed unchanged The chief Ni salts of oxyacids are the antimonate, arsenate and -ite, borate, bromate, carbonate, chlorate, chromate, iodate and periodate, molybdates, nitrates and -ite, phosphates and -ite and hypophosphite, selenate, silicates, sulphates

and -ite, thiosulphate (*v* CARBONATES, NITRATES, &c)

Nickel, selenide of, NiSe A silver white, brittle, crystalline, solid, *SG* 846, obtained by action of Se vapour on finely powdered Ni Melts at red heat, with loss of Se Insoluble in HCl Aq , slowly dissolved by HNO_3Aq , quickly by *aqua regia* (Little, *A* 112, 211)

Nickel, silicides of Commercial Ni generally contains more or less Si For experiments on the quantity of Si taken up by Ni v Gard, *Am S* [3] 14, 274

Nickel, silicofluoride of, $\text{NiSiF}_6 \cdot 6\text{H}_2\text{O}$ Hexagonal rhombohedral, green crystals, *SG* 2109, by dissolving NiCO in $\text{H}_2\text{SiF}_6\text{Aq}$ De composed at red heat, giving NiF and SiF_4 (Berzelius, Mangnac, *Ann M* [5] 15, 262)

Nickel, sulphides of Four sulphides of Ni are known Ni_2S , NiS , Ni_3S_2 , and NiS_2 NiS is somewhat soluble in NH_3 sulphide, it also forms a compound with K_2S The only sulphide produced by the direct union of Ni and S is NiS

NICKEL MONOSULPHIDE NiS Occurs native as *capillary pyrites* or *millierite* Formed by heating Ni with S , by heating NiO with S , or in a stream of H_2S (Tapput, *A Ch* [3] 78, 133, 79, 153), also by heating NiCl_2 with K_2SAq in a sealed tube to c 160° (Sénarmont, *A Ch* [3] 30, 142) As prepared by these methods, NiS is a yellow, brittle, solid, decomposed very slowly by steam at red heat (Regnault, *A Ch* [3] 62, 280), not decomposed by H , slowly acted on by Cl when hot (*v* *P* 42, 540), oxidised by heating in air, acted on by PH_3 , when hot, with formation of Ni_3P_2 (Schrötter, *W A B* 2, 304), insol HCl Aq , sol HNO_3Aq and *aqua regia*

NiS , in combination with water, is ppd from neutral Ni solutions by H_2S , or by NH_4HSAq , also by heating Ni salts with $\text{Na}_2\text{S}_2\text{O}_3\text{Aq}$ preferably in sealed tubes at c 120° (Gibbs, *Am S* [2] 37, 346) Thomsen gives [$\text{Ni}_3\text{S}_2 \cdot \text{H}_2\text{O}$] = 19,400 (*Th* 3, 307) The pp thus obtained is brown black, if ppd from boiling solutions it may be washed and dried without change, but if ppd from cold solutions it oxidises in the air (Clermont, *C R* 84, 714, 85, 73) Decomposed by boiling with water (Geitner, *A* 139, 354) Some what soluble in NH_4Aq and alkali sulphide solutions, the brown solutions thus obtained deposit NiS by standing in air or on addition of a weak acid According to Baubigny (*C R* 94, 1417) the pp produced by H_2S in neutral solutions of Ni salts is a hydrosulphide which is decomposed to NiS and H_2S by filtration A very dilute aqueous solution of a *colloidal form* of NiS was obtained by Winsinger (*Bl* [2] 49, 452) by ppg from a very dilute solution and dialysing

Compound with potassium sulphide, $3\text{NiS} \cdot \text{K}_2\text{S}$ A yellow, lustrous, crystalline solid, obtained by fusing NiSO_4 with K_2CO_3 .

NICKEL DISULPHIDE NiS_2 A dark iron grey powder, obtained by strongly heating NiCO_3 with K_2CO_3 and S , and treating with water (Fellenberg, *P* 50, 75)

NICKEL O NICKELIC SULPHIDE Ni_2S_3 An amorphous greyish-black solid of this composition is obtained by heating NiCl_2Aq with polysulphides of K to 160° (Sénarmont, *A Ch* [8] 80, 142), by heating Ni with SO_2Aq or $\text{Na}_2\text{SO}_3\text{Aq}$ to 200° , Ni_2S_3 is obtained in rhombohedral crystals (Geitner, *A* 139, 354)

tufts, v sol water (Dreser, *Ar Ph* [3] 27, 266) — Nitroprusside sol water (Davy, *Ph* [3] 11, 756) — Chloro-iodide, dark-yellow pp (Dittmar, *B* 18, 1612) — Picrate $B''2C_6H_3(NO_2)_3OH$ [218°] Yellow needles (P a W)

Methylo iodide $B''Me_2I$ (Stahlschmidt, *A* 90, 222) With moist Ag_2O it yields a caustic base. It yields the salts $B''Me, PtCl_3, B''2MeAuCl_3$, and $B''Me_2Cl_4HgCl_2$. When the methylo-iodide is treated with alcoholic KOH at 45° it gives a ruby red colouration, on addition of acids the colour remains red, and, on pouring into a large quantity of water, shows a green fluorescence (O De Coninck, *C R* 104, 1374)

Ethylo-iodide $B''Et_2I$, prisms, v sol water, sl sol alcohol and ether (Von Planta a Kekulé, *A* 87, 2) Yields with Ag_2O a caustic base. It forms the crystalline salts $B''Et, PtCl_3$, and $B''2EtAuCl_3$, and amorphous $B''Et, Cl_3HgCl_2$. An alcoholic solution of the ethylo iodide is coloured garnet red, and finally crimson, by heating with addition of potash for 10 hours on a water bath. After acidifying and pouring into water the colour is still red (De Coninck, *C R* 104, 513)

Isoamylo iodide $B''2C_5H_{11}I$ Yields $B''(C_5H_{11}), PtCl_3$ (Stahlschmidt)

Nicotine dihydride $C_{10}H_7N_2$ *Hydronicotine* (264°) $SG \pm 993$ $[a]_D = -15.40^\circ$ in a 13.7 p c solution. Formed by heating nicotine with fuming $HIAq$ and red phosphorus at 265° for 10 hours (Etard, *C R* 97, 1218) Liquid, with faint odour, miscible with water, alcohol, and ether. Lævorotatory. Its hydrochloride is not ppd by $HgCl_2$. — $B''H_2PtCl_3aq$ pale-yellow crystals, v sl sol water

Isonicotinic v DIPYRIDYL TETRAHYDRIDE

NICOTINIC ACID v PYRIDINE CARBOXYLIC ACID

Homo nicotinic acid v DI METHYL PYRIDINE CARBOXYLIC ACID

NIGRANILINE v ANILINE BLACK

NIGROSINE A name used by Wolff (*Chem Ind* 2, 290, 319) to denote a blue black substance $C_{26}H_{12}N_8$, found among the products of the action of arsenic acid on aniline hydrochloride at 230°. Its hydrochloride $C_{26}H_{12}N_8HCl$ exhibits in solution blood red fluorescence, and is decolourised by reducing agents. The name 'nigrosine' has also been applied to indulines, more especially when obtained by the action of nitro benzene on a mixture of pure aniline and aniline hydrochloride

NIOBATES v p 506

NIOBIMUM *Nb* (*Columbium*) At w 94 Mol w unknown $SG 7.06$ at 15.5° (Roscoe, *C N* 37, 26)

Occurrence — Niobates occur in a few rare minerals, e.g. in *columbite*, *tantalite*, *samaraskite*, *ytro-ilmenite*, *euxenite*, and some varieties of *pitch blende*. Niobates are generally accompanied by tantalates, tungstates, titanates, zirconates, and compounds of Th, Ce, and Yt

History — In 1801, Hatchett found a new oxide in a mineral called *columbite* from Massachusetts, to the metal of the new oxide he gave the name *columbium* (*Crell's Ann* 1, 197, 257, 352). In the following year, Ekeberg (*Scher J.* 9, 597) examined two minerals—one from Finland, the other from Sweden—and announced the discovery of a new oxide, as the oxide was

soluble only in caustic alkalis and was ppd by acids. Ekeberg gave to the metal of this oxide the name *tantalum*. In 1809, Wollaston (*S.* 1, 520) pronounced the oxides discovered by Hatchett and Ekeberg, respectively, to be identical. Berzelius confirmed the decision of Wollaston (*P* 4, 6), he proposed to apply the name *tantalum* to the characteristic metal of *columbite* and the minerals examined by Ekeberg. Oxide of tantalum was recognised as present in several rare minerals (v Hermann, *J pr* 38, 91, H Rosa, *P* 63, 321). In 1844, H. Rose began a series of researches on the minerals containing tantalum compounds (*P* vols 63, 69, 73, 74, 90, 99, 100, 101, 102). Rose concluded that oxides of three distinct metals exist in these minerals: oxide of tantalum in *tantalite* from Finland and Sweden, and oxides of two new metals, which he called niobium and pelopium, in *tantalite* from Massachusetts (formerly called *columbite*) and in a *tantalite* from Bavaria. In 1853, Rose came to the conclusion that the compounds described by him as oxides of niobium and pelopium were really two different oxides of the same metal which was different from tantalum, this metal Rose called niobium (from Niobe, the daughter of Tantalus) (Rose, *P* 63, 317). As different specimens of niobium oxide showed considerable differences of SG, Margnac re examined the ground, and showed that some of Rose's oxides of niobium contained tantalum (*C R* 60, 234, 1355). Margnac also showed that the most probable formulae for the oxide and chloride of Nb are Nb_2O_5 and $NbCl_5$, respectively. Blomstrand (*J pr* 37, 57) confirmed Margnac's results. Determinations of SG of gaseous Nb chloride and oxychloride by Deville a Troost (*C R* 56, 891) have shown the formulae $NbCl_5$ and $NbOCl_3$ to be molecular. H. Rose supposed he had obtained Nb by reducing a compound of Nb, K, and F by Na, Delafontaine showed that Rose's supposed Nb was really NbO , and that the compound from which it was obtained contained O (*Ar Sc* 27, 167). Blomstrand obtained Nb, containing some H, in 1864 by reducing the chloride in H, in 1878 Roscoe prepared approximately pure Nb by the same method (*C N* 37, 25)

The existence of three other metals in niobium containing minerals has been asserted by von Kobell (*J pr* 79, 291, 83, 193, 449), and Hermann (*J pr* 38, 91, 119, *J pr* [2] 3, 373, 4, 178, 15, 105), but the researches of Blomstrand and Margnac (*l c*) make the existence of these metals—*dromium*, *ilmenum*, and *neptunium*—very doubtful

Preparation — Very finely powdered *columbite* is fused with 3 times its weight of $KHSO_4$ in an iron or Pt crucible, until completely dissolved, after cooling, the residue is powdered and treated with boiling water, whereby sulphates of K, Fe, and Mn are removed, the insoluble portion is washed, and digested with yellow NH_3 sulphide, sulphides of Sn and W thus go into solution and FeS remains mixed with Nb_2O_5 and Ta_2O_5 , the residue is washed and digested with $HClAq$ to remove FeS, the insoluble in acid is thoroughly washed with boiling water until white. To separate Nb from this mixture of Nb_2O_5 and Ta_2O_5 , the whole is dissolved in $HFAq$, the solution is heated to boiling, and

$\frac{1}{2}$ part KHF_2 is added for each part of mixed Nb_2O_5 and Ta_2O_5 used, the liquid is evaporated until $\frac{1}{2}$ g of the mixed oxides is present in about 7 cc, and allowed to cool, crystals of K_2TaF_7 separate, these are washed with cold water till the washings give no red, but a pure yellow, pp with tincture of galls, the filtrate is concentrated with addition of KHF_2 , and the second crop of K_2TaF_7 crystals is removed and washed. After one or two repetitions of this process, fine tablets of $\text{NbOF} \cdot 2\text{KF}$ separate on evaporating the filtrate from the K_2TaF_7 crystals, the tablets are collected, pressed, and heated in a Pt dish with H_2SO_4 until HF is completely removed, the residue is boiled with a large quantity of water for some time, when a white pp of $\text{Nb}_2\text{O}_5 \cdot x\text{H}_2\text{O}$ separates out (Berzelius, *v* also Marignac, *Ar Sc* 23, 167, 249, 25, 5). The pp of $\text{Nb}_2\text{O}_5 \cdot x\text{H}_2\text{O}$ is washed, dried, and heated to redness, it is then mixed with a large excess of charcoal, the mixture is heated, then placed in a large hard glass tube (a small quantity being used, as NbCl_5 is very voluminous), heated in dry CO_2 until perfectly dry, and allowed to cool in dry CO_2 , the CO_2 is then completely expelled by dry Cl, and the tube is then heated to redness while dry Cl passes through it, NbCl_5 collects in the tube, and is distilled in a stream of dry Cl. The NbCl_5 is then vapourised in a current of perfectly dry H, with precautions to prevent the entrance of air and moisture, and the mixed vapour is passed through a red hot tube of hard glass. The grey, lustrous crust of Nb which forms in the tube is finally strongly heated in a stream of perfectly dry H (Roscoe, *C N* 37, 25). Nb thus prepared contains about 27 p c H.

Properties and Reactions — A steel grey lustrous metal. SG 7.06 at 15.5° (Roscoe, *lc*). Insol HClAq , HNO_3Aq , or *aqua regia*, sol conc H_2SO_4 . Heated in air, burns to Nb_2O_5 . Heated in Cl forms NbCl_5 .

The at w of Nb has been determined (1) by determinations of V D of NbCl_5 and NbOCl_3 (Deville & Troost, *C R* 56, 891, 60, 1221), and by analyses of NbCl_5 (H Rose, *P* 104, 432, Blomstrand, *Acta Univ Lund* 1864, Marignac, *Bibl Univ Genève*, 1866 and 1866), (2) by analyses of $\text{NbOF} \cdot 2\text{KF}$ aq (Marignac, *lc*).

Nb is metallic in its physical properties. NbO and Nb_2O_5 dissolve in conc H_2SO_4 , but no definite sulphates or other salts of the oxides have been isolated. $\text{Nb}_2\text{O}_5 \cdot x\text{H}_2\text{O}$ forms several niobates, in which Nb forms part of the negative radicle. Nb forms the third member of the even-series family of Group V, it is closely related to Ta, and less closely to N, P, V, As, Sb, Bi, Er, and Bi (*v* NITROGEN GROUP OF ELEMENTS, this vol p 571).

Detection and Estimation. — Niobates dissolve in hot HClAq , on adding water and boiling, $\text{Nb}_2\text{O}_5 \cdot x\text{H}_2\text{O}$ ppts. Solutions in HClAq are coloured blue, then dark brown, by Zn. $\text{K}_2\text{FeCy}_2\text{Aq}$ gives a red pp, and $\text{K}_2\text{FeCy}_2\text{Aq}$ a bright-yellow pp, with aqueous solutions of alkali niobates, gall tincture gives an orange-red pp. Nb is estimated as Nb_2O_5 , the process is sufficiently described under *Preparation* (cf. Rammelsberg, *P* 186, 177, 362, 144, 56, 191).

Niobium, acids of, and their salts Niobic oxide, Nb_2O_5 , reacts with alkali oxides to form salts, these niobates may be regarded as derived

from various hydrates of Nb_2O_5 . Hydrated niobic oxide, $\text{Nb}_2\text{O}_5 \cdot x\text{H}_2\text{O}$, is obtained by fusing Nb_2O_5 with KHSO_4 , washing with water, dissolving in HClAq , and ppg by NH_4Aq , the pp thus obtained by Santesson (*Bl* [2] 24, 52) contained c 84 p c water, which corresponds with the composition $3\text{Nb}_2\text{O}_5 \cdot 4\text{H}_2\text{O}$ ($=\text{Nb}_2\text{O}_{11}(\text{OH})_4$). The hydrate $\text{Nb}_2\text{O}_5 \cdot 7\text{H}_2\text{O}$ is obtained, according to Santesson (*lc*), by reacting on NaNbO_3 with $\text{H}_2\text{SO}_4\text{Aq}$ and drying at 100° (*v* *Hydrates of niobic oxide*, p 509). Niobates have not been obtained by neutralising hydrates of Nb_2O_5 , but either by fusing Nb_2O_5 with basic oxides or carbonates, or by double decomposition from solutions of alkali niobates.

NIOBATES The niobates belong to the form $x\text{Nb}_2\text{O}_5 \cdot y\text{MO}$, where $\text{M} = \text{K}, \text{Ca}, \text{Mg}, \text{Mn}, \&c$. Niobates are known corresponding with the meta- and pyrophosphates, and, besides these, salts have been isolated in which the ratio of the basic to acidic oxide varies from 1.2 to 4.1. The niobates are prepared by fusing Nb_2O_5 with basic oxides, carbonates, and a few other salts, some niobates are obtained by ppg solutions of alkali niobates by solutions of metallic salts. The alkali niobates are soluble in water, the others are insoluble. Solutions of the alkali niobates are decomposed by $\text{H}_2\text{SO}_4\text{Aq}$ with ppg of $\text{Nb}_2\text{O}_5 \cdot x\text{H}_2\text{O}$, CO, ppts acid salts. Solutions of niobates in HClAq are reduced by Zn to Nb_2O_5 (blue), and then to Nb_2O_3 (brown black) (*v* *Niobium oxides*, p 508). *Fluoniobates* and *fluozyniobates* are also known (*v* next page).

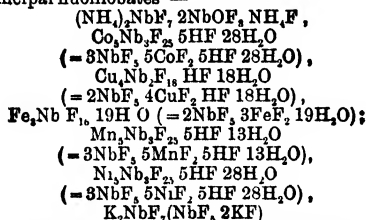
Potassium niobates (1) *Metaniobate*, KNbO_3 . Small rectangular tablets, sol water, obtained by dissolving Nb_2O_5 in molten CaF_2 , fusing the product with K_2CO_3 , in ratio $\text{K}_2\text{CO}_3 \cdot \text{Nb}_2\text{O}_5$, and repeatedly treating the mass (after cooling) with boiling dilute $\text{H}_2\text{SO}_4\text{Aq}$ (Joly, *Fremy's Encyclop Chmique*). (2) *Pyroniobate*, $\text{K}_2\text{Nb}_2\text{O}_7$. Insol water, obtained by melting Nb_2O_5 with a large excess of K_2CO_3 , and washing with water (Santesson, *Bl* [2] 24, 52). (3) $3\text{Nb}_2\text{O}_5 \cdot 4\text{K}_2\text{O} \cdot 16\text{H}_2\text{O}$, and (4) $7\text{Nb}_2\text{O}_5 \cdot 8\text{K}_2\text{O} \cdot 32\text{H}_2\text{O}$. The former salt is obtained by fusing Nb_2O_5 with 2 to 3 times its weight of K_2CO_3 , dissolving in water, and evaporating *in vacuo*, large monoclinic crystals, efflorescent in air, loses $12\text{H}_2\text{O}$ at 100° , and is dehydrated at red heat. The second salt is obtained in quad ratio octahedra by slowly evaporating a solution of the first salt (Marignac, *A Ch* [4] 8, 5, 13, 5). (5) $2\text{Nb}_2\text{O}_5 \cdot 3\text{K}_2\text{O} \cdot 18\text{H}_2\text{O}$, rhombic pyramids, by adding KOH to solution of salt (3) or (4), and evaporating slowly (Marignac, *lc*). (6) $2\text{Nb}_2\text{O}_5 \cdot 2\text{K}_2\text{O} \cdot 11\text{H}_2\text{O}$, the crystalline residue obtained by fusing Nb_2O_5 and K_2CO_3 , in the ratio $\text{Nb}_2\text{O}_5 \cdot \text{K}_2\text{CO}_3$, and treating with water, has this composition (Santesson, *Bl* [2] 24, 52). (7) $4\text{Nb}_2\text{O}_5 \cdot 3\text{K}_2\text{O}$, obtained by strongly heating Nb_2O_5 with twice its weight of KHSO_4 for some hours, and washing with water (Joly, *Fremy's Encyclop Chmique*). (8) $3\text{Nb}_2\text{O}_5 \cdot \text{K}_2\text{O} \cdot 5\text{H}_2\text{O}$, prepared by boiling $\text{KNbOF} \cdot 2\text{KFAq}$ with KHCO_3 , washing the powder which separates, and drying at 100° (Marignac, *lc*).

Sodium niobates (1) *Metaniobate*, $2\text{NaNbO}_3 \cdot 6\text{H}_2\text{O}$. Rhombic prisms, obtained by fusing Nb_2O_5 with 3 pts Na_2CO_3 , allowing to stand in contact with cold water (which dissolves Na_2CO_3), dissolving in hot water, and crystallising

(Joly, *Fremy's Encyclop Chimique*) Santesson (*Bl* [2] 24, 52) obtained this salt by boiling $\text{Nb}_2\text{O}_5 \cdot x\text{H}_2\text{O}$ with NaOH aq, the salt remained insoluble in NaOH aq, slightly soluble in cold water (2) $4\text{Nb}_2\text{O}_5 \cdot \text{Na}_2\text{O} \cdot \text{H}_2\text{O}$, a salt, probably with this composition, was obtained by Santesson (*lc*) as a gelatinous pp by passing CO_2 into solution of NaNbO_3 , (3) $3\text{Nb}_2\text{O}_5 \cdot 2\text{Na}_2\text{O} \cdot 9\text{H}_2\text{O}$. An insoluble amorphous salt, obtained by fusing Nb_2O_5 with NaOH and treating with water (Santesson, *lc*)

Niobates of Ca, Mg, and Mn—viz $\text{Ca}_2\text{Nb}_2\text{O}_7$, $\text{Ca}(\text{NbO}_3)_2$, $\text{Mg}_2\text{Nb}_2\text{O}_7$, $2\text{MgO} \cdot \text{Mg}_2\text{Nb}_2\text{O}_7$, $\text{MgO} \cdot \text{Mg}_2\text{Nb}_2\text{O}_7$, $\text{Mn}(\text{NbO}_3)_2$ —have been obtained by Joly (*lc*) by fusing CaCl_2 , MgCl_2 , and MnCl_2 with Nb_2O_5 . Joly also obtained a niobate of Fe, and a niobate of Fe and Mn. H. Rose (*P* 90, 456) obtained niobates of Cu, Hg, and Ag by adding salts of these metals to solutions of NaNbO_3 .

FLUONIOMATES These salts, which may also be regarded as compounds of NbF_5 with metallic fluorides, and sometimes also with HF , are obtained by dissolving $\text{Nb}_2\text{O}_5 \cdot x\text{H}_2\text{O}$ in large excess of HFAq , adding metallic carbonates, and evaporating, the fluoniomates are also formed by dissolving fluoxyniobates (*u infra*) in HFAq and evaporating (Marnagac, *A Ch* [4] 13, 5, Santesson, *Bl* [2] 24, 52). The following are the principal fluoniomates—



FLUOXINIOMATES These salts are obtained by dissolving Nb_2O_5 with alkali fluorides in HFAq , and evaporating. They may be regarded as derived from the hypothetical acids H_2NbOF_5 , H_3NbOF_6 , and H_4NbOF_7 , they may also be looked on as compounds of NbOF_5 with alkali fluorides. The fluoxyniobates have been examined chiefly by Marnagac (*A Ch* [4] 8, 5, 13, 5).

Ammonium fluoxyniobates 1 $(\text{NH}_4)_2\text{NbOF}_5$ ($=\text{NbOF}_5 \cdot 2\text{NH}_4\text{F}$) Obtained by dissolving Nb_2O_5 and NH_4F in HFAq , and evaporating, easily soluble rhombic prisms, isomorphous with $\text{WO}_3 \cdot 2\text{NH}_4\text{F}$ —2 $(\text{NH}_4)_3\text{NbOF}_6$ ($=\text{NbOF}_5 \cdot 3\text{NH}_4\text{F}$) Obtained similarly to the foregoing salt, forms octahedral crystals, isomorphous with $\text{ZrF}_3 \cdot 3\text{NH}_4\text{F}$ (Baker, *C J* 35, 762)—3 $(\text{NH}_4)_4\text{NbOF}_7$ ($=\text{NbOF}_5 \cdot 4\text{NH}_4\text{F}$) Obtained by dissolving $\text{Nb}_2\text{O}_5 \cdot x\text{H}_2\text{O}$ in conc NH_4FAq , cubic and octahedral crystals of the regular system (Joly, *P* 108, 467)—4 $(\text{NH}_4)_5\text{Nb}_2\text{O}_7 \cdot \text{H}_2\text{O}$ ($=3\text{NbOF}_5 \cdot 5\text{NH}_4\text{F} \cdot \text{H}_2\text{O}$) Obtained by adding less than an equivalent of NH_4F to Nb_2O_5 in HFAq , and evaporating.

Potassium fluoxyniobates—1 $\text{K}_2\text{NbOF}_5 \cdot \text{H}_2\text{O}$ ($=\text{NbOF}_5 \cdot 2\text{KF} \cdot \text{H}_2\text{O}$) Obtained by evaporating a solution of Nb_2O_5 in HFAq after addition of KF . By crystallising from water, the salt separates in such fine tablets that the liquid appears to gelatinise, monoclinic tables are obtained by crystallising from water contain-

ing a little HF . Loses H_2O at 100° , melts at red heat, sol in 12–13 pts water at 17° – 21° , easily soluble in hot water—2 K_3NbOF_6 ($=\text{NbOF}_5 \cdot 3\text{KF}$) Obtained by adding excess of KF to solution of the foregoing salt. Cubical crystals belonging to the regular system (Baker, *C J* 35, 761)—3 $\text{K}_4\text{Nb}_2\text{O}_7 \cdot \text{H}_2\text{O}$ ($=3\text{NbOF}_5 \cdot 5\text{KF} \cdot \text{H}_2\text{O}$) Obtained by adding less than an equivalent of KF to Nb_2O_5 in HFAq , evaporating a little, separating from $\text{K}_2\text{NbOF}_5 \cdot \text{H}_2\text{O}$ which separates, and evaporating the mother liquor—4 $\text{K}_5\text{NbOF}_6 \cdot \text{HF}$ ($=\text{NbOF}_5 \cdot 3\text{KF} \cdot \text{HF}$) Obtained by dissolving Nb_2O_5 in considerable excess of HFAq , and adding excess of KF , isomorphous with $\text{SnF}_6 \cdot 3\text{KF} \cdot \text{HF}$.

Fluoxyniobates of Cu and Zn have also been obtained, $\text{CuNbOF}_5 \cdot 4\text{H}_2\text{O}$, and $\text{ZnNbOF}_5 \cdot 6\text{H}_2\text{O}$.

Niobium, alloys of An alloy of Nb with Al, approximately of the composition Nb_2Al_3 , was obtained by Marnagac (*Ar Sc* 31, 89) by heating $\text{NbF}_5 \cdot 2\text{KF}$ with Al in a carbon crucible, and treating with cold HCl aq. A grey, crystalline, metal like powder, SG 4.45 to 4.52. Soluble in hot HCl aq with evolution of H. Insoluble in HNO_3 aq or dilute H_2SO_4 aq, boiling conc. H_2SO_4 forms SO_2 and S, soluble in HFAq .

Niobium, bromide of NbBr_3 . Formula probably molecular, because of similarity with NbCl_3 , which has been gasified. A purple red solid, obtained by passing CO_2 laden with Br vapour over a heated mixture of Nb_2O_5 and C (H. Rose, *P* 104, 442).

Niobium, carbide of By heating to c 1500° a mixture of 4 pts Nb_2O_5 , 1 pt sugar carbon, and 1 pt Na_2CO_3 , Joly obtained large violet needles of the composition NbC (*Bl* [2] 25, 206).

Niobium, carbonitride of By heating Nb_2O_5 with a mixture of Na_2CO_3 and C to c 1200° , Deville (*C R* 66, 180) obtained a crystalline mass, which evolved NH_3 when heated with molten KOH , according to Joly (*Bl* [2] 25, 206), this substance is either a carbonitride of Nb, or a mixture of carbide, NbC , with nitride NbN .

Niobium, chlorides of Two chlorides of Nb are known, NbCl_3 and NbCl_5 .

NIOBIUM PENTACHLORIDE NbCl_5 . Mol wt 270.85. VD 138.9 (Deville, a Troost, *C R* 60, 1221). Melts at 194° and boils at 240° (D. a. T., *lc*).

Preparation—Perfectly dry Nb_2O_5 is mixed with a large excess of dry sugar or starch, the mixture is completely charred by heating in a closed crucible, and a small quantity is then placed in a rather wide tube of hard glass, narrowed here and there, connected with a CO_2 and a Cl apparatus, the tube is gently heated for some time while a stream of perfectly dry CO_2 is passed through it, and is then allowed to cool in the CO_2 , when cold, perfectly dry Cl is passed through the tube, when every trace of CO_2 is expelled, the tube is gradually heated to bright redness in the stream of Cl, NbCl_5 , mixed with a little NbOCl_3 , collects in the wider parts of the tube. The NbCl_5 is separated from the less volatile NbOCl_3 by distillation in dry Cl.

As NbCl_5 is very voluminous the operation must be conducted in wide tubes and with small quantities of the mixed Nb_2O_5 and C (H. Rose). **Properties and Reactions**—Yellow needles, melts at 194° , beginning to sublime at 125° , boils at 240° (Deville a Troost *C R* 60, 1221). Vapour

is yellow Soluble in alcohol Fumes in air, giving off HCl Decomposed by water to HCl and $\text{Nb}_2\text{O}_5 \cdot x\text{H}_2\text{O}$ Soluble in cold conc HClAq ; Zn produces a blue colour in this solution, on dilution and heating, $\text{Nb}_2\text{O}_5 \cdot x\text{H}_2\text{O}$ separates. Soluble in conc H_2SO_4 with evolution of HCl. Vapour of NbCl_5 is reduced to Nb by heating with H (Blomstrand, Roscoe, *C N* 37, 25) NbOCl_3 is produced by heating with Nb_2O_5 . $\text{Nb}_2\text{O}_5 \cdot \text{S}_3$ is formed by heating in CS_2 vapour (Delafontaine, *Ar Sc* 27, 167)

Niobium trichloride NbCl_5 When vapour of NbCl_5 is slowly passed through a red-hot tube, a dark grey metal like crust forms on the sides of the tube, this crust is NbCl_5 (Roscoe, *C N* 37, 25) Not volatile, non-deliquescent, unchanged by H_2O or NH_3Aq , by HNO_3Aq gives HCl and $\text{Nb}_2\text{O}_5 \cdot x\text{H}_2\text{O}$ Heated in air, gives off white fumes When heated in CO_2 , produces CO and NbOCl_3 .

Niobium, fluoride of No fluoride of Nb has been isolated with certainty $\text{Nb}_2\text{O}_5 \cdot x\text{H}_2\text{O}$ dissolves easily in HFAq , on evaporation a non-crystallisable mass is obtained, which evolves white fumes when heated and leaves Nb_2O_5 Solution of $\text{Nb}_2\text{O}_5 \cdot x\text{H}_2\text{O}$ in HFAq yields *fluoniobates* when mixed with metallic carbonates and evaporated, these fluoniobates may be regarded as compounds of NbF_5 with metallic fluorides (*v Fluoniobates*, p 507)

Niobium, haloid compounds of. The only haloid compound of Nb which has been gasified is NbCl_5 , the trichloride is also known, and the formula NbCl_5 is probably molecular No fluoride or iodide has been isolated, but several compounds are known, which may be regarded as formed by the union of NbF_5 with more positive fluorides (*v Fluoniobates*, p 507) NbCl_5 reduces CO_2 to CO at a high temperature Oxihaloid compounds are known, of the form NbOX_2 , where X = Br, Cl, or F

Niobium, hydride of, ?NbH Margnac (*Ar Sc* 1868) obtained a heavy grey powder, having approximately the composition NbH_2 , mixed with a little Nb_2O_5 , by heating K.NbF_6 , mixed with a little KHF_2 , and covered with NaCl , with excess of Na in an iron crucible The reaction was energetic, the fused mass was broken up, treated with water, then with water containing a little HF, then washed with water, and finally filtered and dried The powder obtained by Margnac had SG 6 to 6.6, it dissolved in conc HFAq with rapid evolution of H, it was insoluble in HClAq , HNO_3Aq , and dilute $\text{H}_2\text{SO}_4\text{Aq}$, sol warm conc H_2SO_4 , also in molten KHSO_4 , heated in air or O to above 100° it burned to Nb_2O_5 and H_2O , it was unchanged when heated in H Krüss a Nilson (*B* 20, 1691) repeated Margnac's experiments, using a quantity of Na equivalent to the K.NbF_6 , they obtained a mixture of c 77 p c NbH , c 21.5 p c Nb_2O_5 , and c 1 p c Fe_2O_3 , K a N gave S.H of NbH as 0.97 at 0° to 100° , 0.92 at 0° to 210.5° , 0.87 at 0° to 301.5° , and 0.83 at 0° to 449° .

Niobium, nitride of, ?NbN NbCl_5 absorbs NH_3 , on heating, NH_4Cl is evolved, and a black mass remains which contains N Heated with KOH.Aq , NH_3 is evolved, it is not attacked by HNO_3Aq ; soluble in HFAq ; heated in air, it oxidises with incandescence The composition of this body is approximately NbN (H Rose;

Deville, *C R* 66, 180, Joly, *Bl* [2] 25, 206). By reducing Nb_2O_5 with a mixture of soda and O, Deville obtained a crystalline mass, probably a mixture of nitride and carbide of Nb

Niobium, nitro-carbide of, v Niobium, carbide of, p 507

Niobium, oxides of Three oxides of Nb have been isolated, NbO , NbO_2 , and Nb_2O_5 , a fourth, Nb_2O_3 , probably exists Nb_2O_5 is formed by heating Nb in air or O, also by decomposing NbOCl_3 by water, and in other ways, Nb_2O_5 is produced by the partial reduction of Nb_2O_5 in H, NbO is obtained by the incomplete reduction of NbOF_3 or NbOCl_3 by Na or Mg, when Nb_2O_5 in HClAq is reduced by Zn the solution becomes brown, and a solid separates, which is probably Nb_2O_3 Moist Nb_2O_5 reacts as an acid forming oxide, niobates are formed by fusing Nb_2O_5 with basic acids or carbonates (*v Niobates*, p 506) The mol wt of none of the oxides of Nb is known with certainty

Niobic oxide Nb_2O_5 (*Niobic anhydride* *Niobium pentoxide*)

Occurrence—Niobates occur in a few rare minerals, e.g. columbite, tantalite, and samarskite.

Preparation—1 NbOCl_3 is agitated with water, the insoluble $\text{Nb}_2\text{O}_5 \cdot x\text{H}_2\text{O}$ is washed till free from HCl, dried at 100° , and heated to incipient redness The solution after treating NbOCl_3 with water contains much Nb_2O_5 , the oxide is obtained by adding slight excess of NH_3Aq , warming till every trace of NH_3 is removed, collecting the pp, washing till free from HCl, and drying—2 Dilute $\text{H}_2\text{SO}_4\text{Aq}$ is added to a boiling solution of NaNbO_3 , the ppd $\text{Nb}_2\text{O}_5 \cdot x\text{H}_2\text{O}$ is thoroughly washed and heated—3 Impure Nb_2O_5 is fused with KHSO_4 , the fused mass is treated with water, and the pp is washed and heated.—4 Nb_2O_5 is obtained in crystals by dissolving in molten borax, heating in a porcelain oven, washing, and drying (Nordenskjöld, *P* 114, 612, Ebelmen, *A Ch* [3] 33, 34, Knop, *Z K* 12, 610, also by strongly heating Nb_2O_5 (from NbOCl_3) in a slow current of HCl (Deville, *C R* 66, 180) For preparation of Nb_2O_5 from columbite v NIOBIUM, Preparation, p 505

Properties—A white powder, becomes yellow when heated, and goes white on cooling Insoluble in water SG 4.4 to 4.53 (Marnagac, *A Ch* [4] 8, 5) The crystals of Nb_2O_5 are flat, right angled tablets, they are optically active (Nordenskjöld, *P* 114, 612, cf Ebelmen, *A Ch* [3] 33, 34, Knop, *A* 169, 56) SH 118 at 0° to 210.5° , 124 at 0° to 301.5° , 134 at 0° to 449° (Krüss a Nilson, *B* 20, 1691)

Reactions—1. Dissolves in hot conc *sulphuric acid*, the solution may be diluted with out ppn, but on heating all the Nb_2O_5 is ppd, the pp contains H_2SO_4 —2 Boiling *hydrochloric acid* dissolves only traces of Nb_2O_5 , the residue is easily soluble in water, and this solution is ppd on boiling with $\text{H}_2\text{SO}_4\text{Aq}$ (Wöhler, *P* 48, 93, Margnac, *A Ch* [4] 8, 15, 13, 20, H Rose, *P* 112, 484).—3. Easily dissolved by cold *hydrofluoric acid*—4. Soluble in *caustic potash* solution—5. *Caustic soda* does not dissolve Nb_2O_5 , but the product is soluble in water Nb_2O_5 , which has been strongly heated is insoluble in H_2SO_4 , HClAq , or HFAq , it is dissolved by molten alkalis.—6. Reduced by *hydrogen* to

NbO₂, at full red heat — 7 Moist Nb₂O₅ dissolved in HClAq is reduced by *zinc* with formation of a blue liquid which then becomes brown and deposits brown flocks, probably of Nb₂O₃ (Magnnac, *A Ch* [4] 18, 5) — 8 Strongly heated with *hydrogen sulphide*, or *carbon disulphide*, an oxysulphide is formed, probably Nb₂OS₂ (Delafontaine, *Ar Sc* 27, 167) — 9 Heated with *ammonia*, Nb nitride (*q. v.*) is formed — 10 Mixed with *carbon* and heated in *chlorine* or *bromine*, NbCl₃ (or NbBr₃) is formed along with some NbOCl₃ (or NbOBr₃) — 11 Fused with *basic oxides* or *carbonates*, niobates (*q. v.*) are produced

Combinations — 1 With *water* to form various hydrates (*v. infra*) — 2 Nb₂O₅ appears to combine with some acids, but no definite compounds have yet been isolated, *e.g.* the pp obtained by decomposing NbOCl₃ by water in presence of Na₂HPO₄ contains H₂PO₄, and the pp obtained by adding water to Nb₂O₅ in H₂SO₄ and boiling contains H₂SO₄ (*v. Blomstrand, Acta Univ Lund* 1864)

HYDRATES OF NIOBIC OXIDE Various hydrates of Nb₂O₅ are known. By fusing Nb₂O₅ with KHSO₄, boiling with water, dissolving the pp in HClAq, and ppg by NH₄Aq, Santesson (*Bt* [2] 24, 52) obtained a flocculent pp containing 8.04 to 8.41 p.c. H₂O, agreeing with the formula 3Nb₂O₅·4H₂O. The solid obtained by ppg NaNbO₄Aq by H₂SO₄Aq and drying at 100° has the composition Nb₂O₅·7H₂O, according to Santesson (*l.c.*) The hydrate obtained by decomposing NbOCl₃ by water is amorphous, that formed by the action of moist air on NbOCl₃ is said to be crystalline (*H. Rose, P* 112, 557). The hydrates of Nb₂O₅ react as weak acids (*v. Niobates*, p. 506)

NIOBOUS OXIDE NbO (Niobium monoxide) By reducing NbOF₃·2KF with Na, H. Rose obtained a black powder which he thought to be Nb (*P* 104, 312). This substance was recognised as an oxide by Delafontaine (*Ar Sc* 27, 167). Prepared by strongly heating NbOF₃·2KF with Na, under KCl, and washing with cold water SG 6.3 to 6.67. Obtained in crystals by passing vapour of NbOCl₃ over heated Mg wire (Deville a Troost, *C R* 60, 1221, *v. also* Deville, *C R* 66, 183). Black, lustrous, regular crystals. Moist NbO is soluble in boiling dilute HClAq, or in HFAq, H is said to be evolved. KOHAq dissolves NbO, forming K niobate. Molten KHSO₄ forms Nb₂O₅, heated in Cl NbOCl₃ is produced

NIOBIUM DIOXIDE Nb₂O₃ (Niobium tetroxide [Nb₂O₃]) A black powder with blue reflection, insol water and acids. Formed by heating Nb₂O₅ in a stream of H to full white heat (Delafontaine, *Ar Sc* 27, 167)

Wöhler (*P* 48, 93) noticed that Zn reduces a solution of Nb₂O₅ in HClAq, with production of blue and then brown coloured substances. Magnnac (*A Ch* [4] 13, 5) obtained a blue-brown pp by boiling Nb₂O₅·xH₂O with HClAq, dissolving the residue in water, and reducing by Zn

Niobium, oxybromide of, NbOBr₃ A voluminous, crystalline, yellowish solid, obtained by passing Br vapour over a heated mixture of Nb₂O₅ with a little charcoal (*H. Rose, P* 104, 442). Sublimes without melting. Heated in CO₂ gives Nb₂O₅ and NbBr₃. Decomposed by water, giving Nb₂O₅·xH₂O and HBrAq

Niobium, oxychloride of, NbOCl₃ Mol. w 216.1. Obtained, along with NbCl₃, by heating Nb₂O₅ mixed with charcoal in a stream of Cl₂, also by heating Nb₂O₅ in a stream of CO₂ charged with vapour of NbCl₃ (Deville a Troost, *C R* 60, 1221). A white, lustrous mass, sublimes at c. 400° without melting. VD at 440° to 810° — 114 (*D a T, l.c.*) Heated strongly in CO₂, NbCl₃ and Nb₂O₅ are formed, the same products are formed by heating in H (Blomstrand, *Acta Univ Lund*, 1864). Sol alcohol, decomposed by water to Nb₂O₅·xH₂O and HClAq

Niobium, oxyfluoride of, NbOF₃ Small crystals, optically active, resemble ZrF₄, obtained by strongly heating Nb₂O₅ mixed with a large excess of CaF₂ in HCl (*Joly, C R* 81, 1266). NbOF₃ forms various compounds with metallic fluorides (*v. Fluoxyniobates*, p. 507)

Niobium, oxysulphide of, Nb₂OS₂ A black powder, obtained by passing H₂S or CS₂ vapour over strongly heated Nb₂O₅. The product of these reactions was supposed by H. Rose to be a sulphide of Nb (*P* 111, 193, *v. also* Rose a Hermann, *J pr* 111, 393). Delafontaine (*Ar Sc* 27, 167) showed the substance to be an oxysulphide, Rammelsberg (*J pr* 108, 95) thought the composition was NbOS or Nb₂O₅S₂

Niobium, salts of No compounds obtained by replacing the H of acids by Nb have yet been isolated. There are indications that Nb₂O₅ combines with some acids (*v. Niobic oxide, Combinations*, No 2, *supra*) M M P M

NITRANILIC ACID v. DI NITRO DI OXYQUINONE

NITRANILINE v. NITROANILINE

NITRATES Salts of *nitric acid*, HNO₃. The greater number of the nitrates are normal salts, many basic nitrates also exist. The general formula for normal nitrates may be written Mⁿ·nNO₃, where Mⁿ denotes a metal of *n* valency. The normal nitrates may also be regarded as composed of a basic and an acidic radicle, on this view, they are classed under the general formulae M₂O N₂O₅, MO N₂O₅, M₂O·3N₂O₅, MO₂·2N₂O₅. The simplest way of looking at the composition of the basic nitrates is to regard them as compounds of the acidic radicle N₂O₅ with more than the normal quantity of base, thus normal lead nitrate is PbO N₂O₅, and basic lead nitrate is 3PbO N₂O₅. Several basic nitrates may be formulated as salts of the hypothetical orthonitric acid H₃NO₃, which bears the same relation to ordinary, or meta, nitric acid that orthophosphoric bears to metaphosphoric acid, thus basic lead nitrate 3PbO N₂O₅ may be written Pb₃(NO₃)₄.

Some nitrates occur native, *e.g.* Ca(NO₃)₂, Mg(NO₃)₂, KNO₃, NaNO₃. Alkali nitrates are found in river, spring, and drainage waters, and in the juices of some plants. With regard to the formation of nitrates in the soil *v. NITRIFICATION*, this vol. Nitrates are prepared by dissolving metals, metallic oxides or carbonates, in nitric acid, also, in some cases, by double decomposition from the alkali nitrates.

Most nitrates are crystalline salts. As no nitrate has been gasified, the formulae of these salts are not necessarily molecular. The normal nitrates are soluble in water, a few, *e.g.* Bi(NO₃)₃, are decomposed by water with production of insoluble basic nitrates. Nitrates are decomposed

by heat, a few give off HNO_3 , but in almost all cases O is evolved, along with oxides of N and H_2O , the final residue is generally a metallic oxide corresponding with the nitrate used, AgNO_3 leaves a residue of Ag. Heated with combustible bodies, nitrates cause deflagration or explosion, if the combustible body be an acid forming element, or a compound capable of forming an acid by oxidation, a salt is formed composed of the metal of the nitrate and the acid produced from the combustible body. Thus K_2SeO_4 is formed by deflagrating KNO_3 with Se, and K_2MnO_4 by deflagrating KNO_3 with an oxide or salt of Mn. Alkali nitrates are reduced to NH_3 by the action of potash and zinc, or by a pair of metals one of which is distinctly more electro-positive than the other, e.g. by Cu and Zn, Fe and Zn, Pt and Zn, &c. Alkali nitrates are also reduced to NH_3 by the action of common putrefactive organisms in presence of peptones and air, also by Pt black charged with O, in the presence of dextrose (v. Loew, *B* 23, 675). Nitrates are reduced to nitrites, N_2O , NO, and N, by organisms present in the soil (v. Warrington, *C* J 45, 669, 53, 742 [references are given here to other memoirs], 59, 484, Munro, *C* J 49, 667).

The greater number of the nitrates are insoluble in conc. nitric acid. A few dissolve in a large quantity of the acid, according to Ditte (*A. Ch.* [5] 18, 320) these nitrates combine with HNO_3 to form acid salts, e.g. $\text{KNO}_3 \cdot 2\text{HNO}_3$, $\text{NH}_4\text{NO}_3 \cdot \text{HNO}_3$, $\text{KNO}_3 \cdot 3\text{HNO}_3$, $\text{RbNO}_3 \cdot 5\text{HNO}_3$. Some other hydrated nitrates dissolve in warm HNO_3 . Aq. when dehydrated, on cooling, hydrates are deposited containing less water than those which crystallise from water, to this class of nitrates belong $\text{Mg}(\text{NO}_3)_2$, $\text{Mn}(\text{NO}_3)_2$, $\text{Zn}(\text{NO}_3)_2$, $\text{Al}(\text{NO}_3)_3$, $\text{Cu}(\text{NO}_3)_2$ (Ditte, *l.c.*).

The methods of detecting and estimating nitrates are numerous, reference must be made to *Manuals of analysis*.

Aluminium nitrates. The normal salt, $\text{Al}(\text{NO}_3)_3 \cdot 9\text{H}_2\text{O}$, obtained by dissolving AlO_3H_3 in HNO_3 Aq. and evaporating, crystallises in oblique rhombic prisms. Melts at 73° , deliquescent, e. sol. H_2O and HNO_3 Aq. (Ordway, *A* 76, 247, Salm Horstmar, *J* 1850 301, Thorey, *Russ. Zeitschr. Pharm.* 10, 321). The salt does not react with HCl gas (Thomas, *C* J 33, 367). Basic Al nitrates are obtained by digesting $\text{Al}(\text{NO}_3)_3$ with AlO_3H_3 (Ordway, *l.c.*).

Ammonium nitrate NH_4NO_3 . According to Tissandier (*C* R 82, 388) this salt is frequently present in rain water. It is obtained by adding a slight excess of NH_3 Aq. to HNO_3 Aq. and evaporating, also by passing the electric discharge through a mixture of H, N, and O, by passing H_2S into dilute HNO_3 Aq. by the interaction of certain metals, e.g. Sn, with HNO_3 Aq. NH_4NO_3 crystallises in various forms according to the temperature, the crystals formed at 36° are trimetric, those formed at 87° are rhombohedral, and monometric crystals are produced at 120° (Lehmann). The specific heats, volume changes, and heats of transformation, of the various modifications have been determined by Bellati a Romanese (*Nuovo Cimento*, [3] 21, 5, abstract in *C* J 54, 106). SG 1.707 (Kopp, *A* 36, 1), 1.709 (Sohiff, *A* 112, 88), for other results v. Clarke's *Specific Gravity Table*, new ed. 110. NH_4NO_3 dissolves in H_2O with a large

disappearance of heat. S at 18° c. 200; saturated solution contains 47.8 p.c. NH_4NO_3 , and boils at 164° . E. sol. alcohol. Deliquesces in air, losing NH_3 and acquiring an acid reaction. Melts at c. 152° , decomposition begins at c. 210° and becomes explosive at c. 300° , products are H_2O and N_2O , but part of salt volatilises (Berthelot, *C* R 82, 932), heated very rapidly, NH_3 , NO, and NH_4NO_3 are also formed (B). According to B (*l.c.*) NH_4NO_3 may be sublimed unchanged, by placing the fused salt in a basin covered with filter paper, over which is a paper cylinder filled with coarse fragments of glass, and heating gently not above 190° – 200° . NH_4NO_3 condenses considerable quantities of NH_3 , forming a liquid varying in composition according to temperature and pressure (v. Divers, *Pr* 21, 107, Raoult, *C* R 77, 788). At -10° , and 760 mm, the liquid $\text{NH}_4\text{NO}_3 \cdot 2\text{NH}_3$ is formed, heated to 28.5° a solid remains, $\text{NH}_4\text{NO}_3 \cdot \text{NH}_3$ (R, *l.c.*), cf. Mendelejeff (*B* 23, 3464), who regards $\text{NH}_4\text{NO}_3 \cdot \text{NH}_3$ and $\text{NH}_4\text{NO}_3 \cdot 2\text{NH}_3$ as amides obtained from NO OH ONH, ONH, and NO(OH), which are the NH, salts of hypothetical ortho-nitric acid NO(OH). The liquid compounds of NH_4NO_3 and NH_3 react with many salts, the reactions generally resembling those of NH_4NO_3 and dry NH_3 combined (for details v. Divers, *l.c.*). NH_4NO_3 absorbs dry HCl, forming NH_4Cl , after a time a little Cl and NO are evolved (Thomas, *C* J 33, 367). The Cu Zn couple reduces NH_4NO_3 Aq. to NH_3 and NH_4NO_3 , at B P NO is evolved (Gladstone a Tribe, *C* J 33, 150).

Antimony nitrate. The compound $\text{Sb}_2\text{O}_5 \cdot \text{N}_2\text{O}_5$ is said to be formed by dissolving Sb_2O_5 in cold fuming HNO_3 (Péligot, *C* R 23, 709).

Barium nitrate $\text{Ba}(\text{NO}_3)_2$. Crystallises in tetartohedral forms belonging to the regular system (Soacchi, *J* 1860 13, Baumhauer, *Z* K 1, 51, Lewis, *P. M.* [5] 3, 453). SG α 2.2 to 3.24 (Kremers, *J* 5, 15, for other determinations v. Clarke's *Specific Gravity Table* (new ed.), 111). H F [Ba , O, N°O Aq.] = 187,020 (?) (*Th* 3, 518). Melts at c. 593° (Carnelley, *C* J 33, 278). S 5 at 0° , 7 at 10° , 9.2 at 20° , 11.6 at 30° , 14.3 at 40° , 17.1 at 50° , 20.3 at 60° , 23.6 at 70° , 27 at 80° , 30.6 at 90° , 32.2 at 100° , saturated solution boils at 101.9° , SG and potge composition of $\text{Ba}(\text{NO}_3)_2$ Aq. are as follows (Mulder) —

SG	P. a. $\text{Ba}(\text{NO}_3)_2$	SG	P. c. $\text{Ba}(\text{NO}_3)_2$
1.009	1	1.05	6
1.017	2	1.06	7
1.025	3	1.069	8
1.034	4	1.078	9
1.042	5	1.087	10

$\text{Ba}(\text{NO}_3)_2$ is only slightly soluble in water containing HCl or HNO_3 , insol. alcohol.

$\text{Ba}(\text{NO}_3)_2$ is prepared by adding to BaCO_3 , or crude BaS, enough HNO_3 Aq. to decompose almost the whole of the salt, filtering, and crystallising, also by mixing equivalent weights of BaCl_2 and NaNO_3 in solution, and recrystallising the $\text{Ba}(\text{NO}_3)_2$, which separates (Bolley, *C* C 1860 830, Kuhlmann, *D* P. J. 150, 57, 108, 416). $\text{Ba}(\text{NO}_3)_2$ melts at a moderate temperature, at red heat it evolves O, N, and NO_2 , and leaves BaO, according to Rammelsberg (*B* 2, 147, 7, 542) the residue

contains more O than BaO , and has the composition $\text{Ba}_2\text{O} \cdot \text{Ba}(\text{NO}_3)_2$, is not acted on by HCl gas (Thomas, *C J* 33, 367)

Beryllium nitrates The composition of these salts is doubtful. They are very soluble in water and difficult to crystallise. By double decomposition from BeSO_4Aq and evaporating, Ordway (*J pr* 76, 22) obtained deliquescent crystals approximating to the composition $\text{Be}(\text{NO}_3)_2 \cdot 8\text{H}_2\text{O}$. By heating at 100° the crystals lost half of their nitric acid, and a basic salt remained, soluble in water. Other soluble basic salts seem to be formed by adding a little NH_3Aq to a solution of the normal salt, and by digesting the normal salt with BeO *xH*₂O.

Bismuth nitrates The normal salt has the composition $\text{Bi}(\text{NO}_3)_3 \cdot 10\text{H}_2\text{O}$ according to Gladstone (*J pr* 44, 179) and Hemtz (*J pr* 45, 102), according to the more recent work of Yvon (*C R* 84, 1164) the crystallised salt has the composition $2\text{Bi}(\text{NO}_3)_3 \cdot 11\text{H}_2\text{O}$. The normal salt is formed by dissolving Bi , Bi_2O_3 , or $\text{Bi}_2(\text{CO}_3)_3$ in HNO_3Aq , filtering through asbestos or powdered glass, and evaporating to the crystallisation-point. Forms large deliquescent crystals, SG 2.823 at 13° (Clarke's *Table of Specific Gravities* (new ed.), 112). Crystals are very caustic, they melt easily in the water of crystallisation, decomposition begins at $c 75^\circ\text{--}80^\circ$ with production of basic salts (*v infra*). Decomposed by HCl gas, giving BiCl , and also much Cl , along with NO , H_2O , and possibly other oxides of N and Cl (Thomas, *C J* 33, 367). Various basic salts have been described. According to Graham (*A* 29, 16) $2(\text{BiO} \cdot \text{NO}_2) \cdot \text{H}_2\text{O}$ is formed by heating the normal salt to 80° , and is not decomposed below 260° (*cf* Ruge, *J* 1862 163). Yvon (*C R* 84, 1164) assigns the composition $4(\text{BiO} \cdot \text{NO}_2) \cdot 3\text{H}_2\text{O}$ to the salt obtained by heating the normal salt to 120° , and also to the product of the action of water on the normal salt. Many basic salts, $x\text{BiO} \cdot y\text{N}_2\text{O} \cdot z\text{H}_2\text{O}$, seem to be produced by decomposing Bi_3NO_3 , or a solution of Bi in HNO_3Aq , by water, the composition of the most stable of these *subnitrates* is $\text{BiO} \cdot \text{NO}_2 \cdot \text{H}_2\text{O}$, in other cases x , y , and z have such values as 5, 4, and 9, or 5, 3, and 8, or 6, 5, and 9. The composition of these basic salts varies with the temperature of the water used, the amount of washing given to the pp, and the length of time the pp is allowed to remain in contact with the acid liquid above it. The compositions of these salts, and the preparation of a salt of constant composition for medicinal use, have been examined chiefly by Phillips (*J Ph* 18, 688), Duflos (*Ar Ph* [2] 23, 307), Herberger (*R P* 55, 289, 306), Ullgren (*B J* 17, 169), Dulk (*R P* 83, 1), Becker (*Ar Ph* 55, 81, 129), Janssen (*Ar Ph* 68, 1, 129), Ruge (*J* 1862 163), and Yvon (*C R* 84, 1164).

Cadmium nitrate $\text{Cd}(\text{NO}_3)_2 \cdot 4\text{H}_2\text{O}$ White, prismatic, deliquescent, needles, by dissolving Cd , CdO , or CdCO_3 in HNO_3Aq and evaporating SG 2.45 at 14° , 2.46 at 20° (Laws, *Am S* [3] 14, 281). H F [$\text{CdO} \cdot \text{N}^\circ\text{O} \cdot 4\text{H}^\circ\text{O}$] = 125,170, [$\text{CdO} \cdot \text{N}^\circ\text{O} \cdot \text{Aq}$] = 86,000 (*Th* 3, 518). Melts at $59^\circ\text{--}5^\circ$, and boils at $c 132^\circ$ (Ordway, *Am S* [2] 27, 14). Reacts with HCl gas to produce CdCl , evolving Cl and NO (Thomas, *C J* 33, 367). Wells (*Am* 9, 304) describes a basic salt, to this salt he has assigned the composition

$2\text{CdO} \cdot \text{N}_2\text{O} \cdot 3\text{H}_2\text{O}$, obtained by digesting hot $\text{Cd}(\text{NO}_3)_2\text{Aq}$ with CdO , and allowing to cool.

Cesium nitrate CsNO_3 . Obtained by dissolving Cs_2CO_3 in HNO_3Aq , and evaporating, the *habitus* of the crystals depends on the rate of evaporation. Melts below red heat, when strongly heated evolves O, and forms CsNO_2 , S 10.68 at 32° , very slightly sol in alcohol (Bunsen, *P* 119, 1).

Calcium nitrate $\text{Ca}(\text{NO}_3)_2 \cdot 4\text{H}_2\text{O}$. Occurs in soils when conditions are favourable to production of HNO_3 , and Ca salts are also present. This salt is prepared in some countries by the slow decomposition of animal and vegetable matter SG 1.79 when liquid, and 1.9 when solid, at $15^\circ\text{--}5^\circ$ (Ordway, *J* 12, 115). SG of $\text{Ca}(\text{NO}_3)_2 = 2.5$ at 17.9° (Favre a Valson, *C R* 77, 579). H F [$\text{CaO} \cdot \text{N}^\circ\text{O} \cdot 4\text{H}^\circ\text{O}$] = 218,440, [$\text{CaO} \cdot \text{N}^\circ\text{O} \cdot \text{Aq}$] = 177,160. Prepared by dissolving CaO or CaCO_3 in HNO_3Aq , and evaporating, if the evaporation is continued to dryness the anhydrous salt is obtained. The hydrated salt crystallises with difficulty in deliquescent, six-sided prisms, melts at 44° , boils at 182° , remaining clear till c one third of the water has gone, when the anhydrous salt is deposited (Ordway, *Am S* [2] 27, 14). The dry salt $\text{Ca}(\text{NO}_3)_2$ melts at 561° (Carnelley, *C J* 33, 278). Decomposed at high temperature, giving off O and NO, the partially decomposed salt is phosphorescent (*Baldwin's phosphorus*), not acted on by HCl gas (Thomas, *C J* 33, 367).

Cerium nitrates *Cerous nitrate*, $\text{Ce}(\text{NO}_3)_3 \cdot 6\text{H}_2\text{O}$, is obtained as a pale rose-coloured, deliquescent, crystalline mass, by dissolving Ce_2O_3 , or CeO , in presence of reducing substances, in HNO_3Aq , evaporating, and drying over H_2SO_4 (Lange, *J pr* 82, 129). Gives off $3\text{H}_2\text{O}$ at 150° , and decomposes at 200° . Forms several double salts with nitrates MNO_3 , and $\text{M}(\text{NO}_3)_2$, e.g. $\text{Ce}(\text{NO}_3)_3 \cdot 2\text{KNO}_3 \cdot 2\text{H}_2\text{O}$, $2\text{Ce}(\text{NO}_3)_3 \cdot 3\text{Mg}(\text{NO}_3)_2 \cdot 24\text{H}_2\text{O}$. These double nitrates have been examined by Lange (*l c*) and Holzmann (*J pr* 84, 76), and more recently by Zschiesche (*J pr* 107, 65). They are obtained by crystallising a mixed solution of $\text{Ce}(\text{NO}_3)_3$ and the other nitrate, and also by dissolving CeO_2 in HNO_3 , adding the other nitrate and a little alcohol, and evaporating. Should the metal of the nitrate which is added be capable of forming a higher oxide than that corresponding to the nitrate used, a little of this oxide is sometimes formed at the expense of the O of the CeO_2 , and the reduction from $\text{Ce}(\text{NO}_3)_3$ to $\text{Ce}(\text{NO}_2)_3$ proceeds without addition of alcohol, thus, addition of $\text{Mn}(\text{NO}_3)_2$ to CeO_2 dissolved in HNO_3Aq produces $2\text{Ce}(\text{NO}_3)_3 \cdot 3\text{Mn}(\text{NO}_3)_2 \cdot 24\text{H}_2\text{O}$, with simultaneous formation of a little MnO_2 . The double cerous nitrates are also formed by dissolving the various metals in an acid solution of CeO_2 in HNO_3Aq , reduction is effected to $\text{Ce}(\text{NO}_2)_3$, *Ceric nitrate*, $\text{Ce}(\text{NO}_3)_4$. Said to be obtained as a reddish yellow mass by evaporating CeO_2 in HNO_3Aq , decomposed by hot water forming a basic salt. Combines with KNO_3 and NH_4NO_3 to form $2\text{Ce}(\text{NO}_3)_4 \cdot 4\text{MNO}_3 \cdot 3\text{H}_2\text{O}$ (Berzelius, *P* 1, 29).

Chromium nitrates The normal salt, $\text{Cr}(\text{NO}_3)_3 \cdot 9\text{H}_2\text{O}$, is obtained by dissolving CrO_3H_2 in HNO_3Aq , evaporating, and crystallising from warm water; the crystals, which form with difficulty, are purple oblique prisms, melting at

87° to a green liquid which boils at 125.5° (Ordway, *Am S* [2] 9, 30, 27, 14) Various basic salts are described by Loewel (*Ph C* 1845, 580), Ordway (*Am S.* [2] 26, 197), and Siewert (*A.* 126, 86), they are formed by heating the normal salt, by dissolving CrO_3H_2 in solution of the normal salt, and by boiling HNO_3Aq with excess of CrO_3H_2 .

Cobalt nitrates The normal salt, $\text{Co}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$, forms red, prismatic, deliquescent crystals, SG 183 at 14° (Boedeker), melts below 100°, at higher temperatures gives off H_2O and oxides of N, and leaves black Co_3O_4 . Produced by dissolving Co, or CoCO_3 , in HNO_3Aq , and evaporating. Easily sol water. Franz (*J pr* [2] 5, 274) gives the following table showing p c $\text{Co}(\text{NO}_3)_2$ in aqueous solutions at 17.5° —

Ra. Co(NO_3),	SG
5	1.0462
10	1.0906
15	1.1378
20	1.1936
25	1.2538
30	1.319
35	1.3896
40	1.4662
Saturated at 17.5°	1.5382

Thomsen gives HF [Co, O', N'O', 6H'O'] 128,880, [Co, O, N'O'Aq] = 84,540. When HCl gas is passed over $\text{Co}(\text{NO}_3)_2 \cdot \text{CoCl}_2$, oxides of N, and Cl are formed (Thomas, *C J* 33, 367). Combines with cerous nitrate to form $\text{Co}(\text{NO}_3)_2 \cdot \text{Ce}(\text{NO}_3)_2 \cdot 8\text{H}_2\text{O}$ (Lange, *J pr* 82, 129).

Basic salts are obtained by adding NH_3Aq to $\text{Co}(\text{NO}_3)_2\text{Aq}$ under different conditions (v Winkelblech, *A* 13, 148, 253, Habermann, *M* 5, 442).

Copper nitrates The normal nitrate, $\text{Cu}(\text{NO}_3)_2 \cdot 3\text{H}_2\text{O}$, is obtained by dissolving Cu or CuO in HNO_3Aq , and evaporating. The solution is at first green owing to production of $\text{Cu}(\text{NO}_3)_2$. Blue prismatic crystals, SG 2.174 (Hassenfratz, *A* 28, 3). The salt $\text{Cu}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ is deposited below 20° HF [Cu, O', N'O', 6H'O'] = 96,950 (*Th* 8, 518). The hexahydrated salt effloresces in air, losing $3\text{H}_2\text{O}$, it melts at 38°, and decomposes at 65°, forming a basic salt. The trihydrated salt melts at 114.5°, and decomposes at 170°. At red heat, $\text{Cu}(\text{NO}_3)_2$ evolves N oxides and leaves CuO , it is deliquescent, easily sol water, but ppd again by conc HNO_3Aq . Franz gives following table (*J pr* [2] 5, 274) —

S.G. of $\text{Cu}(\text{NO}_3)_2\text{Aq}$	P c $\text{Cu}(\text{NO}_3)_2$
1.0942	10
1.2037	20
1.3299	30
1.4724	40
1.5404	44

$\text{Cu}(\text{NO}_3)_2$ is rapidly decomposed by HCl, with formation of CuCl_2 , Cl, NO, and probably N (Thomas, *C J* 33, 367).

Basic nitrates of Cu are produced by boiling $\text{Cu}(\text{NO}_3)_2\text{Aq}$ with KNO_3Aq , or by passing N_2O_4 into H_2O holding $\text{CuO} \cdot \text{H}_2$ in suspension, the product is said to be $4\text{CuO} \cdot \text{N}_2\text{O}_4 \cdot 8\text{H}_2\text{O}$ (Vogel a Reimhauer, *J* 1859 216). For other basic salts v. Graham, *T* 1837 47, Casselmann, *Fzr* 4, 24; Tutschew, *Z.* 6, 109.

Didymium nitrate $\text{Dy}(\text{NO}_3)_3 \cdot 6\text{H}_2\text{O}$. Rose red crystals, by dissolving Dy_2O_3 in HNO_3Aq and crystallising, SG 2.249, loses $6\text{H}_2\text{O}$ at 200°. Easily sol water and alcohol, forms double salts with $\text{Co}(\text{NO}_3)_2$, $\text{Ni}(\text{NO}_3)_2$, and $\text{Zn}(\text{NO}_3)_2$ (v Mariagnas, *A Ch* [3] 88, 148, Hermann, *Rep Chim pur* 1861 53, Ferriehs a Smith, *A* 191, 346, Cleve, *Bl* [2] 43, 861).

Erbium nitrate $\text{Er}(\text{NO}_3)_3 \cdot 6\text{H}_2\text{O}$. Large crystals, sol water and alcohol. Decomposed by heat forming a basic salt $2\text{Er}_2\text{O}_3 \cdot 3\text{N}_2\text{O}_4 \cdot 9\text{H}_2\text{O}$ (Höglund, *Bl* [2] 18, 193, 279, Cleve, *C R* 91, 881).

Gallium nitrate $\text{Ga}(\text{NO}_3)_3$. Obtained by dissolving Ga in HNO_3Aq , evaporating at 100°, drying in an exsiccator, and heating in a dry air stream to 40°. Decomposes at 110°, and at 200° leaves Ga_2O_3 (de Boisbaudran).

Gold nitrates By dissolving $\text{Au}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ in HNO_3Aq , a very unstable salt is obtained, Schottlander gives the formula $5(\text{AuO NO}_3) \cdot \text{H}_2\text{O}$ as approximately correct (*A* 217, 812). The compound $\text{Au}(\text{NO}_3)_3 \cdot 3\text{H}_2\text{O}$, which may be called *auronitric acid*, is obtained in large crystals by dissolving $\text{Au}_2\text{O}_3 \cdot x\text{H}_2\text{O}$ in HNO_3Aq with special precautions. This compound forms shining, yellow, tridinic octahedra, SG 2.84, it is readily decomposed by heat to $2\text{Au}_2\text{O}_3 \cdot \text{N}_2\text{O}_4 \cdot 2\text{H}_2\text{O}$ (Schottlander, *A* 217, 312). Several salts are known derived from auro nitric acid, they are obtained by dissolving HNO_3 , $\text{Au}(\text{NO}_3)_3$, along with various nitrates in HNO_3Aq , and evaporating. The K salts are $\text{KAu}(\text{NO}_3)_4$ and $\text{HK}_2\text{Au}(\text{NO}_3)_6$ (Schottlander, *lc*).

Indium nitrate $2\text{In}(\text{NO}_3)_3 \cdot 9\text{H}_2\text{O}$. Large needle shaped crystals, by dissolving excess of In in HNO_3Aq , and evaporating in an exsiccator. Loses $6\text{H}_2\text{O}$ at 100° at red heat forms a basic salt, then In_2O_3 (Winkler, *J pr* 94, 1, 102, 273).

Iron nitrates According to Scheurer Kestner (*C R* 47, 927), Fe reacts with HNO_3Aq SG 1.034 to form $\text{Fe}(\text{NO}_3)_3$ and NH_4NO_3 , with acid SG 1.073 $\text{Fe}(\text{NO}_3)_3$ is also formed, with acid SG 1.115 only $\text{Fe}(\text{NO}_3)_3$ is produced, and with more conc acid basic salts begin to be formed.

Ferrous nitrate $\text{Fe}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ is best prepared by dissolving FeS in cooled HNO_3Aq SG less than 1.12, the liquid is made as nearly neutral as possible by addition of FeS, decanted, evaporated slowly, and strongly cooled. The crystals may be kept unchanged at low temperatures in the mother liquor. S 200 at 0°, 800 at 25° (Ordway, *Am S* [2] 40, 325).

Ferric nitrate $\text{Fe}(\text{NO}_3)_3 \cdot 9\text{H}_2\text{O}$ (Ordway, *Am S* [2] 26, 197, 27, 14). Prepared by dissolving Fe in HNO_3Aq SG 1.29 till about 10 p c Fe is taken up by the acid, then adding an equal volume of HNO_3Aq SG 1.43, oblique rhombic prisms are deposited on cooling. SG 1.6885 at 20°, nearly colourless, slightly deliquescent, very soluble in water, very slightly soluble in cold HNO_3Aq , melts 47.2°, acid begins to be given off at 100°, boils 125°, completely decomposed at red heat (Ordway, *lc*). Hausmann obtained $\text{Fe}(\text{NO}_3)_3 \cdot 6\text{H}_2\text{O}$ (*A* 89, 109; v also Widenstein, *J pr* 84, 243). Scheurer Kestner obtained a dihydrated salt (*J* 1862, 198).

Numerous *basic ferric nitrates* were obtained by Ordway (*l.c.*) by dissolving FeO_2H_2 in $\text{Fe}(\text{NO}_3)_3\text{Aq}$, and evaporating (*v* also Hausmann, *A* 89, 109, and Scheurer-Kestner, *J*, 1862 193) Basic salts are also produced by heating $\text{Fe}(\text{NO}_3)_3\text{Aq}$ Basic ferric nitrates are slowly resolved by boiling water to normal salt and Fe_2O_3 , the change proceeds most rapidly by heating in a sealed tube

Several *ferric aceto nitrates*, e.g. $\text{Fe}(\text{NO}_3)(\text{C}_2\text{H}_3\text{O}_2)_2 \cdot 3\text{H}_2\text{O}$, $\text{Fe}(\text{NO}_3)_2(\text{C}_2\text{H}_3\text{O}_2) \cdot 8\text{H}_2\text{O}$ have been prepared and described by Scheurer-Kestner (*A Ch* [3] 63, 422)

Lanthanum nitrate $\text{La}(\text{NO}_3)_3 \cdot 6\text{H}_2\text{O}$ Large prismatic, deliquescent, crystals, easily sol water and alcohol May be fused without decomposition at c 40° , but at rather higher temperature HNO_3 is removed and a basic salt formed completely decomposed at red heat (Ordway) Two *double salts*, $2\text{La}(\text{NO}_3)_3 \cdot 3\text{Ni}(\text{NO}_3)_2 \cdot 36\text{H}_2\text{O}$, and $2\text{La}(\text{NO}_3)_3 \cdot 3\text{Zn}(\text{NO}_3)_2 \cdot 69\text{H}_2\text{O}$, are described by Frerichs & Smith (*A* 191, 559)

Lead nitrates The normal salt, $\text{Pb}(\text{NO}_3)_2$, crystallises in octahedra from a solution of PbO or PbCO_3 in boiling very dilute HNO_3Aq *S G* 4 472 at 4° (Playfair & Joule, *C J* 1, 137), 4 41 at 15° (Holker, *P M* [3] 27, 214, *v* also Schroder, *P* 106, 226, Ditte, *B* 15, 1438) *H F* [Pb , N , O] = 105,500, [Pb , O , $\text{N}^\circ\text{O}^\circ$] = 109,470, [Pb , O , $\text{N}^\circ\text{O}^\circ\text{Aq}$] = 68,070 (*Th* 3, 518) *S* 39 at 0° , 48 3 at 10° , 60 6 at 25° , 80 at 45° , 101 at 65° , 120 5 at 85° , 138 9 at 100° (Kremers, *P* 92, 497) *S* in alcohol, *S G* 9282, 4 96 at 4° , 5 82 at 8° , 8 77 at 22° , 12 8 at 40° , 11 49 at 50° (Gerardin, *A Ch* [4] 5, 129) Insol conc HNO_3Aq $\text{Pb}(\text{NO}_3)_2$ is decomposed at low red heat giving PbO , O , and NO_2 In HCl gas, PbCl_2 is formed with evolution of NO and Cl (Thomas, *C J* 33, 367) Forms a compound with lead phosphate, viz $\text{Pb}(\text{NO}_3)_2 \cdot \text{Pb}_3(\text{PO}_4)_2 \cdot 2\text{H}_2\text{O}$ (Gerhardt, *A* 68, 286)

Many *basic lead nitrates* have been described The salt $2\text{PbO} \cdot \text{N}_2\text{O}_5 \cdot \text{H}_2\text{O}$, which formula may be written PbOHNO_3 , is obtained by boiling $\text{Pb}(\text{NO}_3)_2\text{Aq}$ with PbO , filtering hot, and allowing to cool (Berzelius, *P* 19, 312, Pelouze, *J p* 25, 486, Persoz, *A Ch* [3] 68, 191) *S G* 5 93 (Ditte, *C R* 94, 1180) Several other basic salts are known, according to Wakeman & Wells (*Am* 9, 299) the only recrystallisable basic salt, besides PbOHNO_3 , is $10\text{PbO} \cdot 3\text{N}_2\text{O}_5 \cdot 5\text{H}_2\text{O}$

Lithium nitrate LiNO_3 By neutralising HNO_3Aq with LiOH or Li_2CO_3 , and evaporating Rhombic prisms, *S G* 2 334 (Kremers, *P* 92, 520) *H F* [Li , N , O°] = 111,615, [Li , O , NO°] = 113,620, [Li°O , $\text{N}^\circ\text{O}^\circ\text{Aq}$] = 97,005 (*Th* 3, 518)

Melts at 264° (Carnelley, *C J* 33, 275) Easily sol water and alcohol Kremers (*P* 114, 41) gives following table —

<i>S G</i> LiNO_3Aq at $19^\circ 5'$	<i>P c</i> LiNO_3
1 0769	14 2
1 1346	26 7
1 193	40 6
1 255	57 5
1 3154	77 4

LiNO_3 , slowly reacts with dry HCl , a small quantity of Cl and NO being evolved (Thomas, (*J* 33, 370)

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The *hydrated salt* $\text{LiNO}_3 \cdot 5\text{H}_2\text{O}$ is said to be obtained by crystallising below 10° (Troost, *A Ch* [3] 51, 184)

Magnesium nitrate $\text{MgNO}_3 \cdot 6\text{H}_2\text{O}$ Occurs in mother liquor from saltpetre plantations, also in some well-waters of Stockholm, according to Berzelius Prepared by neutralising HNO_3Aq by *magnesia alba*, and evaporating Very deliquescent monoclinic crystals (Maignac, *J* 1856 336) *S G* 1 464 (Playfair & Joule, *C S Mem* 2, 401) *H F* [Mg , O° , $\text{N}^\circ\text{O}^\circ \cdot 6\text{H}_2\text{O}$] = 214,530 [Mg , O , $\text{N}^\circ\text{O}^\circ\text{Aq}$] = 176,480 (*Th* 3, 518) Very soluble water and alcohol Oudemans (*Er* 7, 419) gives the table —

<i>P c</i> $\text{Mg}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$	<i>S G</i> $\text{Mg}(\text{NO}_3)_2\text{Aq}$
1	1 0034
5	1 0202
10	1 0418
15	1 0639
20	1 0869
25	1 1103
30	1 1347
35	1 1649
40	1 1909
45	1 2176
49	1 2397

According to Graham (*T* 1837 47), $5\text{H}_2\text{O}$ are removed from $\text{Mg}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ at c 330° (M.P. of lead), and the monohydrated salt can be fused without change, but is decomposed at red heat, leaving MgO Einbrodt (*A* 65, 115) found that acid began to be evolved before five sixths of the water was removed, hence he regarded the existence of $\text{Mg}(\text{NO}_3)_2 \cdot \text{H}_2\text{O}$ as very doubtful By heating $\text{Mg}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ until water ceases to come off, a *basic salt* $3\text{MgO} \cdot \text{N}_2\text{O}_5$ is obtained, according to Chodnew (*A* 71, 241) Reacts with HCl gas to form MgCl_2 , Cl and O and H_2O being evolved (Thomas, *C J* 33, 370)

Manganese nitrate $\text{Mn}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ Small monoclinic crystals (Hannay, *C J* 33, 269) Obtained by dissolving MnCO_3 in HNO_3Aq , also by dissolving MnO_2 in HNO_3Aq in sunlight or presence of deoxidisers, and evaporating *S G* 1 8199 when solid at 21° , 1 8104 when liquid at 21° (Ordway, *J* 12, 113)

H F [Mn , O , $\text{N}^\circ\text{O}^\circ \cdot 6\text{H}_2\text{O}$] = 157,700, [Mn , O , $\text{N}^\circ\text{O}^\circ\text{Aq}$] = 117,720 (*Th* 3, 518) Decomposed by heat, giving MnO_2 , Mn_2O_3 , or Mn_3O_4 , according to the temperature Reacts with HCl gas to form MnCl_2 , with evolution of Cl and NO (Thomas, *C J* 33, 370) According to Schultz Sellac (*Z* 1870 646) the salt $\text{Mn}(\text{NO}_3)_2 \cdot 3\text{H}_2\text{O}$ crystallises from solution in conc HNO_3Aq

Mercury nitrates *Mercuric nitrate*, $2\text{Hg}(\text{NO}_3)_2 \cdot \text{H}_2\text{O}$, is obtained by dissolving HgO in excess of slightly warmed HNO_3Aq , and evaporating over H_2SO_4 , after some minutes the liquid above the crystals has the composition $\text{Hg}(\text{NO}_3)_2 \cdot 2\text{H}_2\text{O}$ (Millon, *A Ch* [3] 18, 361) Ditte (*J* 1854 366) obtained $\text{Hg}(\text{NO}_3)_2 \cdot 8\text{H}_2\text{O}$ by cooling to -15° a nearly neutral conc solution of HgO in HNO_3Aq

Basic mercuric nitrates are readily formed by heating the normal salt, the chief are $2\text{HgO} \cdot \text{N}_2\text{O}_5 \cdot 8\text{H}_2\text{O}$ (Ditte, *l.c.*), $2\text{HgO} \cdot \text{N}_2\text{O}_5 \cdot 2\text{H}_2\text{O}$ (Maignac, *J* 1855 415), $3\text{HgO} \cdot \text{N}_2\text{O}_5 \cdot \text{H}_2\text{O}$, obtained by the prolonged action of water on any of the other basic salts

Mercuric nitrate forms several *double salts*

Li Li

With HgI_2 , the compounds $\text{Hg}(\text{NO}_3)_2$, HgI_2 , $\text{Hg}(\text{NO}_3)_2 \cdot 2\text{HgI}_2$, and $2\text{Hg}(\text{NO}_3)_2 \cdot 3\text{HgI}_2$ are formed (Preuss, *A* 29, 326; Liebig, *A* 72, 79). These iodo nitrates are decomposed by water, with separation of HgI_2 and solution of $\text{Hg}(\text{NO}_3)_2$. With HgS , the compound $\text{Hg}(\text{NO}_3)_2 \cdot 2\text{HgS}$ is formed, by digesting freshly pptd HgS with $\text{Hg}(\text{NO}_3)_2$ aq, also by passing into $\text{Hg}(\text{NO}_3)_2$ aq less H_2S than suffices to decompose it wholly.

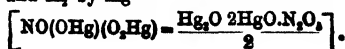
Mercurous nitrate The nitrates derived from Hg_2O have been examined chiefly by Mitscherlich (*P* 9, 387), Lefort (*A* 56, 247), Margnac (*A Ch* [3] 27, 332), and Gerhardt (*A* 72, 74). Hg_2NO_3 is formed by the reaction of excess of Hg with HNO_3 aq, but if the action is continued basic salts are produced. Basic salts are also obtained by decomposing HgNO_3 by water.

The normal salt, $\text{HgNO}_3 \cdot 2\text{H}_2\text{O}$, is obtained in colourless monoclinic crystals by reacting on excess of Hg with cold HNO_3 aq S G c 12. To prevent admixture of basic salts it is advisable to allow the acid and Hg to remain in contact until crystals cease to be formed, then to warm gently, filter, and allow to crystallise S G 478 (Playfair a Joule, *C S Mem* 2, 401). The crystals effloresce somewhat in air, they are dissolved without change in a little water, but dilution produces basic salts. By heating HgNO_3 , with water HgO and NO_2 are formed. HgCl is formed when HCl is passed over HgNO_3 , and Cl and NO are evolved (Thomas, *C J* 33, 370).

Basic mercurous nitrates are formed by warming with Hg the mother liquor from the preparation of the normal salt, and by treating the normal salt with water. To the salt obtained by the first of these methods Gerhardt gave the formula $3\text{Hg}_2\text{O} \cdot 2\text{N}_2\text{O}_5 \cdot \text{H}_2\text{O}$, Margnac gave the formula $4\text{Hg}_2\text{O} \cdot 8\text{N}_2\text{O}_5 \cdot \text{H}_2\text{O}$. The other basic salts are (1) $2\text{Hg}_2\text{O} \cdot \text{N}_2\text{O}_5 \cdot \text{H}_2\text{O}$ (Gerhardt), obtained by the action of a little boiling water on HgNO_3 , Margnac formulates this salt as $5\text{Hg}_2\text{O} \cdot 8\text{N}_2\text{O}_5 \cdot 2\text{H}_2\text{O}$. (2) $2\text{Hg}_2\text{O} \cdot \text{N}_2\text{O}_5 \cdot \text{H}_2\text{O}$ (Gerhardt, Margnac), obtained by adding much water to HgNO_3 aq.

Mercurous nitrate forms double salts with NH_4NO_3 , $\text{Pb}(\text{NO}_3)_2$, $\text{Ba}(\text{NO}_3)_2$, and $\text{Sr}(\text{NO}_3)_2$. The compositions of these salts are expressed by the formulæ $2\text{HgNO}_3 \cdot 4\text{NH}_4\text{NO}_3 \cdot 5\text{H}_2\text{O}$ (Rammelsberg, *P* 109, 397), $2\text{M}(\text{NO}_3)_2 \cdot 2\text{Hg}_2\text{O} \cdot \text{N}_2\text{O}_5$, where $\text{M} = \text{Pb}$, Ba , or Sr (Städeler, *A* 87, 129).

Mercurous-mercuric nitrate, $\text{Hg}_2\text{O} \cdot 2\text{HgO} \cdot \text{N}_2\text{O}_5$, This salt is formed by the gradual oxidation of HgNO_3 in air. It is best obtained by boiling 1½ parts HNO_3 aq, S G 12, with 1 part Hg till all Hg is dissolved, and maintaining the solution near its B P. The salt separates as a yellow powder, after a time a white basic mercurous nitrate begins to form (Wittstock, Gerhardt, *A* 72, 74; Brooks, *P* 66, 63). Rubbed with NaCl , HgCl and oxychloride are formed, and on addition of water HgCl_2 goes into solution. Treated with HCl gas both HgCl and HgCl_2 are formed with evolution of Cl and NO (Thomas, *C J* 33, 370). The salt may be regarded as derived from orthoionic acid— $\text{NO}(\text{OH})_2$ —by replacing H by Hg^1 and H_2 by Hg^{11} .



Nickel nitrates The normal salt, $\text{Ni}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$, is obtained, in emerald-green, deliquescent, monoclinic crystals, by dissolving Ni , NiO , or NiCO_3 , in HNO_3 aq, and evaporating. Melts at 56° , and boils at 136° , remaining clear till $3\text{H}_2\text{O}$ is gone off (Ordway, *Am S* [2] 26, 197, 27, 14). S G 2 066 at 14° , 2 037 at 22° (Clarke's *Specific Gravity Table* [new ed], 112). H F $[\text{Ni}, \text{O}^2, \text{N}^2\text{O}^4, 6\text{H}^1\text{O}^1] = 124,720$, $[\text{Ni}, \text{O}, \text{N}^2\text{O}^4\text{Aq}] = 83,420$ (*Th* 3, 518). Several double salts of $\text{Ni}(\text{NO}_3)_2$ are known—with $\text{Ce}(\text{NO}_3)_3$, $\text{Di}(\text{NO}_3)_4$, and $\text{La}(\text{NO}_3)_3$ (v CERIUM NITRATES, DIDYMIUM NITRATE, LANTHANUM NITRATE). With ammonia forms

$\text{Ni}(\text{NO}_3)_2 \cdot 4\text{NH}_3 \cdot 2\text{H}_2\text{O}$
(Laurent, *A Ch* [3] 36, 354), and
 $\text{Ni}(\text{NO}_3)_2 \cdot 6\text{NH}_3 \cdot 1\frac{1}{2}\text{H}_2\text{O}$
(F Rose, *Ammon Kobaltverbind* [Heidelberg, 1871], 27). Also combines with nickel chloride and ammonia to form

$6(\text{Ni}(\text{NO}_3)_2 \cdot 4\text{NH}_3 \cdot \text{H}_2\text{O}) (\text{NiCl}_2 \cdot 6\text{NH}_3) \cdot 10\text{H}_2\text{O}$
(Schwarz, *W A B* 1850 272).

The basic salt $8\text{NiO} \cdot \text{N}_2\text{O}_5 \cdot 5\text{H}_2\text{O}$ is pptd as a white powder by adding boiling NH_3 aq to a solution of the normal salt (Habermann, *M* 5, 440).

Palladium nitrates The normal salt $\text{Pd}(\text{NO}_3)_2 \cdot x\text{H}_2\text{O}$ forms brown-yellow rhombic prisms, very deliquescent, obtained by dissolving Pd in cold HNO_3 aq, evaporating to a syrup at the ordinary temperature, and allowing to stand in a warm place (Fischer, *P* 10, 607). By dissolving the normal salt in water, and diluting, the Pd is gradually pptd as a basic salt. Basic salts are also obtained by evaporating Pd in HNO_3 aq at c 100° to 120° , and treating the residue with water (Fischer, *lc*, Kane, *B J* 24, 236).

Platinum nitrates A brown salt, probably $\text{Pt}(\text{NO}_3)_4$, is obtained by dissolving $\text{PtO}_2 \cdot x\text{H}_2\text{O}$ in HNO_3 aq, or by decomposing $\text{Pt}(\text{SO}_4)_2$ aq with $\text{Ba}(\text{NO}_3)_2$ aq, and evaporating (Berzelius).

Potassium nitrate KNO_3 (*Nitre Saltpetre*.) Melts at 339° (Carnelley, *C J* 33, 277). S G 2 0958 to 2 1078 at 4° (Playfair a Joule, *C J* 1, 137), 2 059 at 0° (Quinck, *P* 135, 642), 1 072 at M P (Braun, *P* 154, 190). H F

$[\text{K}, \text{O}, \text{NO}] = 121,485$, $\left[\frac{\text{K}^1, \text{O}_2, \text{N}^2\text{O}^4\text{Aq}}{2} \right] = 96,050$

(*Th* 3, 518). Heat of solution = -7967 at 15° -7814 at 34° , -7541 at 53° (Tilden, *Pr* 58, 401). S H 18° to $98^\circ = 23875$ (Regnault, *A Ch* [3] 1, 129). S 133 at 0° , 21 at 10° , 312 at 20° , 445 at 30° , 639 at 40° , 859 at 50° , 1109 at 60° , 139 at 70° , 172 at 80° , 206 at 90° , 247 at 100° (Mulder, *J* 1866 65, v also Tilden a Shen stone, *T* 175, 23). Schiff gives the following tables (*A* 107, 87, 293, for more extended tables v Gerlach, *Fr* 8, 286) —

Weight of alcohol in 100 parts	Weight of KNO_3 in 100 parts solution saturated at 15°
0	20.5
10	13.2
20	8.5
30	5.6
40	4.3
50	2.8
60	1.7
80	0.4

S.G. of KNO_3Aq at 21°	Weight of KNO_3 in 100 parts solution
1.1688	24.93
1.1078	16.62
1.0695	11.08
1.051	8.81
1.0337	5.54
1.017	2.77

S in glycerin S.G. 1.225 = 10 (Vogel, *N. R. P.* 16, 557) KNO_3 is dimorphous, it usually crystallises in trimetric prisms, $a:b:c = 589:1:701$, if a drop of KNO_3Aq is allowed to crystallise slowly under the microscope, rhombohedral crystals are formed (Frankenheim, *P.* 92, 354) If the rhombohedral crystals are touched by a prismatic crystal while the crystallisation is proceeding, they are changed to prismatic, the prismatic may be changed to rhombohedral by heating nearly to the melting point

Occurrence—In small quantities in all vegetable soils, also in most spring and river waters (Boussingault, *C. R.* 44, 108) Nitre is found in the soil of caves, in different parts of the world, wherein animal or vegetable matter undergoes putrefaction, and where alkalis or alkaline earths are present to combine with the nitric acid produced (*v.* NITRIFICATION, p. 521) Nitre is also found as an efflorescence on the surface of the soil in parts of India, Arabia, South America, and other warm countries, the percentage of KNO_3 in a Bengalese soil was found by Davy to be 83 KNO_3 occurs in the juices of certain plants, notably in the leaves of the castor oil plant

Formation—1 By the oxidation of nitrogenous matter in presence of air, moisture, and potash (*cf.* NITRIFICATION, p. 521)—2 By the action of K_2CO_3 or KOH on $\text{Ca}(\text{NO}_3)_2$ or NaNO_3 —3 By the oxidation of NH_3 in presence of moisture, air, and ferric oxide, and combination of the HNO_3 formed with KOH (Pesci, *G.* 1875 307)

In the artificial preparation of nitre by oxidation of nitrogenous matter in soils, the first step is to prepare a soil rich in N containing materials, this is done by mixing porous soil, preferably that left from the lixiviation of a former nitre bed, with farm yard manure, animal and vegetable refuse, and wood ashes or calcareous matter, and watering this with urine This soil is then formed into a mound under a shed, and the process of nitrification is allowed to proceed for perhaps a couple of years, air must be freely admitted, and great care must be taken to keep the soil neither too wet nor too dry About 5lbs crude nitre are obtained, on an average, from 1,000lbs of such soil (for more details *v.* DICTIONARY OF APPLIED CHEMISTRY)

Preparation—1 By purifying crude nitre prepared from the washings of *saltpetre earth* The liquor from the *saltpetre earth* contains $\text{Ca}(\text{NO}_3)_2$, $\text{Mg}(\text{NO}_3)_2$, with smaller quantities of KNO_3 and NaNO_3 , and also alkaline chlorides, K_2CO_3 (wood ashes) is added, the liquid is filtered and evaporated, the crude KNO_3 is then treated with enough boiling water to dissolve all KNO_3 but not all the NaCl present—S of KNO_3 at $100^\circ = 247$, S of NaCl at $100^\circ = 89$ —the liquid is boiled for a considerable time, when NaCl separates with CaCO_3 and MgCO_3 , the liquid is then run off, and while crystallising it

is constantly stirred to insure formation of small crystals containing little mother liquor, the crystals are recrystallised, washed with saturated KNO_3Aq (to dissolve the last traces of alkaline chlorides), and again crystallised.—2 From *Chili saltpetre* (NaNO_3), by adding this salt to hot conc $\text{K}_2\text{CO}_3\text{Aq}$, when Na_2CO_3 separates and KNO_3 remains in solution The mother-liquor is evaporated as long as Na_2CO_3 continues to separate, then run off and allowed to crystallise, with constant stirring—3 By neutralising pure HNO_3Aq with pure KOHAq or $\text{K}_2\text{CO}_3\text{Aq}$, evaporating, and crystallising

Properties—A white, crystalline salt, dimorphous, S.G. 2.1, easily sol water, solution tastes cool and bitter, melts below red heat to a colourless liquid which solidifies on cooling to a white fibrous mass, known as *mineral crystal* or *salprunella* At red heat evolves O, and N as temperature increases Deflagrates when heated with combustible bodies (for physical properties *v.* beginning of article)

Reactions—1 Decomposed by heat, at $c 300^\circ\text{--}400^\circ$ O is evolved, and KNO_3 formed, as temperature increases N is evolved, and finally a mixture of K_2O and K_2O remains—2 Evolves O when heated in presence of combustible substances such as P, S, Zn, C, &c Hence the use of KNO_3 in gunpowder, and as an oxidiser when molten—3 Heated with copper foil, nearly pure K_2O is formed, a similar change occurs when KNO_3 is heated with iron—4 Most elements are oxidised by heating with KNO_3 , if the oxide produced is acidic, a K salt of the corresponding acid is produced—5 Organic compounds are generally burnt (to CO_2 and H_2O) by heating with KNO_3 —6 Reduced by the copper-zinc couple, in presence of water, at first to KNO_2 , and then to NH_3 , a similar reduction takes place by hydrogen occluded by Pd, Pt, or Cu (Gladstone & Tribe, *C. J.* 83, 139, 306)—7 Reacts with hydrogen chloride gas to form KCl , evolving Cl and N oxides (Thomas, *O. J.* 83, 367) For methods by which nitre may be valued approximately, or completely analysed, reference must be made to *Manuals of Analysis*, for an account of the technical applications of nitre reference should be made to DICTIONARY OF APPLIED CHEMISTRY

Rhodium nitrate $\text{Rh}(\text{NO}_3)_3 \cdot 2\text{H}_2\text{O}$ (?) A gum-like, very deliquescent, mass, obtained by dissolving Rh_2O_3 in HNO_3Aq , and evaporating at 100° till HNO_3 ceases to be evolved (Claus, *J. pr.* 34, 428, Berzelius)

Rubidium nitrate RbNO_3 Obtained in long needles, or six sided prisms (according to rate of evaporation) by neutralising HNO_3Aq by Rb_2CO_3 and evaporating S 20.1 at 0° , 43.5 at 10° When heated evolves O (Kirohoff & Bunsen, *P. M.* [4] 22, 55) According to Ditte (*C. R.* 89, 641) an acid salt, $2\text{RbNO}_3 \cdot 5\text{HNO}_3$, is obtained by dissolving RbNO_3 in HNO_3 , H_2O , this salt is decomposed by water or heat

Samarium nitrate $\text{Sm}(\text{NO}_3)_3 \cdot 6\text{H}_2\text{O}$ Pale yellow prisms, e sol water, S.G. 2.375 (Cleve, *C. N.* 48, 74, 51, 145)

Scandium nitrate $\text{Sc}(\text{NO}_3)_3$ (?) Small plates, by evaporating at 100° a solution of Sc_2O_3 in HNO_3Aq Decomposed by heat, giving a basic salt soluble in water (Nilson, *E.* 13, 1444)

Silver nitrate AgNO_3 , (*Lunar caustic*) S G 4 238 to 4 328 (Schröder, *P* 107, 113) S 121.9 at 0°, 227.3 at 19.5°, 500 at 54°, 714 at 85°, 1111 at 110° (Kremers, *P* 92, 497), S 1622.5 at 125°, 1941.4 at 133° (Tilden, *A* Shenstone, *T* 175, 28) Saturated solution boils at 125° S in boiling alcohol = 25 Melts at 218° (Carnelley, *C* J 33, 276) H F [Ag, O, NO] = 30,745, [Ag, N, O] = 28,740 (*Th* 3, 517) Crystallises in trimetric system, $a b c = 9433 \ 1137 \ 8 \text{ H}$ 16° to 99° = 14352 (Regnault, *A* *Ch* [3] 1, 129)

Preparation — Pure Ag is dissolved in HNO_3Aq , the solution is evaporated to dryness, the residue is heated gently till all HNO_3 is removed, when it is dissolved in water and crystallised. If solution of Ag in HNO_3Aq proceeds in the cold the liquid becomes blue from solution of N_2O_5 , but no gas is evolved, on warming, NO escapes rapidly. AgNO_3 may be prepared from Ag which contains Cu by saturating warm fairly conc HNO_3Aq with the metal, adding enough KOHAq to a part of the solution to ppt Ag_2O along with CuO, digesting the pp with the rest of the solution, whereby the remaining CuO is pptd, filtering, and evaporating the filtrate.

Properties — White trimetric crystals, solution in water is perfectly neutral to litmus paper, has a metallic taste, and is poisonous, melts below red heat, is a powerful caustic, at once destroying flesh when applied to it. AgNO_3 is readily reduced by organic matter in light.

Reactions — 1 Decomposed by heat, leaving Ag — 2 With hydrogen chloride gas forms AgCl and HNO_3 , with evolution of a very little Cl and NO (Thomas, *C* J 33, 371) — 3 With organic matter, e.g. paper, dust, &c., forms Ag in presence of sunlight — 4 AgNO_3Aq is slowly and very partially decomposed by hydrogen with ppn of Ag (Russell, *C* J [2] 12, 3) — 5 Uraous oxide, UO_3 , ppts Ag, with production of $\text{UO}_2(\text{NO}_3)_2$ (Isambert, *C* R 80, 1087)

Combinations — 1 With ammonia, to form $\text{AgNO}_2\cdot 2\text{NH}_3$, by supersaturating conc AgNO_3Aq with NH_3 , decomposes above 100°, giving off NH_3 and N (Maignac, *P* 9, 413, Mitscherlich, *A* *Ch* [2] 72, 288, Kane, *P* 20, 153) Dry AgNO_3 absorbs NH_3 to form $\text{AgNO}_3\cdot 3\text{NH}_3$ (H Rose, *J* 1857 256) — 2 With silver bromide, to form $\text{AgNO}_3\cdot \text{AgBr}$, by dissolving freshly pptd AgBr in very conc hot AgNO_3Aq (Risse, *A* 111, 42) AgCl seems to form a similar compound — 3 With silver iodide, to form several compounds. $2\text{AgNO}_3\cdot \text{AgI}$ is obtained by boiling very conc AgNO_3Aq with AgI, pouring off, and allowing to cool (Risse, *A* 171, 23, Riche, *A* 111, 89, cf. Weltzien, *A* 101, 127, Kremers, *J* pr 71, 54, Preuss, *A* 29, 329, Schnauss, *Ar* Ph [2] 82, 260, Hofmann, *A* 171, 23, Sturenberg, *Ar* Ph [2] 143, 12) — 4 With ammonium and potassium nitrates to form $\text{AgNO}_3\cdot \text{MNO}_3$ (M = NH₄ or K) (Ditte, *C* R 101, 878)

Sodium nitrate NaNO_3 , (*Cubic saltpetre*, *Ohln saltpetre*) S G 2 2606 at 4° (Playfair, *A* Joule, *C* S *Mem* 2, 401), 2 246 at 15° (Holker, *P* M [3] 27, 218), 1 878 at melting point (Braun, *P* 154, 190) Melts at c 216° (Carnelley, *C* J 33, 276) H F [Na, N, O] = 111,250, [Na, O, NO] = 118,255,

$\left[\frac{\text{Na}_2\text{O}, \text{N}_2\text{O}_5\text{Aq}}{2} \right] = 91,310$ (*Th* 3, 518) Heat

of solution — 4786 at 16°, — 4255 at 54° (Tilden, *Pr* 38, 401) S 72.9 at 0°, 80.8 at 10°, 87.5 at 20°, 94.9 at 30°, 102 at 40°, 112 at 50°, 122 at 60°, 134 at 70°, 148 at 80°, 162 at 90°, 180 at 100°, 200 at 110°, saturated solution freezes at — 17.5° (Rudorff, *B* 2, 68), contains 216.4 parts NaNO_3 in 100 water, and boils at 119.7° (Mulder, *J* 1866 65, cf. Ditte, *C* R 80, 1164, Maumené, *C* R 58, 81, 81, 107) Schiff (*A* 110, 75) gives a table showing S G and p.c. composition of NaNO_3Aq S in alcohol (61.4 p.c.) at 26° = 21.2 (Pohl, *W* A B 6, 600, v also Wittstein, *Vierteljahr Zeit Pharm* 12, 109) 75 parts NaNO_3 added to 100 parts water at 13.2° lower the temperature through 18.5° 50 parts NaNO_3 mixed with 100 parts snow lower the temperature to — 17.5°

Occurrence — In large quantities in Chili, Peru, and some other parts of South America.

Preparation — 1 The crude salt is purified by repeated solution and crystallisation, or it is recrystallised once or twice, the first portions of each crop of crystals being rejected, then heated with HNO_3Aq , whereby chlorides are transformed into nitrates and again crystallised. 2 By neutralising HNO_3Aq with NaOHAq or $\text{Na}_2\text{CO}_3\text{Aq}$, and evaporating.

Properties — White obtuse rhombohedrons, having much the aspect of cubes, hence the name *cubic saltpetre*. Absorbs water from moist air. Melts c 316°, and solidifies to white mass on cooling, decomposes at higher temperature similarly to, but more readily than, KNO_3 , deflagrates with charcoal, &c., but less quickly than KNO_3 .

Reactions — Decomposed by heat at c 350° — 380°, evolving O, and at higher temperature N also — 2 Heated in presence of oxidisable bodies, produces oxides, which, if acidic, form Na salts of corresponding acids — 3 With hydrogen chloride forms NaCl, and evolves NO and Cl (Thomas, *C* J 33, 367)

Sroutium nitrate $\text{Sr}(\text{NO}_3)_2$ Prepared by adding to SrCO_3 enough HNO_3Aq to dissolve nearly all the salt, filtering, and crystallising. Separates without water of crystallisation from hot conc solutions, from cold and more dilute solutions crystals of $\text{Sr}(\text{NO}_3)_2\cdot 4\text{H}_2\text{O}$ are obtained (Souchay, *A* Lessens, *A* 99, 45) The anhydrous salt crystallises in octahedrons, S G 2 98 at 16.8° (Favre, *A* Valsen, *C* R 77, 579) The hydrate forms triclinic crystals, $a b c = 5895 \ 1 \ 808$, S G 2 249 at 15.5° (F v, *l* c) H F [$\text{Sr}, \text{N}, \text{O}$] = 109,910 (*Th* 3, 517) S anhydrous salt 20 in cold water, 200 in boiling water. Melts at red heat with decomposition, leaving SrO. Does not react with HCl gas (Thomas, *C* J 33, 371) A compound with Sr acetate, viz $\text{Sr}(\text{NO}_3)_2\cdot \text{Sr}(\text{C}_2\text{H}_3\text{O}_2)_2\cdot 8\text{H}_2\text{O}$, is obtained by allowing a mixed solution of the salts to evaporate (von Hauer, *J* pr 74, 432)

Tellurium nitrate The compound $8\text{TeO}_2\cdot 2\text{N}_2\text{O}_5\cdot 8\text{H}_2\text{O}$ is obtained by dissolving Te in excess of hot HNO_3Aq S G greater than 1.15 the salt crystallises in forms which are probably orthorhombic, soluble in HNO_3Aq , easily decomposed by water with separation of TeO_2 , (Klein, *A* *Ch* [6] 5, 59)

Thallium nitrates *Thallous nitrate*, TiNO_3 , is obtained by dissolving Tl in not very conc HNO_3Aq , keeping as small an excess of acid as possible, and evaporating Rhombic prisms, S G 5 5 (Lamy a Des Cloizeaux, *N* 1, 116) S 10 6 at 15°, 43 5 at 58°, 588 at 107° (Crookes, *C* J 17, 141, Lamy, *C* R 54, 1255) Insol alcohol Melts at 205° without decomposition and solidifies to a glass S G 5 8 (Lamy, *l* c) H F $[\text{Ti}, \text{N}, \text{O}] = 58,150$ (*Th* 3, 517) Strongly heated in a crucible leaves TiNO_2 and Ti_2O_3 (Carstanjen, *J* pr 102, 65, 129)

Thallous nitrate $\text{Ti}(\text{NO}_3)_3$ Separates in large deliquescent crystals from solution of TiO OH in HNO_3Aq S G 1 4, according to Strecker (*A* 135, 207) the crystals contain $6\text{H}_2\text{O}$, and according to Willm (*A* Ch [4] 5, 5) $8\text{H}_2\text{O}$ Decomposed at c 100°

Thorium nitrate $\text{Th}(\text{NO}_3)_4 \cdot 12\text{H}_2\text{O}$ Large, translucent tables, very hygroscopic, lose $8\text{H}_2\text{O}$ over H_2SO_4 Forms a very soluble double salt with KNO_3

Tin nitrates No definite salt has been isolated Stannous oxide dissolves in very dilute cold HNO_3Aq , but the solution decomposes on heating with separation of SnO_2 (cf Ditté, *A* Ch [5] 27, 145) Sn dissolves in cold very dilute HNO_3Aq , with production of NH_4NO_3 and probably $\text{Sn}(\text{NO}_3)_2$ Freshly pptd SnO_2 dissolves in HNO_3Aq on heating SnO_2 is pptd

Titanium nitrate By evaporating a solution of TiO_2 in HNO_3Aq over lume, Merz obtained lustrous plates $5\text{TiO}_2 \cdot \text{N}_2\text{O} \cdot 6\text{H}_2\text{O}$, soluble in cold water (*J* pr 99, 157)

Uranium nitrate By dissolving U or an oxide of U in HNO_3Aq , and evaporating, large, yellow, rhombic crystals are deposited having the composition $\text{UO}_2(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O} = \text{uranyl nitrate}$ S G 2 807 (Boedeker) Effloresces somewhat in dry air, melts at 59 5°, and boils at 118° (Ordway, *J* 1859 114) A trihydrated salt, $\text{UO}_2(\text{NO}_3)_2 \cdot 3\text{H}_2\text{O}$, was obtained by Schultz-Sellack (*Z* [2] 6, 646) by evaporating a strongly acid solution of the ordinary salt over H_2SO_4 , and KOH Reacts with HCl to form U oxychloride with evolution of Cl and NO (Thomas, *C* J 33, 371)

Vanadium nitrates No definite salt has been isolated By dissolving VO or VO_2 in HNO_3Aq , a blue solution is obtained which cannot be evaporated without decomposition (Berzelius) By dissolving V_2O_5 in HNO_3Aq , and allowing to evaporate, a reddish residue is obtained (Berzelius)

Yttrium nitrate $\text{Y}(\text{NO}_3)_3 \cdot 6\text{H}_2\text{O}$ Large translucent crystals, by evaporating over H_2SO_4 , a solution of Y_2O_3 in HNO_3Aq (Cleve, *Bl* [2] 21, 344) Heated till NO_2 is evolved, the basic salt $2\text{Y}_2\text{O}_3 \cdot 3\text{N}_2\text{O} \cdot 9\text{H}_2\text{O}$ is obtained (*v* Bahr a Bunsen, *A* 137, 1)

Zinc nitrate $\text{Zn}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ Zn dissolves in HNO_3Aq with formation of NH_4NO_3 and $\text{Zn}(\text{NO}_3)_2$, from very dilute HNO_3Aq , N_2O is evolved The salt is prepared by evaporating a solution of Zn, ZnO , or ZnCO_3 , in HNO_3Aq Large 4 sided prisms, deliquescent, very soluble water and alcohol, melt at 36 4° and boil at 181° (Ordway, *Am* S [2] 27, 14), lose $2\text{H}_2\text{O}$ *in vacuo* over H_2SO_4 (Vogel a Reischauer, *N* J P 11, 137), lose all H_2O by heating to 105° in stream of dry air (Pierre, *A* Ch [3] 16, 247) S G. 2 063 at 18° (Clarke's *Table of Specific*

Gravities (new ed) 110) H F $[\text{Zn}, \text{O}, \text{N}^2\text{O}^4, 6\text{H}^2\text{O}] = 142,180$, $[\text{Zn}, \text{O}, \text{N}^2\text{O}^4\text{Aq}] = 102,510$ (*Th* 3, 518) The S G of $\text{Zn}(\text{NO}_3)_2\text{Aq}$ and p c of $\text{Zn}(\text{NO}_3)_2$ are given by Franz (*J* pr [2] 5, 274) —

S G	P c	S G	P c
1 0496	5	1 3268	30
1 0968	10	1 3906	35
1 1476	15	1 4572	40
1 2024	20	1 5258	45
1 264	25	1 5984	50

$\text{Zn}(\text{NO}_3)_2$ with HCl gas gives ZnCl_2 , evolving Cl and NO (Thomas, *C* J 33, 371) Many basic nitrates of zinc are known, they are obtained by heating the normal salt, by treating HNO_3Aq with a large excess of Zn, and by digesting hot $\text{Zn}(\text{NO}_3)_2$ with ZnO (*v* Schindler, *N* J P 11, 137, Vogel a Reischauer, *ibid*, Ordway, *Am* S [2] 27, 14, Grouvelle, *A* Ch [3] 19, 137, Wells, *Am* 9, 304)

Zirconium nitrates The normal salt $\text{Zr}(\text{NO}_3)_4$ is said to be obtained by evaporating a solution of $\text{ZrO}_2 \cdot x\text{H}_2\text{O}$ in excess of HNO_3Aq , by heating to 100°, the pyro salt ZrN_2O_5 is formed, and by boiling a dilute aqueous solution of this, the basic salt $3\text{ZrO} \cdot 2\text{N}_2\text{O}_5$ is produced

M M P M

NITRATION v NITRO COMPOUNDS
NITRIC ACID HNO_3 (*Spirit of nitre* When dilute, *aqua fortis*) Mol w 62 89 (*v infra*) [c -47°] (Berthelot, *Bl* [2] 29, 3) (86°) (Mitscherlich, *P* 18, 152, Millon, *J* pr 29, 337) S G 1 552 at 12 5° (Mitscherlich, *l* c), 1 552 at 15° (Millon, *l* c) V D (mixed with dry air) at 40 5° = 34 3 (Playfair a Wanklyn, *C* J 15, 142, *v infra*) S H of HNO_3 , $10\text{H}_2\text{O} = 768$, HNO_3 , $20\text{H}_2\text{O} = 849$, HNO_3 , $50\text{H}_2\text{O} = 93$, HNO_3 , $100\text{H}_2\text{O} = 963$, HNO_3 , $200\text{H}_2\text{O} = 982$ (Thomsen, *P* 142, 337) H F $[\text{H}, \text{N}, \text{O}] = 41,510$, $[\text{H}, \text{O}, \text{N}^2\text{O}] = 43,615$, $[\text{H}, \text{O}^2, \text{N}^2\text{O}] = 63,085$,

$$\left[\frac{\text{N}^2\text{O}, \text{H}^2\text{O}}{2} \right] = 7,330,$$

$$\left[\frac{\text{N}, \text{O}, \text{O}^2, \text{H}^2\text{O}}{2} \right] = 16,200;$$

$$\left[\frac{2\text{NO}, \text{O}^2, \text{H}^2\text{O}}{2} \right] = 28,905,$$

$$\left[\frac{2\text{NO}^2, \text{O}, \text{H}^2\text{O}}{2} \right] = 9,335,$$

$[\text{N}^2\text{O}^2, \text{Aq}] = 29,820$, $[\text{N}^2\text{O}^2\text{Aq}, \text{O}^2] = 36,640$, $[2\text{NO}^2, \text{O}, \text{Aq}] = 33,830$, $[\text{N}^2\text{O}, \text{O}^2, \text{Aq}] = 47,560$, $[2\text{NO}, \text{O}^2, \text{Aq}] = 72,970$, $[\text{H}, \text{N}, \text{O}^2, \text{Aq}] = 49,090$, $[\text{HNO}^2\text{Aq}, \text{O}] = 18,820$ (*Th* 2, 199) H V 7250 (Berthelot, *C* R 90, 1510) For E C of HNO_3Aq at different dilutions *v* Ostwald (*J* pr [2] 32, 300) Affinity of HNO_3Aq approximately the same as HClAq M M 1 18 (Perkin, *C* J 55, 680)

Occurrence—In small quantities in rain-water, varying from c 1 to c 16 pts per million pts of rain, for measurements *v* Goppelsröder, (*Fr* 10, 259, 11, 16), Warrington (*C* J 55, 537, references are given to various other results) Ekin (*C* J [2] 9, 64) found HNO_3 in the spring water of an uncultivated hill near Bath, not exposed to contact with organic matter Salts of HNO_3 occur in almost all soils, although not generally in large quantities, and also in the juices of many plants

Nitric acid has been known from early times. Geber mentions it (8th cent), Glauber (17th cent) gave directions for its preparation from

nitric by the use of H_2SO_4 . Lavoisier showed that this acid contained O. Cavendish proved the presence of N in it, and obtained it by passing electric sparks through moist O and N.

Formation—1 By burning H in O in presence of air (Lavoisier, Kolbe, *A* 119, 176, Hofmann, *B* 8, 668). The experiments of L. T. Wright (*C* *J* 85, 42) tend to show, although they do not absolutely prove, that the HNO_2 is formed by oxidation of NH_3 , not of N—2 By passing electric sparks for some time through a mixture of moist N and O (Böttger, *J pr* 73, 494, Perrot, *C R* 49, 204, Buff, a Hofmann, *A* 113, 140)—3 By exploding air with electrolytic gas (H_2 + O) (Bunsen). Hempel's experiments (*B* 23, 1456) show that HNO_2 is formed when air is burnt with O and electrolytic gas under pressure of several atmos. Hempel also proves that considerable quantities of HNO_2 are formed when O is burnt in compressed air in presence of strongly compressed O—4 By burning air in coal-gas (Ilosva, *Bl* [3] 2, 734), or coal gas in air (Wright, *C J* 36, 42)—5 According to Berthelot (*C R* 108, 543), a very little HNO_2 is formed when ether and P are slowly oxidised by air in presence of water—6 Nitrates are formed by the oxidation of nitrogenous animal or vegetable matter in the soil (*v Nitratification*)—7 Ozone produces NH_4NO_3 from NH_3 , and NH_4NO_2 easily oxidises to NH_4NO_3 (Carius, *A* 174, 31, Schönbein, *J pr* 75, 99, Weith, *A* Weber, *B* 7, 1745, Wöhler, *A* 136, 256)—8 By oxidising NH_3 in presence of moisture, e.g. by passing NH_3 and air over Pt black heated to c. 300° , by distilling $(\text{NH}_4)_2\text{SO}_4$ and conc H_2SO_4 with $\text{K}_2\text{Cr}_2\text{O}_7$, or distilling $(\text{NH}_4)_2\text{SO}_4$ with KMnO_4 and dilute H_2SO_4 (Tessie du Mothay, *W J* 1871 260)—9 HNO_2 is a product of the reaction of NO_2 with H_2SO_4 , and NaNO_2 is formed by acting on NaOH with N_2O_4 (Lunge, *B* 12, 1058)—10 By heating MnCl_2 or MnSO_4 with NaNO_2 (Kuhlmann, *W J* 1862 239)—11 By decomposing $\text{Ba}(\text{NO}_3)_2$ or $\text{Pb}(\text{NO}_3)_2$ with H_2SO_4 , filtering, and concentrating by evaporation—12 By heating NaNO_2 with AlO_3H_2 or SiO_2 (Wagner, *D P J* 183, 76).

Preparation—1 A mixture of 101 pts thoroughly purified and dry KNO_3 , in coarse powder, and 98 pts pure H_2SO_4 , is heated in a capacious glass retort, the neck of which passes some way into a glass receiver which is kept cold. The distillate is again distilled till one-third has passed over, when a quantity of conc H_2SO_4 equal to the contents of the retort is added (when the retort is cold), the receiver is changed, and distillation is continued, the distillate is again distilled at as low a temperature as possible, to get rid of H_2SO_4 , the distillate thus obtained is gently warmed, and a current of perfectly dry CO_2 is passed through it until it is quite colourless, oxides of N are thus removed. HNO_2 may be removed by distilling with a little urea, which decomposes HNO_2 , giving CO_2 , H_2O , and N.—2 Commercial acid is distilled, after addition of a little KNO_3 , to decompose H_2SO_4 , until a few drops give no pp, when diluted, with AgNO_3 , the receiver is then changed and distillation continued.

Preparation of fuming nitric acid.—The red fuming acid consists of a solution of NO_2 in HNO_3 , it is a more powerful oxidiser than the

ordinary acid. It is prepared by heating a. 202 pts KNO_3 , with 98 pts H_2SO_4 , and continuing the distillation as long as liquid comes over, the second stage of the process, which is represented by the equation $\text{KHSO}_4 + \text{KNO}_3 = \text{K}_2\text{SO}_4 + \text{HNO}_3$, occurs only at a temperature so high that part of the HNO_3 is decomposed with formation of NO_2 . A better method is to arrange the materials so that a portion of the HNO_3 is reduced to NO , as soon as it is formed; this may be done by mixing 100 pts KNO_3 with $3\frac{1}{2}$ pts starch, placing the mixture in a large retort the beak of which passes inside a glass tube 3 to 4 feet long which tube dips into a glass receiver kept very cold, and adding 100 pts H_2SO_4 , *S G* 1 85. The process goes on almost without the application of heat. About 60 pts red fuming acid are obtained from 100 pts KNO_3 (Brunner, *Rép Chim app* 3, 188).

Properties.—Perfectly pure HNO_3 , has not been isolated, Roscoe obtained an acid with from 99.5 to 99.8 p.c. HNO_3 (*A* 116, 211). Nitric acid is a colourless, highly corrosive liquid, according to Berthelot it solidifies at c. -47° , the B.P. is 86° , but decomposition begins below this temperature, at c. 256° the change $2\text{HNO}_3 = 2\text{NO}_2 + \text{H}_2\text{O} + \text{O}$ is complete. The following table shows the process of decomposition by heat (Carius, *B* 4, 828) —

t	S.G. of vapour		Po decom position	cc of O from 1 g HNO_3
	Air=1	H=1		
86°	2.05	29.6	9.53	8.43
100	2.02	29.1	11.77	10.41
130	1.92	27.6	18.78	16.62
160	1.79	25.8	28.96	26.22
190	1.59	23.0	49.34	43.69
220	1.42	20.4	72.07	63.79
250	1.29	18.6	93.03	82.30
266	1.25	18.0	100.0	88.47
265	1.24	17.9		
312	1.23	17.8		

By mixing the vapour from HNO_3 with dry air, Playfair a Wanklyn (*C J* 15, 142) found the V.D. at $40.5^\circ = 34.3$, that calculated from HNO_3 , being 31.5. HNO_3 is slowly and partially decomposed by sunlight, giving NO_2 , O, and H_2O , so that when exposed to sunlight the acid becomes yellow and contains NO_2 . Solution of HNO_3 in water, and dilution of the liquid, are accompanied by production of heat, Thomsen (*Th* 3, 66) gives the following table —

#	$[\text{HNO}_3, x\text{H}^+ \text{O}]$	$[\text{HNO}_3, x\text{H}^+ \text{O}, x\text{H}^+ \text{O}]$
5	2005	1280
1	3285	
15	4160	1550
25	5276	1389
3	5710	
5	6685	658
10	7818	140
20	7458	-22
40	7436	-15
80	7421	29
100	7439	
160	7450	45
320	7493	

If these results are plotted, and a curve drawn, with the horizontal lines showing molecules of H_2O and the vertical lines showing quantities of heat, the curve shows no signs of irregularity; hence, Thomsen concludes that the heat of solution and dilution of HNO_3 does not indicate the formation of any definite hydrates of HNO_3 . Considering the results which have followed a very close and extended examination of the heat of dilution of H_2SO_4 (v especially Pickering, *C J* 57, 64), it seems inadvisable to draw conclusions in favour of, or against, the formation of hydrates of HNO_3 , from the limited number of thermal observations made by Thomsen. Berthelot (*Bl* [2] 22, 530) has also measured the heat produced on adding water to HNO_3 , already diluted with known quantities of water, and concludes that a hydrate $HNO_3 \cdot 2H_2O$ exists in aqueous solutions of HNO_3 . Perkin has determined the magnetic rotatory power of HNO_3 and $HNO_3 \cdot xH_2O$ (*C J* 55, 680), his results are $-M M HNO_3 = 1.18$, $M M HNO_3 + 2.67H_2O = 3.656$. Now by deducting $2.67 (M M \text{ of } 2.67H_2O)$ from 3.656 , we get 986 as the $M M$ of HNO_3 , in presence of $2.67H_2O$, but pure HNO_3 gave $M M 1.18$, hence the water has reduced the $M M$ of HNO_3 by 194 . From these results, Perkin concludes that HNO_3 and H_2O combine to form H_3NO .

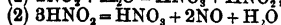
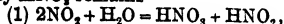
The electrical conductivity of HNO_3 Aq for increasing dilution has been measured by Kohlrausch a Grottrian (*P M* [4] 49, 417) and by Ostwald (*Z P C* 1, 74). Conductivity increases slightly with dilution, but soon reaches a maximum. The behaviour of HNO_3 Aq in this respect is characteristic of that of the strong monobasic acids (cf also Bouty, *C R* 106, 654). Roscoe (*C J* 13, 150) has shown that distilling HNO_3 Aq at 760 mm results in production of an acid containing 68 p.c. HNO_3 , the formula $2HNO_3 \cdot 3H_2O$ requires 70 p.c., and $HNO_3 \cdot 2H_2O$ requires 63.6 p.c. HNO_3 , the liquid of constant B.P. is therefore not a definite hydrate. This liquid boils at 120.5° under 735 mm pressure. By distilling at 150 mm pressure, Roscoe obtained an acid containing 67.6 p.c. HNO_3 , at 70 mm (B.P. $65^\circ-70^\circ$) the acid contained 66.7 p.c. HNO_3 , at pressure greater than 760 mm rather more than 68 p.c. HNO_3 was present in the acid of constant B.P. The percentage of HNO_3 in the liquid obtained by passing dry air into HNO_3 Aq containing from 64 to 68 p.c. HNO_3 , varies with temperature, the higher the temperature the greater the percentage of HNO_3 . For every mixture of HNO_3 and H_2O there is a fixed temperature, whereat HNO_3 and H_2O evaporate in the same proportion as they are present in the residual acid, for 66.2 p.c. HNO_3 , this temperature is 100° , for 64.5 p.c. it is 60° (Roscoe, *l.c.*, for older observations v Millon, *J pr* 29, 349, Smith, *Ph C* 1848 203). When HNO_3 and H_2O are mixed, the maximum contraction takes place for the ratio $2HNO_3 \cdot 3H_2O$ (Kolb, *A Ch* [4] 10, 140).

Kolb (*A Ch* [4] 10, 186) gives the following table, showing weight of HNO_3 is 100 pts of HNO_3 Aq (p) at 0° and 15° . The numbers marked with an asterisk were directly determined by adding a weighed excess of $CaCO_3$, and weighing the residue —

P	S.G.		Contraction
	at 0°	at 15°	
100 00	1 559	1 530	0 0000
99 84*	1 559*	1 530*	0 0004
90 42*	1 558*	1 530*	0 0010
99 52*	1 557*	1 529*	0 0014
97 89*	1 551*	1 523*	0 0065
97 00	1 548	1 520	0 0090
96 00	1 544	1 516	0 0120
95 27*	1 542*	1 514*	0 0142
94 00	1 537	1 509	0 0182
93 01*	1 533*	1 506*	0 0208
92 00	1 529	1 503	0 0242
91 00	1 526	1 499	0 0272
90 00	1 522	1 495	0 0301
89 56*	1 521*	1 494*	0 0315
88 00	1 514	1 488	0 0354
87 45*	1 513*	1 486*	0 0369
86 17*	1 507*	1 482	0 0404
85 00	1 503	1 478	0 0433
84 00	1 499	1 474	0 0459
83 00	1 495	1 470	0 0485
82 00	1 492	1 467	0 0508
80 96*	1 488*	1 463*	0 0531
80 00	1 484	1 460	0 0556
79 00	1 481	1 456	0 0580
77 66	1 476	1 451	0 0610
76 00	1 469	1 445	0 0643
75 00	1 465	1 442	0 0666
74 01*	1 462*	1 438*	0 0688
73 00	1 457	1 435	0 0708
72 39*	1 455*	1 432*	0 0722
71 24*	1 450*	1 429*	0 0740
69 96	1 444	1 423	0 0760
69 20*	1 441	1 419*	0 0771
68 00	1 435	1 414	0 0784
67 00	1 430	1 410	0 0796
66 00	1 425	1 405	0 0806
65 07	1 420*	1 400*	0 0818
64 00	1 415	1 395	0 0830
63 59	1 413	1 393	0 0833
62 00	1 404	1 386	0 0846
61 21*	1 400*	1 381*	0 0850
60 00	1 393	1 374	0 0854
59 59*	1 391*	1 372*	0 0855
58 88	1 387	1 368	0 0861
58 00	1 382	1 363	0 0864
57 00	1 376	1 358	0 0868
56 10*	1 371*	1 353*	0 0870
55 00	1 365	1 346	0 0874
54 00	1 359	1 341	0 0875
53 81	1 358	1 339	0 0875
53 00	1 353	1 335	0 0875
52 33*	1 349*	1 331*	0 0875
50 99	1 341*	1 323*	0 0872
49 97	1 334	1 317	0 0867
49 00	1 328	1 312	0 0862
48 00	1 321	1 304	0 0856
47 18*	1 315*	1 298*	0 0850
46 64	1 312	1 295	0 0848
45 00	1 300	1 284	0 0835
43 53*	1 291*	1 274*	0 0820
42 00	1 280	1 264	0 0808
41 00	1 274	1 257	0 0796
40 00	1 267	1 251	0 0786
39 00	1 260	1 244	0 0755
37 95*	1 253*	1 237*	0 0762
36 00	1 240	1 225	0 0740

P	S G		Contraction
	at 0°	at 15°	
35 00	1 234	1 218	0 0729
33 86*	1 226*	1 211*	0 0718
32 00	1 214	1 198	0 0692
31 00	1 207	1 192	0 0678
30 00	1 200	1 185	0 0664
29 00	1 194	1 179	0 0650
28 00*	1 187*	1 172*	0 0635*
27 00	1 180	1 166	0 0616
25 71	1 171*	1 157*	0 0593
23 00	1 153	1 138	0 0520
20 00	1 132	1 120	0 0483
17 47*	1 115	1 105	0 0422
15 00	1 099	1 089	0 0336
13 00	1 085	1 077	0 0316
11 41*	1 075	1 067*	0 0296
7 22*	1 050	1 045*	0 0206
4 00	1 026	1 022	0 0112
2 00	1 013	1 010	0 0055
0 00	1 000	1 999	0 0000

On adding water to the fuming red nitric acid, the liquid becomes green, then blue, and finally colourless when dilute, red vapour of NO_2 is evolved, the more rapidly the higher is the temperature. The NO_2 present in the fuming acid is slowly decomposed by the added water, giving HNO_3 , which colours the liquid blue (the green colour being the result of the yellow caused by NO_2 and the blue caused by HNO_2), on further dilution NO is evolved, and only HNO_3 remains



Reactions —1 *Heat* decomposes HNO_3 to NO_2 , H_2O , and O , the decomposition is complete at c 256° (Carius, *B* 4, 828) —2 *Light* slowly decomposes HNO_3 in the same way as heat, hence very conc HNO_3 kept in an open place always contains some NO_2 —3 The products of electrolysis of HNO_3 vary with the dilution of the acid, with acid 2HNO_3 , $3\text{H}_2\text{O}$ no H is evolved at first, after a little NO comes off, and then H begins, while the NO slowly ceases, as dilution increases H is evolved and products of reduction — N_2O , NO , N , and NH_3 —are produced, the more rapid the electrolysis the greater is the quantity of H evolved, very dilute acid evolves H only without the formation of reduced products (Bourgois, *J Ph* [4] 13, 266 [abstract in *C J* [2] 9, 885], Gladstone & Tribe, *C J* 35, 172) —4 Conc HNO_3 is rapidly reduced by *hydrogen occluded* by Pt or Pd , with oxidation of the H (G & T , *l c*) —5 HNO_3 is also reduced by many *metals*, the products vary with temperature, concentration of the acid, and the nature of the metal Mg with 68 p.c acid *plus* an equal quantity, or twice the quantity, of water, produces H along with gaseous reduction products (G & T , *C J* 35, 178) The gaseous reduction products are generally N_2O , NO , and N , these gases are formed by the reaction with HNO_3 of *e.g.* Al , Cd , Co , Cu , In , Fe , Pb , Mg , Ni , Ag , Ti , Sn , Zn (Acworth, *C J* 28, 828, Acworth & Armstrong, *C J* 32, 54) It appears to be the case that the greater the heat of formation of a metallic oxide the more completely is HNO_3 reduced by the

metal (*cf* Thomsen, *Th* 3, 547) Al , Cd , Fe , Mg , Pb , Sn , Zn , and the alkali metals produce NH_4NO_3 and NH_4OH , but no nitrous acid or nitrites, Bi , Cu , Hg , and Ag produce nitrites but no NH_4NO_3 or NH_4OH (Divers, *C J* 43, 443, for the combined action of HNO_3 and H_2SO_4 on Zn *v* Divers & Shimidzu, *C J* 47, 597) In most cases nitrates of the metals are formed, but sometimes these are decomposed with the final formation of oxides, *e.g.* Sb , Sn , and W (*v* also Veley, *Pr* 48, 458) Ta , Ti , Au , and most of the Pt metals do not react with HNO_3 —6 All the solid *non metals* are oxidised by nitric acid —7 *Oxidisable compounds* are oxidised by HNO_3 , *e.g.* ferrous compounds are changed to ferric, stannous to stannic, arsenious to arsenic, sulphides generally yield sulphates and nitrates —8 *Hydriodic acid* and *iodides* yield H_2O , NO , and I —9 *Hydrochloric acid* gives H_2O , NOCl , and Cl (*v* *aqua regia* under CHLORHYDRIC ACID, vol II p 8, *Reactions* No 17) —10 *Organic compounds* are oxidised by HNO_3 , straw, hay, cotton, &c., are in flamed by the conc acid (Kraut, *B* 14, 301) Many organic compounds, especially those of the benzenoid class, form nitro derivatives, H being replaced by NO . —11 With *starch*, N_2O_5 or a mixture of this with NO_2 is produced, HNO_3 S G 1.3 to 1.35 gives almost pure N_2O_5 , if the S G is greater than 1.35 NO_2 is also produced, if S G is less than 1.3 the chief gaseous product is NO (Lunge, *B* 11, 1229, 1641)

Combinations —1 With *water*, it is still undecided whether a hydrate or hydrates of HNO_3 are produced when HNO_3 is dissolved in water, *v* *Properties*, p 519 —2 With *ammonia* to form NH_4NO_3 —3 With *dry sulphur dioxide* to form $\text{SO}_2\text{NO}_2\text{OH}$ (*v* NITROGEN DERIVATIVES OF SULPHUR ACIDS in vol IV) —4 With some *metals* to form acid salts, according to Ditté (*A Ch* [5] 18, 320), the nitrates which combine with HNO_3 when dissolved in excess of the acid are those of NH_4 , K , Rb , and Tl —5 With *nitric anhydride* to form $(\text{HNO}_3)_2\text{N}_2\text{O}_5$ ($=\text{H}_2\text{N}_2\text{O}_{11}$, *v* *Dinitric acid*, *infra*)

Detection and Estimation *v* *Manuals of Analysis*

DINITRIC ACID $\text{H}_2\text{N}_2\text{O}_{11}$ ($=(\text{HNO}_3)_2\text{N}_2\text{O}_5$) Weber (*J pr* [2] 6, 342) obtained this compound by adding very conc colourless HNO_3 to melted N_2O_5 , and cooling to c 8° (for details of preparation *v* Weber, *l c*) A bluish yellow liquid, so solidifying at c 5°, S G 1.642 at 18° Fumes in air, dissolves in water with production of much heat, when gently warmed N_2O_5 is evolved. It is very dangerous to keep the compound in sealed tubes as explosions generally occur. Acts as a powerful oxidiser, forms NO_2 derivatives with many carbon compounds. The relation of this acid to nitric acid is probably similar to that of disulphonic to sulphonic acid $\text{H}_2\text{SO}_4 + \text{SO}_3 = \text{H}_2\text{S}_2\text{O}_7$, $2\text{HNO}_3 + \text{N}_2\text{O}_5 = \text{H}_2\text{N}_2\text{O}_{11}$, M M P M

NITRIDES *Compounds of N with one other more positive element* The term is generally applied to the binary compounds of N with B , P , Si , and the metals. The metallic nitrides have for the most part the composition denoted by the formulae R_3N , or R_2N , R = monovalent metal. They are mostly obtained by the action of ammonia on metallic oxides or chlorides; frequently also by the direct action of atmospheric

nitrogen on metals at the moment of separation from their oxides by charcoal. Some metallic nitrides are also obtained by the reaction of metals or oxides of metals with hydrazoic acid, HN_3 (*q v p 559*). Most of them have a metallic aspect, are easily decomposed by heat, sometimes

that nitrates are formed in a fertile soil from NH_3 salts applied to the soil. The following table, taken from the article NITRIFICATION in the first edition of this Dictionary (Supp. in p 1399), exhibits very clearly the oxidation of NH_3 salts to nitrates in the soil —

Nitrogen existing as nitrates in one million parts of drainage water.

Dates on which drainage water was collected	Plots which received no NH_3 salts				Plots which received 400lbs NH_3 salts per acre, between March 10 and 13						
	3 and 4	5	16	17	7	10	11	12	13	14	18
February 16, 1879	34	37	42	33	35	52	48	41	38	48	42
April 7, 1879	36	29	36	23	39 0	45 4	41 8	25 4	29 4	33 0	40 4

with explosion, some of them, however, with stand a very high temperature without decomposition. Many of them are reduced to metals by strongly heating in hydrogen or ammonia gas, heated with water or hydrate of potassium they often yield metallic oxides and ammonia, they burn when heated in the air. They unite with other metallic compounds, especially with metalamides. The chief metallic nitrides are those of Al, Cu, Cr, Fe, Mg, Hg, Mo, Ni, Nb, Pt, K, Na, Ta, Th, ?Sn , Ti, W, U, Y, Zn, and Zr. For descriptions of the nitrides *v* the several metals enumerated above. M M P M

NITRIFICATION Nitre occurs in many parts of the earth. In some places it is found as an efflorescence on the surface of the soil, it is formed on the sides of caves (generally those frequented by animals), on the walls of old stables, and in the soil surrounding some of the villages in North India (*v* Palmer, *C J* [2] 6, 318). Nitre also occurs in Algeria, Spain, &c., and very large deposits of NaNO_3 are found in Peru. Nitrates are found in all, or almost all, soils, especially in those which are fertile and well manured. Nitrates are obtained by the artificial oxidation of nitrogenous matters in soils, by arranging the conditions so that there is first an accumulation of nitrogenous compounds, and then a slow oxidation in presence of moisture and some base which combines with the HNO_3 , as it is produced (*v* Potassium nitrate, Formation, p 515).

That HNO_3 is formed by the oxidation of nitrogen compounds has been abundantly proved (*eg* by Boussingault, *C R* 32, 477). There is no convincing experimental proof that the N of the air is oxidised to HNO_3 , under normal naturally occurring conditions. L T Wright (*C J* 35, 42) found that HNO_3 is formed by burning H in air, but that if special precautions are taken to remove all NH_3 from the air, HNO_3 is not formed. Boussingault (*C R* 76, 22) found that there was no increase in the total N in a quantity of soil rich in N compounds, after keeping in a closed vessel full of air for 11 years, although there was a marked increase in the nitrates in the soil. Various experimenters have proved that NH_3 can be oxidised to HNO_3 , *eg* by ozone, by KMnO_4 and H_2SO_4 , &c. (*v* Nitric acid, Formation, p 518). Experiments conducted at Rothamsted have proved conclusively

In 1862, Pasteur suggested that the oxidation of nitrogen compounds in the soil to nitrates is accomplished by a living organism. In 1877 Schloesing a Muntz (*C R* 84, 301, 85, 1018) showed that nitrification did not take place in sewage slowly filtered through a column of pure sand and limestone until after 20 days, that nitrification then proceeded rapidly, but was completely stopped by allowing vapour of chloroform free access to the surface of the sand and limestone, 7 weeks after the application of CHCl_3 , a little vegetable earth, which was known to nitrify easily, was placed on the surface of the sand, and then nitrification began again. S a M also showed that heating soil to 100° destroyed the power of that soil to produce nitrates from nitrogenous matter.

Warrington has conducted, at Rothamsted, an extensive, laborious, and accurate, series of experiments on nitrification in soils, his memoirs are to be found in *C J* 33, 44, 35, 429, 45, 637 [numerous references here to other workers], 47, 758, 51, 118, 53, 751, *cf* also Munro, *C J* 49, 632. The results fully confirm the view that the production of nitrates in soils from nitrogenous compounds is due to the activity of a living organism. Summarising the results, Warrington says (*C J* 45, 461) 'Nitrification in soils and waters is found to be strictly limited to the range of temperature to which the vital activity of low organisms is confined. Nitrification is also dependent on the presence of plant-food suitable to organisms of low character. Further proof is afforded by the fact that antiseptics are fatal to nitrification. The action of heat is equally confirmatory. By raising sewage or soil to the temperature of boiling water, its nitrifying power is entirely destroyed. Finally, nitrification can be started in boiled sewage, or in other sterilised liquids of suitable composition, by the addition of a few particles of surface soil, or a few drops of a solution which has lately nitrified, while, if no such addition is made, these liquids may be freely exposed to filtered air without nitrification taking place. Schloesing has apparently given a final blow to the theory that air exists in a condensed state in the pores of a soil, and may on that account exert special oxidising powers, by his recent experiments (*Ann de la Science agronomique*, 1884 1), showing that the gases in a vegetable soil simply occupy the normal

volume due to temperature and atmospheric pressure.

The result of Warrington's examination of the classes of bodies that undergo nitrification in soils is, that all nitrogenous substances which yield ammonia by the action of organisms existing in fertile soils are nitrifiable (*C J* 45, 658).

The conditions under which nitrification proceeds in soils, sewage, &c., are as follows (*W. C J* 45, 654 *et seq.*)—(1) A sufficient quantity of some base must be present to combine with the nitric acid. When the whole of the bases present are neutralised the nitrification stops, although nitrifiable compounds may still be present. There is a limit to the quantity of alkaline carbonate which can be present in a liquid undergoing nitrification. This fact renders impossible the nitrification of urine, except when diluted with water, because the first product of the action on urine of organisms in the soil is NH_3 , carbonate, and unless this be present in very moderate quantity nitrification cannot proceed. Warrington's experiments showed that 14 parts urine in 100 water did not nitrify when kept in contact with soil containing nitrifying organisms for 172 days. Addition of gypsum allows nitrification to proceed in cases where it would either not occur or be stopped by the presence of NH_3 , carbonate (*W. C J* 47, 758, Pickard, *Ann. de Science agronomique*, 1884 302, Joule, *ibid* 1884 289). Thus, whereas a 14 p.c. urine solution did not nitrify after 172 days, a 80 p.c. solution, to which gypsum was added, nitrified after 78 days, and a 50 p.c. solution containing gypsum nitrified after 151 days. (2) In order that a nitrogenous liquid shall nitrify, the concentration of the liquid must not exceed a certain degree (for experiments *v C J* 45, 662). (3) A sufficient quantity of the nitrifying organism must be present. The vigour of the organisms obtained in different cultivations differs considerably. A liquid which refuses to nitrify when seeded, because its concentration is excessive, may be caused to undergo nitrification by seeding it with a very vigorous organism, &c. one which has been obtained by cultivation in a fairly conc. solution well supplied with nutriment. 'The plan that has proved most effective in producing rapid nitrification is to make use of the sediment lying at the bottom of the vessel in which the nitrification of a fairly strong solution has been conducted' (*W. C J* p. 665). (4) Stagnant liquids of considerable depth nitrify much less quickly than a shallow liquid of similar composition, or than a liquid which is poured over a porous solid and so brought into contact with fresh supplies of oxygen. (5) Some quantity of carbon in combination—'organic carbon'—is needed for the nourishment of the nitrifying organism, but no advantage accrues from the presence of more C than is required for this purpose (*cf* Munro, *C J* 49, 661 *et seq.*). (6) Nitrification proceeds within certain limits of temperature, the organism seems to be destroyed at, or somewhat below, 100° , at 8° – 5° nitrification proceeds much more slowly than at somewhat higher temperatures. (7) Light tends to decrease, or even stop, the process of nitrification (*v W. C J* 88, 44). The product of nitrification is sometimes a nitrite, sometimes a nitrate, and sometimes both. The exact con-

ditions under which nitrite and nitrate are formed have not yet been determined satisfactorily. In a later communication (*C J* 59, 484) Warrington describes the isolation of two organisms one of these oxidises ammonia to nitrous acid and has no effect on nitrites, the other produces neither nitrites nor nitrates in ammoniacal solutions, but in absence of ammonia rapidly converts nitrites into nitrates. P. F. Frankland and G. O. Frankland (*Pr* 47, 296) seem to have isolated a bacillus, which grows slowly in broth, and which converts ammoniacal salts into nitrates.

With regard to the distribution of the nitrifying organisms in the soil, Warrington's experiments show that the organisms are not evenly distributed below a depth of about 9 inches in clay-soils (*C J* 45, 649), and that nitrification occurs chiefly, if not altogether, in the surface soil, and rarely in a clay-subsoil 2 or 3 feet from the surface (*C J* 51, 118). M. M. P. M.

NITRILES. Compounds of the form RCN where R is an organic radicle.

Formation—1 By distilling potassium alkyl sulphates (KRSO_3) with potassium cyanide (Dumas, *C R* 25, 474).—2 From alkyl iodides and potassium cyanide in presence of dilute alcohol (Schlagdenhauffen, *C R* 48, 228, Henry, *C R* 104, 1181).—3 By dehydrating amides by distillation with P_2O_5 or P_2S_5 (Dumas, Malaguti, & Leblanc, *A* 64, 333).—4 By distilling organic acids with potassium or lead sulphocyanide. This reaction is most successful with aromatic acids (Krusch, *B* 17, 1766).—5 By boiling the formyl derivatives of aromatic amines (*eg* formanilide) with zinc dust (Gasiorowski, *B* 17, 73).—6 By the action of bromine and NaOH on the amide of the acid containing one C atom more $\text{XCH}_2\text{CONH}_2 + 3\text{Br}_2 + 8\text{NaOH} = \text{XCN} + 6\text{NaBr} + \text{Na}_2\text{CO}_3 + 6\text{H}_2\text{O}$. This reaction, which gives a means of descending the series, is particularly applicable to the higher homologues (yield from monoamide = 30 p.c.), but the lower the homologue the smaller is the yield, till in the case of valeramide hardly any nitrile at all is formed, the chief product of the reaction being butylamine (Hofmann, *B* 17, 1406).—7 By distilling the formyl derivatives of aromatic monamines with zinc dust, the yield being 10 p.c.–20 p.c. (Gasiorowski & Merz, *B* 18, 1008).—8 By warming the oxim of the corresponding aldehyde with Ac_2O .—9 Aromatic nitriles may be formed from diazo-compounds by Sandmeyer's reaction, using cuprous cyanide (Sandmeyer, *B* 17, 2653).

Reactions—1 Readily converted into NH_3 , and the corresponding acid by heating with acids or alkalis. Cold conc. HCl aq. forms the intermediate amides.—2 Alcohol (1 mol.) and gaseous HCl form the salt of an imido-ether, *eg* $\text{CH}_3\text{CH}(\text{OEt})\text{NH}_2\text{Cl}$.—3 Zinc and dilute acids yield the corresponding amine (Mendius, *A* 121, 129).—4 H_2S unites, forming a thioamide, *eg* CH_3CSNH_2 .—5 Sodium polymerises many fatty nitriles. Sodium acting on an alcoholic solution of aromatic nitriles often reduces them to the aromatic hydrocarbons or their hydrides (Bamberger & Lotter, *B* 20, 1702).—6 HBr combines with nitriles. The product is decomposed by water, fatty nitriles yielding the acid, while benzotrile yields benzamide—

7 *Hydroxylamins unites, forming amidoxims*
 $R\text{C}(\text{NH}_2)\text{NOH}$

Isonitriles v. CARBAMINES

NITRILIO DIACETONAMINE v. ACETON-AMINE

NITRILIO-PROPIONITRILE A name for hydrocyanaldimine *v* vol 1 p 104

NITRILES *Salts of nitrous acid, HNO₂, v. Nitrous acid and Nitrites, under Nitrogen, p 567*

NITRO- *Use of this prefix applied to inorganic compounds for nitro-compounds and nitro salts v the element the nitro compound of which is sought for, or the salts to the names of which nitro is prefixed. Thus, nitroferrocyanides are described under ferrocyanides, a section of the group CYANIDES, nitrochromate of potassium is described under CHROMATES*

NITRO-ACENAPHTHENE $\text{C}_{12}\text{H}_7\text{NO}_2$, [155°] (J), [102°] (A) Formed by nitrating acenaphthene dissolved in HOAc (Jandrier, C R 104, 1858, Quincke, B 20, 609, 21, 1454) Pale-yellow needles (by sublimation) Easily reduced by zinc and HCl to an amido derivative, which yields a bluish-violet product on oxidation

Dinitro-acenaphthene $\text{C}_{12}\text{H}_5(\text{NO}_2)_2$, [206°] Yellow needles, formed at the same time as the preceding (Q, cf Berthelot, B 1, 8, 250)

NITRO ACETIC ACID $\text{CH}_3(\text{NO}_2)\text{CO}_2\text{H}$ or $\text{CH}_3(\text{ONO})\text{CO}_2\text{H}$

Ethyl ether EtA' (c 155°) SG ± 1.133 Formed from bromo or iodo acetic ether and silver nitrate at 130° Extracted with ether Forrand, B 1 [2] 31, 536, Steiner, B 15, 1605, Lewkowsch, J pr [2] 20, 163 Oil, smelling like nitrous ether Yields amido acetic ether on reduction Splits up when boiled for a long time, yielding CO_2 , oxalic ether, and NO

NITROACETAMIDO-DI METHYL-HYDRO-QUINONE v Acetyl-di methyl derivative of Nitro amido-hydroquinone

NITRO ACET ANILINE v Acetyl derivative of Nitro ANILINE

NITRO-ACET-NAPHTHALIDE v Acetyl derivative of Nitro NAPHTHYLAMINE

NITRO ACETONITRILE A name formerly given to fulminic acid (*v* vol 11 p 817)

NITRO ACETONYL-UREA

$\text{CMe}_2\text{CO}\text{NH}\text{N}(\text{NO}_2)\text{CO}$ (?) [141°] Obtained by nitrating acetonyl urea (Franchimont & Klobbie, R T C 7, 241) Slender needles, sl sol water and benzene, sol alcohol and ether Decomposed by boiling water, forming a oxy-isobutyric acid

NITRO-ACETOPHENONE $\text{C}_6\text{H}_5\text{NO}_2$, *v* $\text{CH}_3\text{CO}\text{C}_6\text{H}_4\text{NO}_2$ [12] Nitro phenyl methyl ketone Formed, together with the *m*-isomeride, by nitration of acetophenone, especially at 40° (Engler, B 18, 2238) Prepared by boiling *o*-nitro-benzoyl-acetoacetic ether with dilute H_2SO_4 for 8 hours, and extracting with ether (Guthzeit, A 221, 325) Oil, v sol alcohol With PCl_5 it yields chloro-*o*-nitro-styrene Boiling alcoholic ammonium sulphide converts it into indigo KMnO_4 yields *o*-nitro benzoic acid Tin and HCl yield *o*-amido-acetophenone (242°-252°).

***m* Nitro-aceto-phenone**
 $\text{CH}_3\text{CO}\text{C}_6\text{H}_4\text{NO}_2$ [18] [81°] (Buchka, B 10, 1714) Formed by the action of *m* nitro-benzoyl chloride on sodio acetacetic ether and digesting the product with water (Gevekoht, B 15, 2084)

Obtained also by nitrating acetophenone in the cold Needles, volatile with steam Yields *m*-nitro-benzoic acid on oxidation

Oxim [132°] Forms a methyl-ether [84°] (Gabriel, B 15, 5063)

***p*-Nitro-acetophenone** $\text{CH}_3\text{CO}\text{C}_6\text{H}_4\text{NO}_2$, [144] [81°] Formed by the action of *p* nitro-benzoyl chloride on sodio acetacetic ether, and digesting the product with water (Gevekoht, B 15, 2084) Formed also by warming *p*-nitro phenyl-propionic acid with dilute H_2SO_4 , or by allowing its ether to stand with conc H_2SO_4 at 40°, pouring the *p* nitro-benzoyl acetic acid which is formed into water, and expelling CO_2 by boiling (Drewson, A 212, 160, Engler & Zulke, B 22, 203) Yellow prisms Yields chloro-*p*-nitro-styrene with PCl_5 Yields *p*-amido acetophenone [106°] on reduction by tin and HCl

Phenyl-hydrazide

NITRO-ACET TOLUIDE v Acetyl derivative of Nitro TOLUIDINE

NITRO ACET-XYLIDE v Acetyl derivative of Nitro XYLIDINE

DI-NITRO-ACETYLENE-DI-UREA $\text{C}_2\text{N}_4\text{H}_2\text{O}_4$, *v* $\text{CO}\text{C}(\text{NH}\text{C}(\text{NO}_2)\text{NH})\text{CO}$ *Di-nitroglycoluril* Formed by nitrating acetylene-di-urea (Franchimont & Klobbie, R T C 7, 18) Charas at 180° On boiling with water it is decomposed into CO_2 , water, and an isomeride of hydantonic acid.

NITRO-ACIDS v Nitro-compounds

(a) **NITRO-ACRIDINE** $\text{C}_{12}\text{H}_7\text{N}_3\text{O}_2$, *v* $\text{C}_{12}\text{H}_7(\text{NO}_2)_2\text{N}$ [214°] Formed, together with the two following bodies, by nitrating acridine (Graebe & Caro, A 158, 275) Golden-yellow plates (by sublimation), insol water, sl sol alcohol and ether Its solution in dilute acids shows blue fluorescence The hydrochloride crystallises in yellow prisms

(B) Nitro-acridine [154°] Plates, v sol. hot alcohol Forms salts with acids

Di-nitro acridine $\text{C}_{12}\text{H}_5(\text{NO}_2)_2\text{N}$. Reddish-yellow tables, sl sol alcohol and ether Does not dissolve in dilute acids

TRI NITRO ACRIDINE CARBOXYLIC ACID
 $\text{C}_{12}\text{H}_5(\text{NO}_2)_3\text{NCO}_2\text{H}$ Formed by boiling methyl-acridine with HNO_3 (SG 133) (Bernthsen, A. 224, 40) Yellow prisms

NITRO ALDEHYDO BENZOIC ACID

$\text{C}_6\text{H}_4(\text{CHO})(\text{NO}_2)\text{CO}_2\text{H}$ [124] [160] Formed, together with a small quantity of $\text{C}_6\text{H}_3(\text{CHO})(\text{NO}_2)_2\text{CO}_2\text{H}$ [184°] by nitrating *p* aldehyde benzoic acid (Löw, A 231, 869) Four sided prisms (from water), v sol alcohol and ether With acetone and NaOH it forms indigo carboxylic acid—AgA' sl sol water

Ethyl ether EtA'

NITRO ALDEHYDO-CINNAMIC ACID

$\text{C}_6\text{H}_5\text{NO}_2$, *v* $\text{C}_6\text{H}_4(\text{CH}\text{CH}\text{CO}_2\text{H})(\text{NO}_2)(\text{CHO})$ [124] [194°]. Formed by nitration of aldehydo-cinnamic acid (Löw, A 231, 376) Prisms *v*. sol glacial acetic acid, acetone, and hot water, hardly sol. ether or chloroform Does not give the indigo reaction with acetone and NaOH, so that NO_2 is probably not *ortho* to CHO —AgA' aq.

Ethyl ether EtA' [80°] Prisms

NITRO ALIZABIN v Nitro-DI-oxi ANTHRA-QUINONE

DI NITRO ALLYL-ANILINE $\text{C}_8\text{H}_7\text{N}_2\text{O}_4$, *v* $\text{C}_8\text{H}_7\text{NHC}_6\text{H}_4(\text{NO}_2)_2$ [76°] Formed from bromo-

m di nitro-benzene, allylamine, and alcohol (Romburgh, *R T C* 4, 192) Yellow needles

Tri-nitro-allyl-aniline $C_9H_7N_3O_6$ *ze* $C_9H_7N_3O_6$ [80°] Formed by the action of allylamine on chloro tri nitro benzene (peryl chloride) (R)

p-NITRO-ALLYL-BENZOYL-ACETIC ETHER $C_{12}H_{11}(NO_2)COCH(C_2H_5)CO_2Et$ [46°] Formed (from sodium *p* nitro benzoyl acetate ether and allyl iodide (Perkin & Bellenot, *C J* 49, 452)

NITRO-AMIDO-ACETAMIDE $C_8H_8N_2O_4$ *ze* $CH_3(NHNO_2)CONH_2$ Formed by decomposing nitro hydantoin by boiling water (Franchimont & Klobbie, *R T C* 7, 239) Long prisms, decomposing at 130°

NITRO-DI-AMIDO-BENZENE *o* Nitro-phenylene diamine

NITRO-AMIDO-BENZENE SULPHONIC ACID $C_8H_7N_2SO_3$ *ze* $C_8H_7(NO_2)(NH_2)(SO_3H)$ [2 1 4] *o* Nitraniline sulphonic acid Formed by heating (1, 2, 4) bromo nitro benzene sulphonic acid with alcoholic NH_3 at 180° (Goslich, *A* 180, 103) and by nitrating acetyl *p* amido benzene sulphonic acid (Nietzki, *B* 18, 294, 21, 3220). The same acid appears to be formed by sulphonating *o*-nitro aniline (Post & Hardtung, *B* 13, 88) Yellow needles, extremely sol water, *m* sol alcohol and $HClAq$ Boiling aqueous KOH converts it into $C_8H_7(NO_2)(OK)(SO_3K)$ $HClAq$ at 180° yields *o*-nitro aniline

Salts — KA' *aq* *S* 5 at 6° — NH_4A' *S* 13 at 6° — BaA' , 2½ *aq* *S* 5 at 9° — CaA' , 2½ *aq* (*P a H*) — PbA' , 2½ *aq* *S* 2 at 6°

Chloride $C_8H_7(NO_2)(NH_2)SO_2Cl$ [60°] **Amide** [156°] Yellow needles

Nitro-amido-benzene sulphonic acid $C_8H_7(NO_2)(NH_2)(SO_3H)$ [1 2 4] Formed by the action of HNO_3 (1 mol) on acetyl amido benzene *m*-sulphonic acid (1 mol) dissolved in conc H_2SO_4 (Eger, *B* 21, 2579, 22, 847) Yellow needles (from water), *sl* sol alcohol, almost *insol* ether — KA' golden yellow plates (from water) — NaA' yellow needles, *v* sol water

Nitro-amido-benzene sulphonic acid $C_8H_7(NO_2)(NH_2)(SO_3H)$ [3 1 6] Formed by heating *m*-nitroaniline sulphate at 120°–170° with $ClSO_3H$ (Lumprecht, *B* 18, 2186) Long colourless prisms or glistening plates Easily soluble in hot water, more sparingly in cold

Salts — $A'K$ *aq* long thin glistening red prisms or yellowish-red plates — $A'Na$ 2½ small yellow plates — $A'Ca$ 4½ easily soluble large orange tables or thick red prisms — $A'Ba$ *aq* red tablets or prisms, sparingly soluble in cold water

Nitro-amido-benzene sulphonic acid $C_8H_7(NO_2)(NH_2)(SO_3H)$ [3 1 4] ? Formed by heating *m* nitro aniline with fuming H_2SO_4 at 160° (Post & Hardtung, *A* 205, 102, *B* 13, 40) Yellowish-brown prisms — BaA' 2½ long spikes *S* 14 at 100° — CaA' , 4½ small needles, *v* sol water This acid is perhaps identical with the preceding

Nitro-amido-benzene sulphonic acid $C_8H_7(NO_2)(NH_2)(SO_3H)$ [4 1 8] Formed in small quantity by heating $C_8H_7Br(NO_2)(SO_3H)$ [1 4 8] with alcoholic ammonia at 160° (Thomas, *A* 186, 132) Needles, *v* sol water — BaA' , 1½ *aq* *S* 15 at 15°

Nitro-amido-benzene disulphonic acid $C_8H_7(NO_2)(NH_2)(SO_3H)_2$ Formed by the action

of ammonium sulphide on di nitro benzene di-sulphonic acid obtained from nitro benzene *m* sulphonic acid (Lumprecht, *B* 8, 289) Very deliquescent mass — BaA' 2½ *aq*

(a) **NITRO-AMIDO-BENZOIC ACID** $C_8H_7N_2O_4$ *ze* $C_8H_7(NO_2)(NH_2)CO_2H$ [2 5 1]. Mol *w* 182 Formed by boiling (a) di nitro-*m*-uramidic benzoic acid with water (Griess, *B* 5, 198, 11, 1734) Yellow needles or prisms, *m* sol hot water, *v* sol hot alcohol Yields nitro oxy benzoic acid [169°] Yields on reduction a diamido benzoic acid which forms *p* phenylene-diamine on distillation — BaA' , 3½ *aq*

(b) **Nitro-amido-benzoic acid** $C_8H_7(NO_2)(NH_2)CO_2H$ [4 3 1] [298°] Formed by boiling (b) di nitro *m* uramidic benzoic acid with water (Griess) Formed also by saponifying its acetyl derivative (Kaiser, *B* 18, 2946) Red plates or needles Yields on reduction a diamido benzoic acid which forms *o* phenylene-diamine on distillation — CaA' , *aq* red crystals, *sl* sol water — BaA' , 2½ *aq*

Ethylether EtA' [139°] Red needles **Formyl derivative** [221°] (Zehra, *B* 23, 3634)

Acetyl derivative

$C_8H_7(NO_2)(NHAc)CO_2H$ [206°] Formed, together with the (2, 3, 1) isomeride, by nitrating *m* acetamido benzoic acid below 0° (K) Yellow tables — CaA' , 7½ *aq* — BaA' , 7½ *aq*

γ **Nitro-amido-benzoic acid**

$C_8H_7(NO_2)(NH_2)(CO_2H)$ [2 3 1] [157°] Formed by boiling (γ) di nitro *m* uramidic benzoic acid with water (Griess, *B* 2, 435, 5, 199) Formed also by saponifying its acetyl derivative which is prepared as above (K) Long yellow needles, *v* sol hot water Yields *o* nitro benzoic acid on elimination of NH_3 (Griess, *B* 11, 1734) Yields on reduction a di amido benzoic acid which forms *o* phenylene diamine on distillation — KA' 2½ *aq* — BaA' , 7½ *aq* — $HA'HCl$, white crystals, decomposed by water

Acetyl derivative

$C_8H_7(NO_2)(NHAc)CO_2H$ [241°] Colourless crystals — CaA' , 6½ *aq* — BaA' , 2½ *aq*

(δ) **Nitro amido benzoic acid**

$C_8H_7(NO_2)(NH_2)CO_2H$ [3 4 1] [284°] Formed by heating di-nitro-*p* uramidic benzoic acid with water (Griess, *B* 5, 855) Formed also by heating $C_8H_7(NO_2)(OMe)CO_2H$ [3 4 1] with aqueous ammonia at 140°–170° (H Salkowski, *A* 173, 52) Reddish yellow needles (from alcohol), *sl* sol hot water Yields, on reduction, di amido benzoic acid [210°] Converted into *m* nitro benzoic acid by the diazo reaction — KA' *aq* orange prisms — BaA' , 5½ *aq*

Ethyl ether EtA' [145°] Formed by heating $C_8H_7Br(NO_2)CO_2Et$ [4 3 1] with alcoholic NH_3 for 3 hours at 150° (Grohmann, *B* 23, 3449) Yellow crystals

Amide $C_8H_7(NO_2)(NH_2)CONH_2$ [227°] Formed by heating $C_8H_7Br(NO_2)CONH$ [4 3 1] with alcoholic NH_3 at 180° (G) Lemon yellow crystals, *insol* water, *sl* sol alcohol

Acetyl derivative

$C_8H_7(NO_2)(NHAc)CO_2H$ [221°] Formed by nitration of acetyl *p* amido benzoic acid below 10° (Kaiser, *B* 18, 2948) Thick yellow tables, *v* *sl* sol cold water — CaA' , 2½ *aq* — BaA' , 6½ *aq*

(e) **Nitro-amido-benzoic acid**

$C_8H_7(NO_2)(NH_2)(CO_2H)$ [5 2.1] [263°].

Formation — 1 By boiling di nitro-*o* uramido benzoic acid with water (Griess, *B* 11, 1730) — 2 By heating $C_6H_3(NO_2)_2(OEt)CO_2Et$ with alcoholic NH_3 at 140° and boiling the resulting amide with baryta water (Hubner, *A* 195, 21) — 3 By heating $C_6H_3Br(NO_2)CO_2H$ [2 5 1] with conc NH_3 aq at 145° (Rahlis, *A* 198, 112) — 4 From its amide which is formed when nitro isatoic acid is warmed with aqueous ammonia (Kolbe, *J pr* [2] 30, 477)

Properties — Slender yellow needles, v sol boiling water. On elimination of NH_3 , it yields *m* nitro benzoic acid

Salts — $KA'2aq$ — $CaA'3aq$ — $BaA'3aq$ v sol cold water, sl sol hot water — $PbA'2aq$ — $HA'HCl$ needles, decomposed by water

Amide $C_6H_3(NO_2)_2(NH_2)CONH_2$ [200°–210°] Yellow needles (from acetone)

(ζ) Nitro-amido-benzoic acid
 $C_6H_3(NO_2)(NH_2)(CO_2H)$ [3 2 1] [204°] Formed by heating $C_6H_3(NO_2)(OEt)(CO_2Et)$ [3 2 1] with alcoholic NH_3 at 130° – 160° and saponifying the resulting amide (Hubner, *A* 195, 37) Yellow needles (from water). On elimination of NH_3 , it yields *m* nitro benzoic acid — $KA'2aq$ — $CaA'2aq$ — $SrA'2aq$ — $BaA'2aq$ purple needles, sl sol cold water — $Pb(OH)A'$ — $CuA'2$ — AgA' — $HA'HCl$ needles

Ethyl ether EtA' [204°] Plates
Amide $C_6H_3(NO_2)(NH_2)(CONH_2)$ [109°] Yellow plates, almost insol water and alcohol

(η) Nitro-amido-benzoic acid
 $C_6H_3(NO_2)(NH_2)CO_2H$ [5 3 1] [208°] Formed by reducing *s* di nitro benzoic acid with NH_3 and H_2S (Hubner, *A* 222, 81) Small golden prisms (from water). On elimination of NH_3 , it yields *m* nitro benzoic acid. Reduces to di amido benzoic acid which, when distilled with steam, forms phenylene *m* diamine — NaA' aq red needles — $NH_4A'3aq$ bright yellow needles — $BaA'4aq$ — $CaA'5\frac{1}{2}aq$ — $PbA'3\frac{1}{2}aq$ — AgA' aq

Ethyl ether EtA' [155°] Yellow needles
Di-nitro-*o*-amido-benzoic acid $C_6H_3N_2O_5$ *ie* $C_6H_3(NO_2)_2(NH_2)CO_2H$ Mol w 227 [256°] Formed, together with its methyl ether, by heating $C_6H_3(NO_2)_2(OEt)CO_2Me$ with aqueous ammonia (H Salkowski, *B* 4, 870, *A* 173, 40) Golden yellow scales (from alcohol) — NH_4A' aq

Methyl ether MeA' [165°] Needles

Ethyl ether EtA' [135°] Laminæ

Di nitro *p* amido benzoic acid
 $C_6H_3(NO_2)_2(NH_2)(CO_2H)$ [5 3 4 1] **Chrysamic acid** [260°] Formed by the action of aqueous ammonia upon $C_6H_3(NO_2)_2(OMe)CO_2H$, which is a product of the action of warm fuming HNO_3 on anisic acid (Cahours, *A Ch* [3] 27, 454, Beilstein *A Kellner*, *A* 128, 104) Formed also by oxidation of di nitro *p* toluidine by chromic acid mixture (Friedrich, *B* 11, 1976) Plates (from alcohol), sl sol cold Aq — NH_4A' — AgA'

Methyl ether MeA' [144°] Plates

Ethyl ether EtA' [114°] Plates

Acetyl derivative

$C_6H_3(NO_2)_2(NHAc)CO_2H$ [270°] Silky needles

Nitro-di-amido-benzoic acid $C_6H_3N_2O_5$ *ie* $C_6H_3(NO_2)(NH_2)_2CO_2H$ [5 4 3 1] Formed by reducing chrysamic acid with H_2S and alcoholic NH_3 (*B* *a K*) Minute red crystals, v sol alcohol, sl sol hot water — NH_4A' aq Monoclinic prisms, *a b c* = 1.073 : 1 : 809, β = $77^\circ 32'$

DI - NITRO - AMIDO - BENZYL - METHYL KETONE $C_6H_3N_2O_5$ [214°] Formed by reducing tri nitro benzyl methyl ketone in alcoholic solution with the theoretical quantity of $SnCl_4$ and HCl (Ditttrich, *B* 23, 2724) Groups of golden yellow needles (from alcohol)

NITRO AMIDO-ISOBUTYL-BENZENE
 $C_6H_3C_4H_7(NO_2)(NH_2)$ [1 2 3] Nitro isobutyl aniline [124°] Formed by saponifying its acetyl derivative (Gelzer, *B* 21, 2941) Yellow crystals, v sol boiling water

Acetyl derivative
 $C_6H_3C_4H_7(NO_2)(NHAc)$ [106°] Obtained by nitrating $C_6H_3C_4H_7NHAc$ Yellow needles, v sol sol boiling water

Nitro amido isobutyl benzene
 $C_6H_3C_4H_7(NO_2)(NH_2)$ [1 3 4] [106 5°] Formed from its acetyl derivative and alcoholic potash (Gelzer, *B* 20, 3254) Orange crystals, sl sol hot water

Acetyl derivative
 $C_6H_3C_4H_7(NO_2)(NHAc)$ [105°] (252°) Yellow needles. Obtained from $C_6H_3C_4H_7NHAc$ [1 4] and fuming HNO_3 at 0°

Di nitro amido isobutyl benzene
 $C_6H_3(C_4H_7)(NO_2)_2(NH_2)$ [127°] Got by heating di nitro isobutyl phenol [93°] with NH_3 aq at 175° (Barr, *B* 21, 1544) Yellow needles

p NITRO DI AMIDO - DI-ISOBUTYL - TRI PHENYL METHANE

$C_6H_3(NO_2)CH(C_6H_5)(NH_2)C_6H_5$ [126°] Formed from *p* nitro benzoic aldehyde, *p* amido isobutyl benzene, and conc H_2SO_4 (Bischler, *B* 21, 3207) Yellow needles. Its hydrochloride and platinochloride are both crystalline. Its di acetyl derivative melts at 114° , and its di benzoyl derivative at 126°

m Nitro di *p* amido di isobutyl tri phenyl methane $C_6H_3N_2O_5$ [65°] Formed from *m* nitro benzoic aldehyde, *p* amido isobutyl benzene, and conc H_2SO_4

Di benzoyl derivative [114°] Plates

(α) NITRO-*o*-AMIDO CINNAMIC ACID

$C_6H_3N_2O_5$ *ie* $C_6H_3(NH_2)(NO)CHCHCO_2H$ [240°] Formed from *o* amido cinnamic acid (1 pt), conc H_2SO_4 (15 pts), and KNO_3 (3 pts) at 0° . On dilution with water (β) nitro *o* amido cinnamic acid separates as brownish needles, while the (α) compound may be ppd by nearly neutralising the mother liquor with $NaOH$ (Friedlander *a Lazarus*, *A* 229, 241) Brown needles. Insol benzene, ether, or petroleum, sl sol water, v sol alcohol and acetone. Dissolved by mineral acids, but ppd by sodic acetate. With HCl at 150° it gives (α)-nitro carbostyryl (nitro oxy quinoline), which forms slender white needles, v sl sol alcohol, glacial acetic acid, or acetone, and does not melt below 320°

Ethyl ether EtA' [160°] Formed, along with (β) nitro-carbostyryl, by nitrating *o*-amido cinnamic ether. Compact brown needles

(β) Nitro-*o* amido cinnamic acid

$C_6H_3(NH_2)(NO_2)CHCHCO_2H$ [254°] Prepared as above. Brownish yellow needles. Insol dilute mineral acids, and in water. Sol alkalis. With HCl at 150° it gives (β)-nitro carbostyryl [260°], which crystallises from glacial acetic acid in compact yellow needles

(3 4 1)-Nitro amido cinnamic acid

$C_6H_3(NO_2)(NH_2)C_4H_7CO_2H$ [225°] Formed by saponifying its acetyl derivative, which is

formed by nitration of acetyl-*p*-amido cinnamic acid (Gabriel & Herzberg, *B* 16, 2021) Red needles Sol hot alcohol and acetic acid, less sol water, nearly insol. benzene and ligroin

Acetyl derivative [261°-266°]

NITRO-AMIDO-*o*-CRESOL

$C_6H_4Me(NO_2)(NH_2)(OH)$ [1 3 5 2] [118°] Obtained by nitrating $C_6H_4Me(CO_2H)(NHAc)(OH)$ [1 3 5 2] and saponifying the resulting acetyl derivative (Nietzki & Ruppert, *B* 28, 8478) Brownish red needles (from alcohol)

Di-acetyl derivative

$C_6H_4Me(NO_2)(NHAc)(OAc)$ [146°]

Nitro-amido-cresol Methyl ether

$C_6H_4Me(NO_2)(NH_2)(OMe)$ [1 2 4 5]. [182°]

Formed from its acetyl derivative. Needles On elimination of NH_2 and reduction it yields $C_6H_4Me(NH_2)(OMe)$ [111°] Reduction yields $C_6H_4Me(NH_2)_2OMe$ [166°]

Acetyl derivative

$C_6H_4Me(NO_2)(NHAc)(OMe)$ [156°] Formed from $C_6H_4Me(NHAc)(OMe)$ in $HOAc$ by treatment with HNO_3 (S G 1 48) (Limpach, *B* 22, 89) Needles

*Nitro-*o*-amido cresol Methyl ether of the acetyl derivative*

$C_6H_4(CH_3NHAc)(NO_2)(OMe)$ [1 3 4] *Methyl derivative of acetyl-nitro oxy benzylamine* [137°]

Formed by nitrating $C_6H_4(CH_3NHAc)(OMe)$ in the cold (Goldschmidt & Polonowska, *B* 20, 4410) Prisms, v sl sol hot water Yields nitro amide acid on oxidation

Di-nitro amido m cresol

$C_6H_4Me(NO_2)_2(NH_2)OH$ [151°] (L & D), [156°] (E & O) Formed by reducing tri-nitro-cresol $C_6H_4Me(NO_2)_3OH$ [1 2 4 6 8] with H_2S and alcoholic NH_3 (Kellner & Beilstein, *A* 128, 166, Liebermann & Dorp, *A* 163, 104, Emmerling & Oppenheim, *B* 9, 1094) Thin yellow needles (from hot water)

Di nitro amido-m-cresol

$C_6H_4Me(NO_2)_2(NH_2)(OH)$ [1 2 5 4 6 8] [160°] Formed by nitrating $C_6H_4Me(CO_2H)(NHAc)(OH)$ (Nietzki & Ruppert, *B* 23, 8479) Large red needles Perhaps identical with the preceding

Mono-acetyl derivative [225°]

Di-acetyl derivative [175°]

NITRO AMIDO ISO CYMENE

$C_6H_4(NO_2)(NH_2)PrMe$ [x 50r 6 3 1] Formed from its phthalyl derivative by heating with conc HCl for 24 hours at 180° (Kelbe & Warth, *A* 221, 176) Oil Volatile with steam

Benzoyl derivative

$C_6H_4(NO_2)(NHBz)PrMe$ [177°] Formed by nitration of the benzoyl derivative of amido-isocymene

Phthalyl derivative

$(C_6H_4(NO_2)PrMe)_2C_2O_2C_6H_4$ [167°] Formed by nitration of the phthalyl derivative of amido isocymene

Di-nitro-amido-cymene $C_6H_4MePr(NO_2)_2(NH_2)$ [1 4 2 6 8] [118°-115°] From the ethyl ether of di-nitro-thymol and alcoholic NH_3 at 180° (Mazzara, *G* 19, 160) Yellow tables (from dilute alcohol), sl sol hot water

NITRO-*p* AMIDO-ETHYL-BENZENE

$C_6H_4(C_2H_5)(NO_2)(NH_2)$ [1 8 4] [47°] Yellowish-red prisms Sol. alcohol, ether, benzene, chloroform, and CS_2 , more sparingly in ligroin The acetyl compound is obtained by careful nitration of acetyl-*p*-amido-ethyl-benzene.

Acetyl derivative

$C_6H_4(C_2H_5)(NO_2)(NHAc)$ [47°], long yellow silky needles, extremely soluble in alcohol, ether, &c, less easily in ligroin (Paucksoh, *B* 17, 769)

*Di-nitro-*p*-amido-ethyl-benzene*

$C_6H_4(C_2H_5)(NO_2)_2NH_2$ [1 3 5 4] [185°] Orange-yellow prisms Sol benzene and chloroform, less easily in alcohol and ether The acetyl derivative is obtained by nitration of acetyl *p* amido ethyl-benzene

Acetyl derivative

$C_6H_4(C_2H_5)(NO_2)_2NHAc$ [182°], needles

NITRO AMIDO-HYDROQUINONE

Acetyl di-methyl derivative

$C_6H_4(NO_2)(NHAc)(OMe)_2$ [164°], yellow needles Formed by nitration of acetyl-amido-*o*-methyl-hydroquinone (Baessler, *B* 17, 2121)

NITRO-AMIDO HYDROTOLUQUINONE

$C_6HMe(NO_2)(NH_2)(OH)$ Formed by reducing di nitro hydrotolugunone with $SnCl_2$ (Kehrmann & Brasch, *J pr* [2] 39, 389) Its hydrochloride crystallises in long yellow needles changing to small plates

NITRO AMIDO MESITYLENE

$C_6HMe_2(NO_2)(NH_2)$ [73°] *Nitro-mesidine*

Formed by reducing di-nitro mesitylene with alcoholic ammonium sulphide (Maule, *C J* 2, 116, *A* 71, 137, Knecht, *A* 215, 98, Klobbie, *R T C* 6, 81) $-B^+HCl - B^+H.PtCl_6 - B^+H_3PO_4$

Acetyl derivative $C_6HMe_2(NO_2)(NHAc)$ [191°] Formed by nitrating acetyl mesidine (Biedermann & Ledoux, *B* 8, 58, Ladenburg, *B* 7, 1133, *A* 179, 163) Silky needles, m sol alcohol Feeble base

Benzoyl derivative $C_6HMe_2(NO_2)(NHBz)$ [168 5°] Formed by nitrating benzoyl mesidine (Schack, *B* 10, 1711)

Nitro-di amido mesitylene $C_6Me_2(NO_2)(NH_2)_2$ [184°] Formed, together with di nitro amido mesitylene by reduction of tri nitro mesitylene by alcoholic ammonium sulphide (Fittig, *A* 141, 139) Orange laminae (from water) or monoclinic crystals (from alcohol), $abc = 1.625 : 1.417$, $\beta = 60^\circ 4'$ (Lang, *A* 141, 140, Hintze, *A* 235, 183), v sl sol cold water $-B^+HCl_2$ tables

Di-nitro amido mesitylene $C_6Me_2(NO_2)_2(NH)$

Di nitro mesidine [194°] Formed as above Sulphur yellow crystals (from alcohol), insol water Very feeble base, its hydrochloride being decomposed by water

Acetyl derivative $C_6Me_2(NO_2)_2(NHAc)$ [275°] S (alcohol) 5 at 78° Formed by nitrating the acetyl derivative of nitro amido-mesitylene Needles

NITRO AMIDO - DI - METHYL ANILINE v

NITRO PHENYLENE DI-METHYL DIAMINE

TETRA NITRO DI-METHYL-DI AMIDO-DI-PHENYL-METHANE *Di-nitro- derivative*

$CH_3(C_6H_4(NO_2)_2NMeNO_2)_2$ Formed by the action of HNO_3 (S G 1 48) on $CH_3(C_6H_4NMe)_2$, dissolved in $HOAc$ (Bomburgh, *R T C* 7, 226)

Decomposes at 218° On boiling with aqueous KOH it gives off methylamine CrO_3 in $HOAc$ forms $CO(C_6H_4(NO_2)_2NMeH)_2$

NITRO-AMIDO-METHYL-QUINOLINE

$O_2H \cdot C(NH_2)C(NO_2) \cdot NMe$ [201°]. Formed by heating chloro-nitro-(*Py* 8)-methyl-quinoline with alcoholic NH_3 at 190° (Conrad & Limpach, *B* 21, 1965). Yellow needles (from alcohol).

NITRO AMIDO (a) NAPHTHOIC ACID

$C_{10}H_7N_2O_4$, i.e. $C_{10}H_7(NO_2)(NH_2)CO_2H$ [c 110°] Formed by reducing di nitro naphthoic acid [215°] with H_2S and NH_3 (Ekstrand, *J pr* [2] 38, 271, *B* 19, 1985) Needles (from water)

Nitro-amido-(β) naphthoic acid

$C_{10}H_7(NO_2)(NH_2)CO_2H$ [235°] Formed by reducing di nitro (β) naphthoic acid [226°] (Ekstrand, *J pr* [2] 42, 301) Stellate groups of small needles— HA/HCl needles

Nitro amido (α) naphthoic acid Acetyl derivative $C_{10}H_7(NO_2)(NHAc)CO_2H$ [259°] Formed by nitrating the acetyl derivative of (4', 1) amido naphthoic acid (Ekstrand, *J pr* [2] 38, 247) Yellow needles, v sol alcohol

Nitro amido-naphthoic acid

$C_{10}H_7(NO_2)(NH_2)CO_2H$ [4' 1' 1] *Anhydride* $C_{10}H_7(NO_2)<\frac{NH}{CO}>$ Nitro naphthostyryl [300°]

Formed by nitrating naphthostyryl (Ekstrand, *J pr* [2] 38, 180) Orange needles (from $HOAc$)

Di-nitro-amido-naphthoic acid Anhydride

$C_{10}H_4(NO_2)_2<\frac{NH}{CO}>$ [above 290°] Formed by nitrating the preceding anhydride (E) Plates, v sol $HOAc$ and alcohol

NITRO-AMIDO (α) NAPHTHOL $C_{10}H_7N_2O_4$

i.e. $C_{10}H_7(NH_2)(NO_2)OH$ [180°] Formed by reducing di nitro-(α) naphthol with ammonium sulphide (Ebell, *B* 8, 564) Small yellowish needles, insol water, sol alcohol

Benzoyl derivative $C_{10}H_7BzN_2O_4$ [158°] Small red needles (from alcohol) (Hubner, *A* 208, 332)

Nitro di-amido (α) naphthol Tri-acetyl derivative $C_{10}H_4(NO_2)(NHAc)_3OAc$ [235°] Formed by nitrating $C_{10}H_7(NHAc)_3OAc$ [280°] (Meerson, *B* 21, 1195) Yellow powder, yielding phthalic acid on oxidation by potassium permanganate Fuming $HClAq$ converts it into $C_{10}H_4(NO_2)(NH_2)<\frac{O}{N}>CMe$, whence boiling dilute

potash forms $C_{10}H_4(NO_2)(OH)<\frac{O}{N}>CMe$ crystallising in brown needles [163°]

NITRO-AMIDO-NAPHTHOL SULPHONIC ACID $C_{10}H_7N_2SO_4$, i.e. $C_{10}H_7(NO_2)(NH_2)(OH)SO_3H$ Formed by reducing di-nitro-naphthol sulphonic acid with $SnCl_2$ (Lauterbach, *B* 14, 2029) Golden yellow plates, m sol hot water Its alkaline solution is blood red

NITRO-AMIDO-TETRA OXY-BENZENE

$C_6(NO_2)_2(NH_2)_2(OH)_2$, Formed by partial reduction of di nitro di oxy quinone (nitraniline acid) with $SnCl$ Small violet needles, nearly insol alcohol, ether, and benzene (Nietzki & Benckiser, *B* 16, 2094, 18, 500) Its alkaline solution is readily oxidised by the air to nitro amido di-oxy quinone $C_6(NO_2)(NH_2)(OH)_2O_2$ Nitrous acid converts it into nitro-diazo di-oxy quinone $C_6(NO_2)(N.OH)(OH)_2O_2$

NITRO-α-AMIDO-PHENOL $C_6H_7N_2O_4$, i.e. $C_6H_7(NO_2)(NH_2)OH$ [6 2 1] [111°] Formed by reducing the corresponding di-nitro-phenol with ammonium sulphide (Post & Stuckenberg, *A* 205, 85) Red needles, sl sol water, v col alcohol and ether Gives a dirty-green colour with $FeCl_3$ — $B'H_2SO_4$ colourless pyramids

Nitro-α-amido-phenol. Methyl ether $C_6H_7(NO_2)(NH_2)(OMe)$ [3 2 1] [76°]. Formed

from $C_6H_7(NO_2)_2(OMe)$ [118°] by heating with alcoholic ammonia at 190° (Bantlin, *B* 11, 2106) Long yellow needles.

Nitro-α-amido-phenol $C_6H_7(NO_2)(NH_2)OH$ [4 2 1] [142°] Obtained by reducing the corresponding di-nitro-phenol with ammonium sulphide (Laurent & Gerhardt, *Compt Chim* 1849, 468, *A* 75, 68, Post & Stuckenberg, *A* 205, 71) Orange prisms (containing aq) melting at 80° to 90° or anhydrous crystalline crusts, melting at 142°, v sol hot water, alcohol, and ether— $C_6H_7KN_2O_4$, dark red nodules— $AgHA'$ brownish yellow pp (L & G)

Benzoyl derivative $C_6H_7BzN_2O_4$ [above 200°] Yellow needles, sl sol alcohol

Nitro-benzoyl derivative [218°]

Methyl ether $C_6H_7(NO_2)(NH_2)(OMe)$ Nitro anisidine Formed by reducing $C_6H_7(NO_2)_2(OMe)$ by alcoholic ammonium sulphide (Cahours, *A* 74, 301) Long garnet red needles, insol cold, sol boiling water— $B'HCl$ — $B'H_2PtCl_6$ — $B'HBBr$ — $B'HNO_3$ — $B'H_2SO_4$

Benzoyl derivative of the methyl ether $C_6H_7(NO_2)(NHBz)(OMe)$ Needles (from alcohol), m sol boiling alcohol

Ethyl ether $C_6H_7(NO_2)(NH_2)(OEt)$ [97°] Formed by heating the di-ethyl ether of di nitro di oxy s di phenyl hydrazine (hydrazonitro phenetole) with conc $HClAq$ (Andraee, *J pr* [2] 21, 818) Yellow needles, m sol water Yields p nitro phenetole on elimination of amidogen— $B'HCl$

Nitro-α-amido phenol

$C_6H_7(NO_2)(NH_2)(OH)$ [5 2 1] *Carbonyl derivative* $C_6H_7(NO_2)<\frac{NH}{O}>CO$ [241°] Obtained by nitrating carbonyl α amido phenol (Chelmicki, *J pr* [2] 42, 441) Long yellow needles (from water) Converted by boiling $KOHAq$ into nitro pyrocatechin [170°]

Nitro-α-amido phenol. Acetyl derivative of the methyl ether $C_6H_7(NO_2)(NHAc)(OMe)$ [143°] Formed by nitrating $C_6H_7(NHAc)(OMe)$ [1 2] (Muhlhauser, *A* 207, 242) Yellow needles (from alcohol)

Nitro-amido-phenol [134°]. Formed by boiling nitrated m phenylene diamine with aqueous KOH (Barbaglia, *B* 7, 1259) Orange plates

Nitro m amido phenol Methyl ether $C_6H_7(NO_2)(NH_2)(OMe)$ [4 3 1] [129°] Formed by heating (4, 3, 1) di nitro anisole with alcoholic NH_3 at 190° (Bantlin, *B* 11, 2106) Yellow leaflets (by sublimation)

Nitro-p-amido-phenol $C_6H_7(NO_2)(NH_2)(OH)$ [3 4 1] [148°] Got by saponifying its acetyl derivative (Hahle, *J pr* [2] 43, 63) Dark-red prisms, forming a violet solution in alkalis— KA' HA/HCl plates or prisms

Di-acetyl derivative [147°] Formed from di acetyl p amido phenol and fuming HNO_3 , Pale yellow prisms (from dilute alcohol)

Methyl ether $C_6H_7(OMe)(NO_2)(NH_2)$ [123°] Formed in small quantity by distilling $C_6H_7(OMe)(NO_2)NMe_2OH$, a crystalline compound got by mixing nitro-p amido-phenol with NMe_2OH . Crystals Yields $C_6H_7(OMe)(NO_2)(NHAc)$ and $C_6H_7(OMe)(NO_2)(NH_2)Cl$

Nitro-p-amido-phenol $C_6H_7(NO_2)(NH_2)OH$ [206°] Formed by boiling its m nitro-benzoyl derivative with alkalis (Hübner, *A* 210, 883) Colourless needles (containing aq) melting at

188° or anhydrous golden needles, melting at 206° —KA' 1:1aq red silky needles —NaA' 2aq —BaA' 4aq

Nitro benzoyl derivative

$C_6H_5(NO_2)(NH_2)O \cdot CO \cdot C_6H_5NO$ [225°] Formed by nitrating benzoyl-*p*-amido phenol [228°] Yellow needles (from HOAc)

Nitro-di-amido-phenol $C_6H_4N_2O_4$ 1, 2

$C_6H_4(NO_2)(NH_2)_2OH$ [4 6 2 1] Formed by reducing picric acid with aqueous ammonium sulphide (Griess, *A* 154, 202) Long dark yellow needles (containing aq) or narrow plates, sl sol water and alcohol, v sl sol ether —(HA') $_2$ H $_2$ SO $_4$ 5aq yellowish needles —BaA' $_2$ 2aq ruby red needles

Nitro-di-amido phenol Dibenzoyle derivative $C_6H_4(NO_2)(NHBz)_2OH$ [167°-170°] Formed by nitrating di benzoyl (a) diamido phenol

Nitro-di-amido phenol Dibenzoyle derivative $C_6H_4(NO)(NHBz)_2OH$ [201°] Formed by nitrating (6, 2, 1) di-amido phenol (Post a Stuckenberg, *A* 205, 79) Long brown needles, sl sol alcohol

Di-nitro-o-amido phenol $C_6H_3N_2O_4$ 1, 2 $C_6H_3(NO_2)_2(NH_2)(OH)$ [6 4 2 1] *Pyramic acid* [170°] S 14 at 22° (Darney, *Am* 5, 36)

Formation—1 By reduction of picric acid (Wöhler, *P* 13, 488, Girard, *A* 88, 281, *J* 1855, 535, Pugh, *A* 96, 83, Lea, *J* 1861, 637 2 By nitration of nitro *o* amido phenol (Stuckenberg, *A* 205, 75), or of benzoyl-*o* amido phenol (Hubner, *A* 210, 392)

Properties—Red needles. Converted by the diazo reaction into $C_6H_3Cl(NO_2)_2$ [110°] The salts do not explode when struck Cyanogen passed into its alcoholic solution forms 'ethoxycarbamidamido - dinitrophenol' $C_6H_3N_2O_4$ (Griess, *B* 15, 448), a crystalline body converted by boiling HClAq into uramidodinitro phenol

Salts—NaA' aq S 206 at 15° Dark red crystalline crusts (Smolka, *M* 8, 391) —NH $_4$ A' orange red tables —KA' —BaA' $_2$ —CuA' $_2$ —MgA' $_2$ 3aq S 558 at 17° —ZnA' $_2$ aq S 017 at 23° —CdA' $_2$ aq S 08 at 23°, 314 at 100° —Hg $_2$ A' $_2$ red powder —HgA' aq yellow needles S 032 at 18°, 08 at 100° —PbA' $_2$ red needles S 038 at 20°, 067 at 100° —MnA' $_2$ 2aq S 1026 at 19° —CoA' S 081 at 100° —NiA' S 0286 at 100° —AgA' —HA'HCl reddish brown needles (Petersen, *Z* 1868, 378) —(HA'HCl).PtCl $_4$

Acetyl derivative $C_6H_3(NH_2)(NO_2)_2(OAc)$ [193°] (Schiff, *B* 19, 849)

Methyl ether $C_6H_3(NH_2)(NO_2)_2(OMe)$ Dark violet needles (from alcohol), insol cold Aq

Nitro-m-amido-phenol

$C_6H_3(NO_2)(NH_2)(OH)$ [6 4 3 1] [225°] Formed by warming di nitro aniline with alcoholic KOy (Lippmann a Fleissner, *M* 7, 96) Brownish-red crystals, v sl sol water Yields di nitro-resorcin on warming with aqueous alkalis Its salts explode on heating —KA' —BaA' $_2$ —HgA' aq

Di-nitro-amido-phenol $C_6H_3(NO_2)_2(NH_2)(OH)$ [202°] Formed by the action of aqueous NH $_3$ on (3) tri nitro-phenol (Henriques, *A* 215, 334) —KA' aq

Di-nitro amido-phenol

$C_6H_3(NO_2)_2(NH_2)(OH)$ [6 2 4 1] *Isopiramic*

acid [170°] S 082 at 22°, 81 at 100°. Formed by heating its benzoyl derivative with HClAq (Dabney, *Am* 5, 33) Yellowish brown needles (from water) —KA' bluish black crystals (from alcohol) Explodes when heated

Benzoyl derivative

$C_6H_3(NO_2)(NHBz)(OH)$ [250°] Formed by heating $C_6H_3(CO_2H)(NHBz)(OH)$ [15 2] dissolved in HOAc with HNO $_3$ at 80° (D) Yellow plates (from alcohol) —KA' aq —BaA' $_2$ 3aq —CaA' $_2$ 4 $\frac{1}{2}$ aq —PbA' $_2$

Di-nitro-o-amido-phenol Benzoyl derivative $C_6H_3(NO_2)(NHBz)OH$ [4 3? 2 1] [220°] Formed by nitrating benzoyl *o* amido phenol in HOAc at -4° (Hubner, *A* 210, 387) Greenish yellow needles, insol water On treatment with POCl $_3$ it yields $C_6H_3(NO_2)_2 \begin{smallmatrix} <O \\ >N \end{smallmatrix} C_6H_5$ [219°]

—KA' 2aq —NH $_4$ A' aq —BaA' $_2$ 5aq —ZnA' $_2$ 3aq. —AgA' red needles

Di nitro-o-amido phenol Acetyl derivative of the methyl ether $C_6H_3(NO_2)_2(NHAc)(OMe)$ [157°] Formed by nitration of *o* acetanilidine (Mühlhauser, *B* 13, 921, *A* 207, 234) Prisms

Tri nitro-amido-phenol Ethyl ether $C_6H_3(NO_2)_3(NH_2)(OEt)$ Formed by heating $C_6H_3(NO_2)_3(OEt)NHCO_2Et$ with dilute H $_2$ SO $_4$ (Kohler, *J pr* (2) 29, 283) Small yellow needles (from alcohol)

o **NITRO-*p*-AMIDO-DIPHENYL** $C_{12}H_9N_2O_4$ 1, 2 $C_6H_4(NO_2)C_6H_4NH_2$ [98°] Formed by reducing *op* di nitro diphenyl with ammonium sulphide (Schultz, *A* 174, 225, 207, 350) Reddish brown monoclinic crystals, *abc* = 1 52 1 2 19, β = 69° 31' —B'HCl needles

p Nitro *p*-amido diphenyl

$C_6H_4(NO_2)C_6H_4NH_2$ [198°] Formed by reducing *pp* di nitro diphenyl with alcoholic ammonium sulphide in the cold (Fittig, *A* 124, 273, Schultz, *A* 174, 222) Small red needles (from alcohol) Gives *p*-nitro benzoic acid on oxidation —B' H PtCl $_6$

Nitro p amido diphenyl Benzoyl derivative $C_6H_4C_6H_3(NO_2)(NHBz)$ [1 3 4] [143°] Formed by nitrating benzoyl *p* amido diphenyl (Hubner, *A* 209, 339) Needles (from HOAc) Reduced by tin and HOAc to

$C_6H_5C_6H_3 \begin{smallmatrix} <N \\ >NH \end{smallmatrix} C_6H_5$ [198°]

Di nitro-p amido diphenyl Benzoyl derivative $C_6H_4(NO_2)_2NHBz$ [206°] Formed by nitrating benzoyl *p* amido diphenyl (Hubner) Dark yellow needles (from HOAc)

Nitro-di-amido-diphenyl

[4 1] $C_6H_3(NH_2)C_6H_3(NO_2)(NH_2)$ [1 2 4] [143°] Formed by mixing benzidine sulphate (28 g) in H $_2$ SO $_4$ (300 g) with KNO $_3$ (10 g) (Tauber, *B* 23, 796) Long red needles —B'H $_2$ SO $_4$ 3aq

Di-nitro di-p amido-diphenyl $C_{12}H_9N_2O_4$ 1, 2 $C_6H_3(NO_2)NH_2$ [1 3 4]

Di-nitro benzidine

$C_6H_3(NO_2)NH_2$ [1 3 4] [221°] Formed by nitration of di acetyl benzidine and saponification of the product with KOH (Brunner a Witt, *B* 20, 1024, cf. Strakosch, *B* 5, 237) Obtained also by hydrolysis of di nitro di phthalyl benzidine (Bandrowski, *M* 8, 471) Red needles Sol phenol, v sl sol alcohol, insol water Its tetrazo compound combines with α -naphthylamine *p* sulphonic acid to form a dye stuff, which dyes

unmordanted cotton the shade of alizarine-violet. By SnCl_2 and HCl it is reduced to tetra-amido diphenyl

Di-acetyl derivative [above 300°]

Di-nitro-di-p-amido-diphenyl [197°]. Obtained, together with the preceding, by hydrolysis of di nitro di phthalyl-benzidine (Bandrowski) Yellow needles (from alcohol)

Di-nitro-di-amido-diphenyl

$[\text{4 } 8 \text{ 1}] \text{C}_6\text{H}_4(\text{NH}_2)(\text{NO}_2) \text{C}_6\text{H}_4(\text{NO}_2)(\text{NH}_2) [\text{1 } 2\text{or } 3 \text{ 4}]$
Di nitro benzidine [214°] Formed by stirring KNO_3 (20 g) into benzidine sulphate (28 g) dissolved in H_2SO_4 (300 g) (Tauber, *B* 23, 795) Yellow plates (from alcohol) Its azo compounds do not dye cotton

NITRO-AMIDO-PHENYL-ACETIC ACID

$\text{C}_6\text{H}_4\text{N}_2\text{O}_6$, $\pm \text{C}_6\text{H}_4(\text{NO}_2)(\text{NH}_2) \text{CH CO}_2\text{H}$ [2 4 1] [186°] Formed by reducing (4,2,1) di nitro-phenyl acetic acid with aqueous ammonium sulphide (Gabriel & Meyer, *B* 14, 824) Reddish yellow needles, v sol hot water and alcohol, sl sol ether. Forms salts with acids and bases — $\text{HA}'\text{HCl}$ colourless needles

Methyl ether MeA' [94°]

Ethyl ether EtA' [100°] Yellow needles

Nitro-amido-phenyl-acetic acid

$\text{C}_6\text{H}_4(\text{NO}_2)(\text{NH}_2)\text{CH}_2\text{CO}_2\text{H}$ [3 4 1] [144°] Prepared by saponification of its nitrile (nitro amido benzyl cyanide) by boiling with HCl (Gabriel, *B* 15, 836) Orange-yellow plates or needles Sol alcohol and ether, insol CS_2 . By the action of amyl nitrite and HCl it gives (3,4,1) nitro diazo- nitroso toluene $\text{C}_6\text{H}_4(\text{N}_2\text{Cl})(\text{NO})(\text{CH NO})$

Nitrile $\text{C}_6\text{H}_4(\text{NO}_2)(\text{NH}_2) \text{CH CN}$ [118°] Formed by saponifying its acetyl derivative with potash Orange plates, sol water and alcohol

Acetyl derivative of the nitrile

$\text{C}_6\text{H}_4(\text{NO}_2)(\text{NHAc}) \text{CH CN}$ [113°] Formed by nitrating $\text{C}_6\text{H}_4(\text{NHAc}) \text{CH}_2\text{CN}$ (Gabriel) Flat yellow needles or plates, sol alcohol and hot water

m-Nitro- α -amido-phenyl-acetic acid

$\text{C}_6\text{H}_4(\text{NO}_2) \text{CH}(\text{NH}_2) \text{CO}_2\text{H}$ [172°] Formed by adding HNO_3 (1 mol) to a cold solution of α -amido phenyl acetic acid in H_2SO_4 (Plöchl & Loé, *B* 18, 1179) Silky needles, v sol hot water, insol alcohol — CuA' , pale blue needles

DI-NITRO-AMIDO-DIPHENYLAMINE

$\text{C}_{12}\text{H}_{10}\text{N}_4\text{O}_6$, $\pm \text{C}_6\text{H}_4(\text{NH}_2) \text{NH C}_6\text{H}_4(\text{NO}_2)_2$ [1 2 4] [177°] Formed by the action of chloro di nitro benzene on *p* phenylene diamine in alcoholic solution in presence of NaOAc (Nietzki, *B* 23, 1852) Brownish red plates, sl sol alcohol — $\text{B}'\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OH}$ brown needles

Acetyl derivative $\text{C}_{12}\text{H}_8\text{AcN}_4\text{O}_6$ [238°] Red needles

Di nitro-amido diphenylamine [172°] Formed from *m* phenylene-diamine and $\text{C}_6\text{H}_5\text{Cl}(\text{NO}_2)_2$ (Leymann, *B* 15, 1287)

NITRO - AMIDO - PHENYL - ISOBUTYRIC

ACID $\text{C}_6\text{H}_4\text{N}_2\text{N}_2\text{O}_6$, $\pm \text{C}_6\text{H}_4(\text{NO}_2)(\text{NH}_2) \text{CH}_2\text{CHMe CO}_2\text{H}$ [138°] Formed by reducing di-nitro isobutyric acid with ammonium sulphide (Edeleanu, *C. J* 53, 559) Bright red plates Reduced by long boiling with ammonium sulphide to the compound

$[\text{1 } 5] \text{C}_6\text{H}_4(\text{NH}_2) \text{CH}_2\text{CHMe CO}_2\text{H}$ [216°]

p NITRO - o AMIDO-PHENYL-CARBAMIC ETHER $\text{C}_6\text{H}_4(\text{NO}_2)(\text{NH}_2) \text{NH.CO}_2\text{Et}$ [4 2 1]

[162°] Formed by reduction of di-nitro phenyl-urethane with hot aqueous NH_4HS (Hager, *B* 17, 2630) Orange red needles or prisms. V sol alcohol, v sl sol water On heating above its melting point it loses EtOH , and is converted into nitro phenylene urea with the formula $\text{C}_6\text{H}_4(\text{NO}_2) \text{<NH>CO}$

NITRO - AMIDO - PHENYL - ETHANE ν

NITRO AMIDO ETHYL BENZENE

NITRO-AMIDO DI-PHENYL-ETHYLENE $\text{C}_{12}\text{H}_{12}\text{N}_2\text{O}_6$, $\pm \text{C}_6\text{H}_4(\text{NO}_2) \text{CH CH C}_6\text{H}_4\text{NH}_2$ [230°] Formed by reducing di nitro-di-phenyl-ethylene with alcoholic ammonium sulphide (Strakosch, *B* 6, 329) Purple plates (from nitrobenzene) — BHCl silky needles, decomposed by water

Nitro-amido-di-phenyl-ethylene

$[\text{2 } 1] \text{C}_6\text{H}_4(\text{NO}_2) \text{CH CH C}_6\text{H}_4(\text{NH}_2)$ [1 2] Formed by reducing di o nitro stilbene, formed by the action of alcoholic potash on o nitro benzyl chloride (Bischoff, *B* 21, 2077) Amorphous mass, sol ether and alcohol

Di-nitro-amido phenyl ethylene ν **DI NITRO-AMIDO STYRENE**

p NITRO - DI-p-AMIDO-TRI - PHENYL - METHANE $\text{C}_6\text{H}_4(\text{NO}_2) \text{CH}(\text{C}_6\text{H}_4\text{NH}_2)_2$ Prepared by heating aniline sulphate (28 pts) with *p*-nitro benzoic aldehyde (15 pts) and ZnCl_2 (20 pts) at 100° (Fischer, *B* 15, 677) Large garnet red crystals (containing C_6H_5) Yields paralucaniline on reduction with zinc and HCl — $\text{B}''\text{H}_2\text{Cl}_2$ needles

m Nitro-di-p amido-tri-phenyl-methane

$[\text{3 } 1] \text{C}_6\text{H}_4(\text{NO}_2) \text{CH}(\text{C}_6\text{H}_4\text{NH}_2)_2$ [136°] Prepared by heating *m* nitro benzoic aldehyde with aniline hydrochloride and ZnCl_2 (Fischer & Ziegler, *B* 13, 671) Light yellow crystals, sol ether Crystallises with C_6H_5 in yellow concentric crystals [81°]

NITRO AMIDO-PHENYL-(α)-NAPHTHYLAMINE $\text{C}_{16}\text{H}_{13}\text{N}_3\text{O}_6$, $\pm \text{C}_{10}\text{H}_7\text{NH C}_6\text{H}_4(\text{NO}_2) \text{NH}_2$ [1 4 2] [147°] Formed by reducing di nitro phenyl (α) naphthylamine with alcoholic ammonium sulphide (Heim, *B* 21, 2302) Dark yellow needles (from dilute alcohol) Gives a dark green solution in H_2SO_4

Nitro-amido phenyl-(β)-naphthylamine

$\text{C}_{16}\text{H}_{13}\text{NH C}_6\text{H}_4(\text{NO}_2) \text{NH}_2$ [1 4 2] [195°] Formed by reducing di nitro phenyl (β) naphthylamine (Heim, *B* 21, 590) Needles or prisms, m sol alcohol Dyes silk golden yellow Conc H_2SO_4 forms a yellow solution turned green by heating

Acetyl derivative $\text{C}_{16}\text{H}_{11}\text{AcN}_2\text{O}_6$ [200°] Orange red needles (from alcohol)

p NITRO - α AMIDO-PHENYL-PROPIONIC ACID $\text{C}_6\text{H}_4\text{N}_2\text{N}_2\text{O}_6$, $\pm \text{C}_6\text{H}_4(\text{NO}_2) \text{CH}_2\text{CH}(\text{NH}_2) \text{CO}_2\text{H}$ Formed from α amido phenyl propionic acid, H_2SO_4 , and HNO_3 (Erlenmeyer & Lipp, *A* 219, 213) Fluffy white mass (from alcohol), or prisms in stars (containing 14aq) (from water) Sl sol alcohol, m sol water, insol ether, v sol ammonia. Neutral to litmus Bitter-sweet taste Turns brown at 220°, decomposes at 240°-245° Gives, on oxidation by chromic mixture, *p*-nitro-benzoic acid Boiled with KOH it gives off NH_3 — $\text{HA}'\text{HCl}$ Needles in rosettes — CuA' , 2aq

Nitro amido- β -phenyl-propionic acid

$[\text{3.4.1}] \text{C}_6\text{H}_4(\text{NO}_2)(\text{NH}_2) \text{C}_6\text{H}_4 \text{CO}_2\text{H}$ **Nitro amido- β -phenyl-propionic acid**

hydrocinnamic acid. [145°] Red crystals
Sol water, alcohol, ether, and benzene The
acetyl derivative is formed by nitration of *p*-
amido β phenyl-propionic acid

Acetyl derivative [174°] Long yellow
needles Sol alcohol and benzene, sl sol cold
water and ether (Gabriel, *B* 15, 844)

Nitro-amido-phenyl-propionic acid
[2 4 1] $C_6H_4(NO_2)(NH_2)CH_2CH_2CO_2H$. [189°].
Prepared by reduction of di nitro phenyl propi-
onic acid with aqueous ammonium sulphide
(Gabriel & Zimmernann, *B* 12, 601) Plates
or flat needles Sol alcohol, ether, and acetic
acid, insol CS₂

Di-nitro-amido β -phenyl-propionic acid
[5 3 4 1] $C_6H_4(NO_2)_2(NH_2)CH_2CH_2CO_2H$ [194°]
Formed by heating $C_6H_4(NO_2)(OMe)CH_2CH_2CO_2H$
with ammonia in sealed tubes at 100° (Stöhr, *A*
225, 87) Yellow needles, v sl sol water
Does not form salts with acids.— NH_4A' —
BaA', 1:4aq

Methylether MeA' [102°]

Ethyl ether EtA' [95°]

DI-NITRO-AMIDO-PHENYL-TOLYL-AMINE
 $C_6H_4(NH_2)NHOC_6H_4(NO_2)_2$ [184°] Formed
from tolylene-diamine [99°] and $C_6H_4Cl(NO_2)_2$
(Leymann, *B* 15, 1287). Red tables

Formyl derivative [157°]

Acetyl derivative [164°]

***p*-NITRO-DI-*o*-AMIDO-PHENYL-DI-TOLYL-
METHANE** $C_6H_4(NO_2)CH(C_6H_4NH_2)_2$

(α)-Isomeride [172°] Formed from *p* nitro-
benzoic aldehyde, *p* toluidine, HClAq and alcohol
(Bischler, *B* 20, 3302) Crystallises from
benzene in needles (containing C_6H_6), v sl sol
cold alcohol —B'H₂PtCl₆

(β)-Isomeride [127°] Formed from *p*-nitro-
benzoic aldehyde, *p* toluidine, and conc H₂SO₄
(Bischler, *B* 20, 3304) Yellow plates, v sol
benzene and warm alcohol, sl sol ligroin —
B'H₂Cl₆—B'H₂PtCl₆

Di-acetyl derivative [136°]

Di-benzoyl derivative [152°]

***m*-Nitro-di-*o*-amido-phenyl-di-tolyl-methane**
[3.1] $C_6H_4(NO_2)CH(C_6H_4NH_2)_2$

(α)-Isomeride [128°] Formed by the action
of HCl on a mixture of *m* nitro benzoic aldehyde
and *p*-toluidine

(β)-Isomeride [86°] Formed from *m*-nitro-
benzoic aldehyde, *p* toluidine and H₂SO₄
(Bischler, *B* 21, 3207) Yellowish needles, v
sol hot alcohol —B'H₂PtCl₆

Di-acetyl derivative [104°]

Di-benzoyl derivative [148°]

***p*-Nitro-di-*m*-amido-phenyl-di-tolyl-methane**
[4 1] $C_6H_4(NO_2)CH(C_6H_4NH_2)_2$ Prepared by
heating *o*-toluidine sulphate with *p* nitro-benzoic
aldehyde and ZnCl₂ at 100° (Fischer, *B* 15,
679) Small yellow crystals (containing C_6H_6)

***m*-NITRO-DI-AMIDO-PHENYL-DI-XYLYL-
METHANE** $C_6H_4(NO_2)CH(C_6H_4MeNH_2)_2$ [92°].
Formed by condensing *m* nitro benzoic alde-
hyde with *m*-xyldine (Bischler, *B* 21, 3216).
Plates —B'H₂Cl₆, yellow plates —B'H₂PtCl₆

Acetyl derivative [132°] Needles

Benzoyl derivative [186°]

***p*-Nitro-di-*o*-amido-phenyl-di-xylyl-methane**
 $C_6H_4(NO_2)CH(C_6H_4MeNH_2)_2$ [90°] Formed
by condensing *p*-nitro-benzoic aldehyde with
m-xyldine by H₂SO₄ (B) Yellow needles.—
B'H₂Cl₆—B'H₂PtCl₆

Di-acetyl derivative. [88°]

Di-benzoyl derivative [192°] Needles

**DI-NITRO-DI-AMIDO-DIPHENYL SUL-
PHONIC ACID**

$C_6H_4(NO_2)(NH_2)C_6H_4(NO_2)(NH_2)SO_3H$ [1 3 4 5]
Obtained from di-acetyl di amido-diphenyl sul-
phonic acid by nitration and saponification
(Zehra, *B* 23, 3460) —KA' aq

Di-acetyl derivative Yellow needles

***s*-DI-NITRO-DI-AMIDO-QUINONE**

$C_6(NO_2)_2(NH_2)_2O$ [1 4 2 5 3 6] Prepared by dis-
solving *s* di amido di-amido benzene nitrate
 $C_6H_4(NH_2)(NH_2)(HNO_2)_2$ (1 pt) in conc H₂SO₄
(15 to 20 pts) at about 10° On adding lumps
of ice to the mixture, the compound crystallises
out in dark-yellow needles It is practically
insoluble in all indifferent solvents Very weak
base, whose salts are readily decomposed by
water By warming with dilute KOH, ammonia
is evolved, and the K salt of nitranilic acid
 $C_6(NO_2)_2(OK)_2O$ separates out Bystannouschlor-
ide it is reduced to tetra amido hydroquinone
 $C_6(NH_2)_4(OH)_2$ [1 2 4 5 3 6] (Nietzki, *B* 20, 2115)

NITRO-AMIDO-RESORCIN $C_6H_4N_2O_4$ *res*
 $C_6H_4(NO_2)(NH_2)(OH)_2$ [170°] Formed by re-
duction of di nitro resorcin with ammonium
sulphide (Benedikt & Hubl, *M* 2, 325) Dark-
brown crystals, sl sol water, v sol alcohol.—
B'H₂SO₄, needles

Di nitro-amido resorcin

$C_6H_4(NO_2)_2(NH_2)(OH)_2$ [190°] Obtained by re-
ducing tri nitro-resorcin (styphnic acid) with
alcoholic ammonium sulphide (B a H) Cop-
pery leaflets, insol water, sl sol alcohol

DI-NITRO-*p*-AMIDO-STYRENE

$C_6H_4(NO_2)(NH_2)CHCH(NO_2)$ Formed by ni-
trating *p* amido cinnamic acid, CO₂ being split
off (Friedländer & Lazarus, *A* 229, 247)
Slender reddish brown needles (from alcohol).
Conc H₂SO₄ gives off CO₂ forming a nitro amido-
benzoic aldehyde

Acetyl derivative [252°]

Di nitro-amido styrene

$C_6H_4(NO_2)(NH_2)C_6H_4$ *Acetyl derivative*.
[212°], needles, sl alcohol and acetic acid, sl
sol hot water, nearly insol ether, formed by
nitration of acetyl *p*-amido cinnamic acid
(Gabriel & Herzberg, *B* 16, 2041)

NITRO-AMIDO-TOLUENE *v* NITRO TOLU-
IDINE

**NITRO-AMIDO-TOLUENE *exo* SULPHONIC
ACID** $C_6H_4(NO_2)(NH_2)CH_2SO_3H$ Formed by
reducing $C_6H_4(NO_2)_2CH_2SO_3H$ by NH₃ and H₂S
(Mohr, *A* 221, 226) Needles, sol water —KA'.
—BaA', 2aq

Nitro-amido toluene sulphonic acid

$C_6H_4Me(NO_2)(NH_2)SO_3H$ [1 2 4 5] *S* 17 at
11° Obtained by sulphonating (2,1,4)-nitro-
toluidine (Foth, *A* 230, 300) Yellow needles
(from water) —KA' aq —BaA', 4aq

Nitro-amido-toluene sulphonic acid

$C_6H_4Me(NO_2)(NH_2)SO_3H$ [1 3 2 5] Formed from
acetyl-*o*-toluidine by successive sulphonation and
nitration (Nietzki & Pollini, *B* 23, 138)

NITRO-AMIDO-*p*-TOLUIC ACID

$C_6H_4Me(NO_2)(NH_2)CO_2H$ [1 2 3 4] [245°].
Formed by saponifying its acetyl derivative
(Niementowski, *J* pr [2] 40, 27) Yellow needles,
sol hot water —KA' 2aq reddish-yellow needles

Acetyl derivative [210°] Formed from

acetyl-amido toluic acid and HNO_3 in the cold. Yellow needles, insol water

Nitro-amido toluic acid

$\text{C}_6\text{H}_4\text{Me}(\text{NO}_2)(\text{NH}_2)\text{CO}_2\text{H}$ [1 2 4 5] [236^o]. Formed by heating bromo nitro-toluic acid with alcoholic NH_3 at 180° (Fileti & Cross, *G* 18, 298). Silky yellow needles (containing aq)

DI-NITRO-DI-AMIDODE-DITOLYL [3 5 4 1]

$\text{C}_6\text{H}_4\text{Me}(\text{NO}_2)(\text{NH}_2)\text{C}_6\text{H}_4\text{Me}(\text{NO}_2)(\text{NH}_2)$ [1 3 5 4] [267^o]. Formed by saponifying its acetyl derivative (Gerber, *B* 21, 746). Garnet red needles

Di-acetyl derivative

Obtained from di-amido-ditolyl Crystals, decomposing at 320°

NITRO-AMIDO-XYLENE v Nitro-XYLIDINE

NITRO-AMIDO-XYLENE SULPHONIC ACID

$\text{C}_6\text{H}_4\text{N}_2\text{SO}_3$ *vs* $\text{C}_6\text{H}_4\text{Me}_2(\text{NO}_2)(\text{NH}_2)\text{SO}_3\text{H}$ [1 3 2or5 4 6]. Formed by nitration of *m* xylidine sulphonic acid (Sartig, *A* 230, 838). Slender needles (from water), sl sol cold water, insol alcohol— $\text{KA}'1\frac{1}{2}\text{aq}$ — $\text{BaA}'1\frac{1}{2}\text{aq}$ — $\text{PbA}'2\text{aq}$

NITRO-AMYLENE $\text{C}_6\text{H}_4\text{NO}_2$ *vs*

$\text{CH}_3\text{CH}(\text{NO}_2)\text{C}_6\text{H}_4$. Formed from allyl iodide and potassium nitro ethane (Gal, *J* 1873, 833). Oil. May be reduced to $\text{C}_6\text{H}_5\text{NH}_2$ (85°)

Nitro amylene $\text{CH}_3\text{C}(\text{NO}_2)\text{CMe}_2$ [166°–170°]. Formed from di methyl ethyl carbinol and conc HNO_3 (Hattinger, *M* 2, 289). Oil, sol alcohol and ether. Dissolves in alkalis and gives a blue colour with KNO_3 and H_2SO_4 . On heating with HClAq it gives NH_3 , hydroxylamine, and acetic acid. On heating with water it yields nitro ethane and a ketone. NaOEt gives a yellow pp

NITRO ANILIC ACID v DI-NITRO DI OXYQUINONE

o NITRO ANILINE $\text{C}_6\text{H}_4\text{N}_2$ *vs*

$\text{C}_6\text{H}_4(\text{NO}_2)\text{NH}_2$ [2 1] *o* Nitraniline Mol w 138 [71 57]

Formation—1 By heating *o* bromo nitro benzene with alcoholic NH_3 (Walker & Zincke, *B* 5, 114)—2 Together with *p* nitro-aniline by nitration of acetanilide and saponification of the product (Körner)—3 By heating *o*-nitro-anisole $\text{C}_6\text{H}_4(\text{NO}_2)(\text{OMe})$ with ammonia at 200° (Salkowski, *A* 174, 278)—4 By reduction of *o* di-nitro benzene (Rinne & Zincke, *B* 7, 1374)—5 By nitration of benzanilide and saponification of the product (Lellmann, *A* 221, 6)

Preparation—1 By splitting off the HSO_3 group from *o* nitro-aniline *p*-sulphonic acid by heating it with HCl under pressure. The sulphonic acid can be very readily prepared by sulphonation and nitration of acetanilide (Nietzki & Benckiser, *B* 18, 294)—2 Twelve pts of *o*-nitro phenol are heated with 20 pts of aqueous NH_3 (35 p c) at 160°–170° for 16 hours, the product is crystallised from water, the yield being about 60 p c of the *o* nitro phenol employed (Merz & Ruz, *B* 19, 1749)

Properties—Orange needles, m sol hot water, v sol alcohol, v e sol ether. Volatile with steam. Does not combine with chlorinated quinones (Niemeyer, *A* 228, 322)

Salts— B'HOI plates. Decomposed by water into HCl and *o*-nitro aniline

Formyl derivative $\text{C}_6\text{H}_4(\text{NO}_2)(\text{NHCHO})$

[122°]. Needles (Hübner & Herff, *A* 209, 867).

Acetyl derivative $\text{C}_6\text{H}_4(\text{NO}_2)(\text{NEAc})$

[93°]. Yellow plates, m sol cold water

Propionyl derivative [65°] (Smith, *Am*

6, 179)

Benzoyl derivative [94°] Needles

Oxalyl derivative v Oxalic acid

m Nitro-aniline $\text{C}_6\text{H}_4(\text{NO}_2)\text{NH}_2$ [3 1] [114°] (285°) *S* 114 at 20°, *S* (alcohol) 705 at 20° (Carnelley & Thomson, *C* *J* 53, 786)

Formation—1 By reducing *m*-di nitro-benzene with H_2S and alcoholic ammonia (Hofmann & Muspratt, *A* 57, 204; Beilstein & Kurbatoff, *A* 176, 44)—2 Together with *p* nitro-aniline by adding HNO_3 to a solution of aniline in H_2SO_4 (Hübner, *A* 208, 299)

Preparation—A solution of SnCl_2 (3 mols) in alcohol saturated with HCl is slowly allowed to drop into a well cooled alcoholic solution of *m*-di nitro benzene (1 mol) with continual agitation (Anschutz & Heusler, *B* 19, 2161)

Properties—Long yellow needles. Colours pine wood yellow. Gives no colour with bleaching powder

Reactions—1 Cyanogen passed into its alcoholic solution forms a compound with formula $\text{C}_6\text{H}_4(\text{NO}_2)\text{NH}(\text{CN})(\text{NH})(\text{CN})\text{C}_6\text{H}_4\text{NO}_2$ (Senf, *J* pr [2] 35, 530)—2 Cyanogen iodide forms a green pp of $(\text{C}_6\text{H}_4(\text{NO}_2)\text{NH})_2\text{C}$ [286°] (Hübner, *B* 10, 1719)—3 Silver nitrate forms a compound $(\text{C}_6\text{H}_4(\text{NO}_2)(\text{NH}_2))_2\text{AgNO}_3$ [125°] when added to its alcoholic solution (Mixer, *Am* 1, 239)—4 Chlorinated quinones dissolved in benzene form dark green crystalline additive compounds (Niemeyer, *A* 228, 322)

Salts— B'HOI Pearly crystals, v e sol water— $\text{B'H}_2\text{PtCl}_6$ Yellow powder, v e sol water and alcohol— B'HB'r plates (Staedel & Bauer, *B* 19, 1940)— $\text{B'H}_2\text{C}_2\text{O}_4$ crystals

Acetyl derivative $\text{C}_6\text{H}_4(\text{NO}_2)(\text{NHAc})$ [150°] (Meldola & Salmon, *C* *J* 53, 778, [143°] (Meyer & Stuber, *A* 165, 183) Prisms

Benzoyl derivative [156°] Plates

p Nitro-aniline $\text{C}_6\text{H}_4(\text{NO}_2)(\text{NH}_2)$ [4 1] [147°] *S* 077 at 20°, *S* (alcohol) 584 at 20° (Carnelley & Thomson, *C* *J* 53, 786)

Formation—1 By nitration of the anilides of tartaric, succinic, or acetic acid, the product being saponified (Arppe, *A* 90, 147, 93, 157, Hofmann, *Fr* 10, 589, 12, 639), the *o* nitro aniline, which is formed at the same time, may be removed by steam distillation (Körner)—2 By heating [4 1] $\text{C}_6\text{H}_4(\text{NO}_2)(\text{OMe})$ with ammonia at 200° (Salkowski, *A* 174, 281)—3 By reduction of *p*-di nitro-benzene (Zincke & Rinne, *B* 7, 871)—4 By heating *p* chloro nitro-benzene with ammonia (Engelhardt & Latschinoff, *Z* 1870, 232)—5 By heating *p*-nitrophenol (6 pts) with aqueous NH_3 (20 pts) at 190°–200° for several hours, the yield being 58 p c (Merz & Ruz, *B* 19, 1753).

Preparation—1 Equal volumes of HNO_3 (*S* *G* 142) and H_2SO_4 are mixed and cooled. Acetanilide is gradually added as long as it will dissolve. The cold solution is set aside for half an hour, then poured into water, and the nitro compound crystallised from boiling water. The nitro acetanilide is saponified by boiling NaOH , and the nitraniline crystallised from water (Meldola, *C* *J* 43, 427)—2 1 kilo of acetanilide is slowly dissolved in 4 kilos of H_2SO_4 , kept cool by standing the vessel in a stream of cold water 590 grms of HNO_3 of *S* *G* 1478 (= 85 p c), or the corresponding quantity of ordinary HNO_3 (142) diluted with 1200 grms of H_2SO_4 is then very gradually run in, taking care that the tem-

perature does not rise above 20° After standing, the product is poured into cold water, and the yellow pp of nitroacetanilide which separates is filtered off and saponified by boiling with strong HCl The yield is very satisfactory (Nöling & Collin, *B* 17, 262, *B* J Friewell, *priv com*)

Properties—Long monoclinic needles (from water) May be readily sublimed Not volatile with steam Weak base, its salt being decomposed by water Does not combine with chlorinated quinones Cyanogen iodide at 120° forms $(C_6H_4(NO_2)_2NH)_2C$ (above 300°) (Hubner)

Salts— $BHCl$ — $B \cdot H_2PtCl_4$

Formyl derivative $C_6H_4(NO_2)(NHCHO)$ [194°] Formed from formanilide and fuming HNO_3 at -17° (Osborn & Mixer, *Am* 8, 346)

Acetyl derivative $C_6H_4(NO_2)(NHAc)$ [207°] Formed by nitrating acetanilide Prisms Gives *p* nitro phenol when boiled with conc KOHAq (Wagner, *B* 7, 76)

Benzoyl derivative [199°] Needles

Di nitro aniline $C_6H_3N_2O_4$, *ie* $C_6H_3(NO_2)_2(NH_2)$ [6 2 1] Mol w 188 [138°] S (95 p c alcohol) 52 at 12° Obtained by heating the methyl or ethyl ether of *c*-di-nitro phenol with aqueous ammonia (Salkowski & Rehs, *B* 7, 370, *A* 174, 273) Yellow needles Yields *m*-di nitro benzene on elimination of NH_3

Acetyl derivative $C_6H_3(NO_2)_2(NHAc)$ [197°]

Di-nitro aniline $C_6H_3(NO_2)_2(NH_2)$ [4 2 1] [176°] (Barr), [188°] (Hentschel, *J pr* [2] 34, 427) S (95 p c alcohol) 76 at 21° (S), (88 p c alcohol) 58 (Rudneff, *Z* 1871, 202)

Formation—1 By the action of alkalis on di nitro phenyl citraconimide (Gottlieb, *A* 85, 17) —2 By heating (1,2,4) chloro di nitro benzene with alcoholic ammonia (Clemm, *J pr* [2] 1, 145) —3 By heating [4 2 1] $C_6H_3(NO_2)_2(OMe)$ with NH_3 Aq at 100° (Salkowski, *B* 5, 872, 6, 139) —4 By heating (4,2,1) di-nitro phenol (3 g) with ammonia (10 c c of 27 p c) for 16 hours at 175° (Barr, *B* 21, 1542)

Properties—Light yellow prisms, v sl sol boiling water Does not form salts

Reactions—1 Elimination of NH_3 yields *m*-di nitro-benzene —2 Conc KOHAq forms di-nitro phenol [114°] (Willgerodt, *B* 9, 979) —3 Alcoholic KCy added slowly forms di nitro amido phenol [225°] (Lippmann & Fleissner, *M* 7, 95)

Acetyl derivative $C_6H_3(NO_2)_2(NHAc)$ [120°] Formed by nitration of acetanilide (Rudneff, *Z* 1871, 202, Ladenburg, *B* 17, 148)

Tri-nitro aniline $C_6H_2N_3O_6$, *ie* $C_6H_2(NO_2)_3(NH_2)$ [6 4 2 1] *Picramide* Mol w 228 [188°] Formed by the action of ammonia on (1,2,4,6)-chloro-tri nitro benzene (Pisani, *A* 92, 836) or on picric ethers (Liebemann & Palm, *B* 8, 278) Formed also by dissolving *p*-bromo aniline in cooled HNO_3 (S G 1 5) (Hager, *B* 18, 2578) Yellow plates with blue shimmer (from alcohol) or monoclinic tables (from HOAc) Tin and HCl reduce it to tri-amido phenol (Hepp, *A* 215, 350) Nitrous ether does not attack it. Boiling potash yields picric acid

Combinations— $(C_6H_3N_3O_6)C_6H_5$ Yellow prisms, which separate from its solution in benzene (Mertens, *B* 11, 843)— $B \cdot C_6H_5$ — $B \cdot C_6H_4H_2$ — $B \cdot C_6H_4NH_2$ [124°] (Hepp, *Bl* [2] 80, *A* 215, 350)

References—BROMO, CHLORO, and IODO

NITRO-ANILINE

NITRO ANILINE-SULPHONIC ACID *v*.

NITRO-AMIDO BENZENE-SULPHONIC ACID

NITRO-ANISIC ACID *v* *Methyl derivative of* NITRO OXY BENZOIC ACID

NITRO-ANISOLE *v* *Methyl ether of* NITRO-PHENOL

NITRO ANTHRAQUINONE $C_{14}H_8N_2O_4$, *ie*

$C_6H_4 \begin{smallmatrix} \diagup CO \\ \diagdown CO \end{smallmatrix} C_6H_4(NO_2)_2$ [2 3] Mol w 253

[220°] Prepared by nitrating anthraquinone dissolved in H_2SO_4 by HNO_3 in the cold (Roemer, *B* 15, 1786, Liebermann, *B* 16, 54) Yellow plates (by sublimation) or prismatic needles (from HOAc), sol benzene, aniline, and chloro form, sl sol alcohol and ether On reduction and treatment with nitrous acid it yields erythro oxy anthraquinone

Nitro anthraquinone $C_6H_4 \begin{smallmatrix} \diagup CO \\ \diagdown CO \end{smallmatrix} C_6H_4NO_2$

[1 4] [230°] Formed by boiling anthraquinone for half an hour with HNO_3 (S G 1 5) (Böttger & Petersen, *J pr* [2] 6, 367, *B* 6, 20, *A* 166, 147) Formed also by nitration of di bromo anthracene (Claus & Hertel, *B* 14, 978) Yellow needles (by sublimation), insol water, v sl sol ether and alcohol, m sol benzene and HOAc Yields alizarin on fusion with potash Conc H_2SO_4 (12 pts) at 200° forms 'imido oxy anthraquinone' $C_{14}H_7N_2O_4$, which sublimes in rose coloured needles

Di-nitro-anthraquinone $C_{14}H_6N_2O_6$, *ie*

$[3 \frac{1}{2}] C_6H_3(NO_2)_2 \begin{smallmatrix} \diagup CO \\ \diagdown CO \end{smallmatrix} C_6H_3(NO_2)_2$ [2 6] Mol w 298 [above 300°] Prepared by allowing anthraquinone (10 g) dissolved in H_2SO_4 mixed with HNO_3 (10 g of S G 1 48) to stand for several days It is also formed by nitrating *o* nitro anthraquinone (Roemer, *B* 16, 363) Yellow crystals (by sublimation), sol nitro benzene, sl sol xylene and HOAc, nearly insol alcohol and ether On reduction and treatment with nitrous acid it yields di oxy anthraquinone (anthrarufin) On heating with H_2SO_4 at 200° it yields four colouring matters, $C_{14}H_7N_2O_5$, $C_{14}H_7N_2O_6$, $C_{14}H_6N_2O_7$, and $C_{14}H_5N_2O_8$

Di nitro anthraquinone $C_{14}H_6N_2O_6$ [256°–260°] Formed by boiling anthraquinone with a mixture of equal volumes of H_2SO_4 and HNO_3 (S G 1 5), or by boiling anthracene with fuming nitric acid (Böttger & Petersen, *A* 160, 147, 166, 154) Minute monoclinic, almost colourless, crystals, insol water, sl sol alcohol, v sl sol ether H_2SO_4 converts it at 200° into violet 'di imido di oxy-anthraquinone' $C_{14}H_4N_2O_8$

Di nitro-anthraquinone $C_{14}H_6N_2O_6$, *Fritzsche's Reagent* [280°] Formed, together with anthraquinone, by heating anthracene with di lute nitric acid at 90° On crystallisation from alcohol it separates first (Fritzsche, *N Petersb Acad Bull* 22, 43, *Z* 1869, 114, *cf* Anderson, *A* 122, 302) Prepared by adding HNO_3 (30 g) to a solution of chrysene (50 g) containing anthracene in alcohol (5,000 c c), and heating on a water-bath The crystals of the chrysene compound (*v infra*) which then separate are oxidised by CrO_3 in HOAc, which attacks the chrysene and leaves the di nitro anthraquinone (Schmidt, *J pr* [2] 9, 268)

Properties—Yellow needles (from boiling HOAc), v sl sol alcohol and ether. Sublimes in colourless serrated plates. Forms very characteristic compounds with aromatic hydrocarbons. H_2SO_4 at 200° forms $\text{C}_{12}\text{H}_8\text{N}_2\text{O}_4$, a black powder.

Combinations— $\text{C}_{12}\text{H}_8\text{N}_2\text{O}_4(\text{PhCH CHPh})$: orange red plates— $\text{C}_{12}\text{H}_8\text{N}_2\text{O}_4(\text{C}_{10}\text{H}_8)$ violet monoclinic laminæ, obtained by dissolving di-nitro anthraquinone (9 pts) and anthracene (10 pts) in crude xylene (100 pts)—With chrysene $\text{C}_{18}\text{H}_{12}\text{N}_2\text{O}_4(\text{C}_{18}\text{H}_{12})$ Slender red needles [294°], sl sol boiling HOAc

NITRO ANTHRAQUINONE CARBOXYLIC ACID $\text{C}_{14}\text{H}_6\text{O}_4(\text{NO}_2)_2\text{CO}_2\text{H}$ [above 300°] Formed by nitration of anthraquinone carboxylic acid (Liebemann a Glock, B 17, 891) Small needles. Dissolves in H_2SO_4 with a violet colour

(a) **NITRO ANTHRAQUINONE SULPHONIC ACID** $\text{C}_{14}\text{H}_6\text{NSO}_6$, H_2NSO_6

$\text{C}_6\text{H}_4 \begin{array}{c} \text{CO} \\ \diagup \quad \diagdown \\ \text{CO} \end{array} \text{C}_6\text{H}_4(\text{NO}_2)(\text{SO}_3\text{H}) \left[\begin{array}{c} 1 \\ 6 \end{array} \begin{array}{c} 2 \\ 3 \end{array} \right]$ [255° uncor] Prepared by nitration of anthraquinone sulphonic acid with a mixture of HNO_3 (15) and H_2SO_4 , the (β) isomeric acid which is formed simultaneously remains dissolved whilst the α acid separates out (Claus, B 15, 1514, 17, 1276, Lufschütz, B 17, 899, cf Liebemann, B 16, 55) Small white plates (from dilute HNO_3) or very fine silky needles (from hot water) Strong acid. On fusion with KOH it gives alizarin

Salts—A'Na aq long needles, sol hot water, nearly insol cold water and alcohol—A'K small needles—A'NH₄ aq felted needles—A'Ca microscopic needles, sl sol water—A'Ba needles

Chloride [194° uncor] Yellow concentric needles. Nearly insol alcohol and ether

(β) **Nitro-anthraquinone sulphonic acid** $\text{C}_{14}\text{H}_6\text{O}_4(\text{NO}_2)(\text{SO}_3\text{H})$ [250° uncor] Crystalline powder. V sol water and alcohol. Strong acid. Formed as above. Fusion with KOH gives no alizarin

Salts—K, Na, and NH₄ salts are extremely soluble—A'Ba 3½ aq needles—A'Pb 2 aq white needles (Claus, B 15, 1516)

Nitro anthraquinone (α) di sulphonic acid [182°] Formed by nitration of the lead salt of anthraquinone-(α) di sulphonic acid with HNO_3 and H_2SO_4 (Claus a Schneider, B 16, 907) Yellow prisms. Sol water, alcohol, and acetic acid, insol ether, chloroform, and ligroin

DI NITRO-ANTHERONE $\text{C}_{14}\text{H}_8\text{N}_2\text{O}_4$ [116°] Formed as a by-product in the preparation of hydroanthracene-nitrite by the action of HNO_3 on an acetic acid solution of anthracene-dihydride (Liebemann a Landshoff, B 14, 472) Insol alkalis, sol benzene

DI NITRO DIANTHRYL $\text{C}_{22}\text{H}_{12}\text{N}_2\text{O}_4$, $\text{H}_2\text{N}_2\text{O}_4$
 $\text{NO}_2 \cdot \text{O} \begin{array}{c} \diagup \quad \diagdown \\ \text{C}_6\text{H}_4 \end{array} \text{C} \begin{array}{c} \diagup \quad \diagdown \\ \text{C}_6\text{H}_4 \end{array} \text{CNO}_2$ [337°]

Formed by nitration of dianthryl in acetic acid solution (Gumbel, B 20, 2433) Yellow stellate needles. V sol benzene and chloroform, sl sol alcohol and acetic acid. On reduction it gives di-amido-dianthryl [309°] By CrO_3 and acetic acid it is oxidised readily to anthraquinone. Bromine forms $\text{C}_{22}\text{H}_{10}\text{Br}_2$ [above 300°] (Sechse, B 21, 2512)

NITRO-ARACHIC ACID $\text{C}_{22}\text{H}_{40}(\text{NO}_2)_2\text{O}_2$ [70°] Formed by mixing arachic acid with HNO_3 and H_2SO_4 (Tassinari, B 11, 2031) Sl sol cold alcohol, v sol ether

NITRO-ARBUTIN v ARBUTIN

TETRA-NITRO-AURINE $\text{C}_{11}\text{H}_8(\text{NO}_2)_4\text{O}_8$ [c 140°] Formed by nitration of aurine (Acker mann, B 17, 1625) Brownish-yellow microscopic needles. V sol alcohol, nearly insol water, benzene, chloroform, and ether. Dissolves in alkalis with a dark red colour

Salts—A'Ag, brown pp—A'Ba black powder

Ethyl ether A'Et₂ [c 105°], yellow crystals, v sol alcohol and benzene, insol water and carbonated alkalis

NITRO-AZO- compounds v Azo compounds

Nitro-diazo- compounds v Di-azo compounds

NITRO-AZOXY- compounds v Azoxy compounds

NITRO-BARBITURIC ACID $\text{C}_4\text{H}_4\text{N}_2\text{O}_4$, $\text{H}_4\text{N}_2\text{O}_4$
 $\text{CO} \begin{array}{c} \text{NH CO} \\ \diagup \quad \diagdown \\ \text{NH CO} \end{array} \text{CHNO}_2$ *Dihydruric acid* Formed by the action of nitric acid on hydriuric acid and on barbituric acid (Schlieper, A 56, 23, Baeyer, A 127, 211, 130, 140) Colourless dimetric efflorescent prisms (containing 3 aq), v sol hot water forming an intense yellow solution, m sol alcohol, insol ether. Bromine and water at 100° decompose it into di-bromo barbituric acid and HNO_3 . HIAq reduces it to amido barbituric acid. Its solution gives a white pp with ammonium salts

Salts—The salts are very stable, the acid not being separated by mineral acids— $\text{NH}_4\text{H}_2\text{A}'''$ crystalline pp, v sl sol cold water— $\text{NaH}_2\text{A}'''$ 2 aq silky needles— $\text{KH}_2\text{A}'''$ Ppd by adding HCl to a solution of the acid in potash— $\text{K}_2\text{HA}'''$ yellow needles, insol alcohol and conc KOHAq. Explodes when heated— $\text{BaH}_2\text{A}'''$ Cl aq— $\text{Ca}(\text{H}_2\text{A}')_2$ 4 aq— $\text{Cu}(\text{H}_2\text{A}')_2$ 6 aq— $\text{Fe}(\text{H}_2\text{A}')_2$ 8 aq— $\text{Fe}(\text{H}_2\text{A}')_2$ 9 aq— $\text{AgH}_2\text{A}'''$ aq— $\text{Ag}_2\text{A}'''$

NITRO-BENZALDOXIM v Oxim of Nitrobenzoic aldehyde

NITRO-BENZAMIDE v Amide of Nitrobenzoic acid

m-NITRO BENZAMIDINE

$\text{C}_7\text{H}_7(\text{NO}_2)\text{C}(\text{NH})\text{NH}_2$ Formed from nitrobenzoic imido ethyl ether (Tafel a Enoch, B 23, 1552) Colourless mass (from ether), v sol water—B'HCl [240°] Tables, v sol water

m-NITROBENZAMIDOXIM $\text{C}_7\text{H}_7\text{N}_2\text{O}_4$, $\text{H}_2\text{N}_2\text{O}_4$
 $\text{C}_6\text{H}_4(\text{NO}_2)\text{C}(\text{NOH})\text{NH}_2$ [174°] Formed from m nitro benzonitrile and hydroxylamine (Schopff, B 18, 1063) Orange needles, v sol warm water. ClCO₂Et forms $\text{C}_7\text{H}_7(\text{NO})\text{C}(\text{NH}_2)\text{NO CO}_2\text{Et}$ [158°] Acetic anhydride produces the azoxim

$\text{C}_7\text{H}_7(\text{NO}_2)\text{C} \begin{array}{c} \diagup \quad \diagdown \\ \text{N} \end{array} \text{CMe}$ [109°]—B'HCl—B'H₂PtCl₄

Ethyl ether $\text{C}_7\text{H}_7\text{EtN}_2\text{O}_4$ Prisms—B'HCl **Benzyl ether** $\text{C}_7\text{H}_7(\text{OH}(\text{Ph})\text{N}_2\text{O}_4$ [58°]

p Nitro-benzamidoxim

$\text{C}_7\text{H}_7(\text{NO}_2)\text{C}(\text{NH}_2)\text{NOH}$ [169°] Formed from p-nitro-benzonitrile and hydroxylamine (Weise, B 22, 2418) Yellow needles, sol acids and alkalis. Reduces Fehling's solution and ammoniacal AgNO₃. M sol hot water and alco-

hol Can be distilled With Ac_2O it yields $\text{C}_6\text{H}_4(\text{NO}_2)\text{O} \left\langle \begin{smallmatrix} \text{N} \\ \text{O} \end{smallmatrix} \right\rangle \text{CMe}$ [144°], while aldehyde

forms $\text{C}_6\text{H}_4(\text{NO}_2)\text{C} \left\langle \begin{smallmatrix} \text{N} \\ \text{NH} \end{smallmatrix} \right\rangle \text{O} \text{CH}_2\text{CH}_3$ [158°]

ClCO_2Et produces $\text{C}_6\text{H}_4(\text{NO}_2)\text{C}(\text{NH}_2)\text{NOCO}_2\text{Et}$ [169°], which on heating yields the compound

$\text{C}_6\text{H}_4(\text{NO}_2)\text{C} \left\langle \begin{smallmatrix} \text{N} \\ \text{NH} \end{smallmatrix} \right\rangle \text{CO}$ [286°] Sodium diazo

benzene sulphonate produces the compound

$\text{C}_6\text{H}_4(\text{NO}_2)\text{C} \left\langle \begin{smallmatrix} \text{N} \\ \text{NH} \end{smallmatrix} \right\rangle \text{C}(\text{NH}_2)\text{C}_6\text{H}_4\text{NO}_2$ [151°]

(Stieglitz, *B* 22, 3157) COCl_2 reacts in benzene solution forming $\text{C}_6\text{H}_4(\text{NO}_2)\text{C}(\text{NH}_2)\text{NO}_2\text{CO}$ [283°]— B^*HCl [185°]

Ethyl ether $\text{C}_6\text{H}_4(\text{NO}_2)\text{C}(\text{NH}_2)\text{NOEt}$, [60°] Formed from the Na salt and EtI With nitrous acid it yields $\text{C}_6\text{H}_4(\text{NO}_2)\text{C}(\text{O NO})\text{NOEt}$ [55°]— B^*HCl

NITRO BENZENE $\text{C}_6\text{H}_5\text{NO}_2$. Mol w 123 [8°] (209°) at 745 mm (Brühl, *A* 200, 188) SG ρ 1.2039 (*B*) SV 121.9 (Lossen, *A* 254, 78), 121.5 (Ramsay) $\mu_s = 1.5712$ $R_\infty = 52.64$ Dispersive power Barbier & Roux, *C* R 108, 1249) Formed by nitration of benzene (Mitscherlich, *P* 31, 625) The rate of nitration has been studied by Lothar Meyer (*B* 22, 18) In small quantity by the action of ozonised air on a mixture of sulphuric acid and benzene (*L* Maquenne, *Bl* [2] 37, 298) Formed also by the action of ppd . Cu_2O (1 mol) in the cold upon an aqueous solution of diazo benzene nitrite (1 mol), obtained by slowly adding a solution of 15 g NaNO_2 in 50 cc of water to a mixture of 9 g of aniline, 20 g HNO_3 (1.4), and 50 cc of water When the evolution of nitrogen has ceased the nitro benzene is distilled off with steam, the yield is 42 p.c. of the theoretical (Sandmeyer, *B* 20, 1494)

Preparation—A mixture of nitric acid (100 pts) and H_2SO_4 (115 pts) is run into benzene The product is washed with alkali and distilled

Properties—Oil, v sol alcohol and ether Not attacked by chlorine or bromine in the cold

Reactions—1 For bromination the presence of carriers (FeBr_3 or FeCl_3) are necessary Thus nitrobenzene (10 g) heated with FeBr_3 (3 g) and bromine (4.3 cc) in sealed tubes for 10 hours at 105° gives 65 p.c. of the theoretical amount of *m*-bromo nitro benzene When the same mixture is heated for 30 hours at 120° the product is tetra bromo nitro-benzene (Scheufelen, *A* 231, 158) Bromine at 250° forms $\text{C}_6\text{H}_4\text{Br}_2$ and a little $\text{C}_6\text{H}_3\text{Br}_3$ (Kekulé, *A* 187, 169)—2 *Chlorine* in presence of FeCl_3 forms *m* chloro-nitro benzene and (5,2,1) di chloro-nitro-benzene—3 Not attacked by boiling dilute potash or ammonia. Boiling alcoholic potash forms azoxybenzene—4 Readily reduced to aniline—5 *Chromic oxychloride* forms a powder $\text{C}_6\text{H}_4(\text{NO}_2)(\text{CrO}_2\text{Cl})_2$ decomposed by water with reproduction of nitro benzene (Henderson & Campbell, *C* J 57, 253, cf Étard, *A* Ch [5] 28, 272)—6 Exposed to sunlight in alcoholic solution it is reduced to aniline, aldehyde being formed (Giamman & Silber, *B* 19, 2899, *G*. 16, 536)—7 Conc HClAq at 245° forms di-chloro-aniline (Baumbauer, *A* Suppl 7, 204)—8 Reduced in alcoholic solution by sodium-amalgam

to azo benzene (Werigo, *A* 185, 176, Alexieff, *Bl* [2] 1, 324)

o Di-nitro-benzene $\text{C}_6\text{H}_3(\text{NO}_2)_2$ [1 2] Mol w 168 [118°] S (alcohol) 3.8 at 24.8°, 3.3 at 78°. Formed in small quantity in the preparation of the *m* isomeride, and purified by successive crystallisations from alcohol and HOAc (Rinne & Zincke, *B* 7, 869, Körner, *G* 4, 354, Lobry, *R T C* 2, 239) Long colourless needles (from hot water) or monoclinic tables (from alcohol) (Bodewig, *J* 1884, 464) May be sublimed. Yields on reduction *o* nitro aniline [71°] and *o*-phenylene diamine [99°]

m Di-nitro-benzene $\text{C}_6\text{H}_3(\text{NO}_2)_2$ [1 3] [90°] (Reissert, *B* 23, 2243) (297° cor) (Meyer & Stadler, *B* 17, 2649 note) S (alcohol) 5.9 at 24.8° The chief product of the action of boiling fuming HNO_3 on benzene or nitrobenzene (Dewille, *A* Ch [3] 3, 187, Muspratt & Hofmann, *A* 57, 214) Formed also from (4,2,1)-di nitro aniline by elimination of NH_3 (Rudneff, *Z* 1871, 203) Prepared by adding benzene to a mixture of equal volumes of H_2SO_4 and fuming HNO_3 , and heating until a sample of the oily layer solidifies on cooling The product is poured into water, and the solid crystallised from alcohol (Körner, Beilstein & Kurbatoff, *A* 176, 43)

Properties—Colourless flexible needles, v e sol boiling alcohol Detonates when projected into a red hot tube filled with nitrogen (Berthelot, *A* Ch [6] 16, 24) A mixture with KClO_3 is a powerful explosive (*rackarock*)

Reactions—1 On reduction it yields *m*-nitro aniline [114°] and *m* phenylene diamine (Hofmann, *Pr* 11, 518)—2 Alcoholic KCy forms $\text{C}_6\text{H}_3(\text{NO})_2(\text{OEt})\text{CN}$ (Lobry de Bruyn, *R T C* 2, 205)—3 Alkaline K_2FeCy_4 forms (4,2,1)- and (6,2,1) di nitro phenols (Hepp, *B* 18, 2347, *A* 215, 355)

p-Di-nitro-benzene $\text{C}_6\text{H}_3(\text{NO}_2)_2$ [1 4] [172°] Deposited from the alcoholic mother liquor from which the *m* isomeride has crystallised (Körner) Flat monoclinic needles, sl sol cold alcohol May be sublimed Yields *p*-nitro aniline [146°] and *p* phenylene diamine [140°] on reduction Forms a sparingly soluble compound with naphthalene

o-Tri-nitro-benzene $\text{C}_6\text{H}_2(\text{NO}_2)_3$ [1 2 4] [57.5°] SG ρ 1.173 S (alcohol) 5.4 at 15.5°, S (benzene) 141 at 15.5° Formed by heating *p* di nitro benzene with a mixture of HNO_3 and H_2SO_4 (Hepp, *A* 215, 362, Lobry de Bruyn, *R T C* 9, 190) Yellow crystals Forms with aniline a compound $\text{C}_6\text{H}_4(\text{NO}_2)_3\text{C}_6\text{H}_4\text{NH}_2$ [84°]

Reactions—1 NaOMe in HOMe forms $\text{C}_6\text{H}_3(\text{NO}_2)_2(\text{OMe})$ [4 2 1] [88°]—2 NaOEt forms $\text{C}_6\text{H}_3(\text{NO}_2)_2(\text{OEt})$ [4 2 1] [86°]—3 Boiling aqueous Na_2CO_3 forms (4,2,1) di nitro phenol [112°] 4 Alcoholic NH_3 forms di nitro aniline [175°]

s-Tri-nitro-benzene $\text{C}_6\text{H}_2(\text{NO}_2)_3$ [1 3 5] [122°] Prepared by heating *m* di nitro benzene (40 g) with HNO_3 (120 g) and fuming H_2SO_4 (300 g) for one day at 80° and then for two days at 120°, the yield being 50 p.c (Hepp, *A* 215, 347, Claus, *B* 16, 1597) Trimetric plates, $a b c = 954.1 : 758.1$ sl sol. cold alcohol, v e sol benzene (forming a compound with C_6H_5) Not volatile with steam With aniline it forms $\text{C}_6\text{H}_3(\text{NO}_2)_3\text{NH}_2\text{Ph}$ crystallising from benzene in red plates [124°] With naphthalene it forms

$C_6H_5(NO_2)_2C_6H_5$ [158°], and with di methyl-aniline $C_6H_5(NO_2)_2NMe_2Ph$ [108°]

Reactions—1 Alkaline K_2FeO_4 oxidises it to picric acid—2 $NaOMe$ dissolved in $HOME$ forms, in the cold, $C_6H_5(NO_2)_2(OMe)$ [105°] (De Bruyn, *R T C* 9, 208)—3 Reduced by tin and $HClAq$ to tri amido benzene

References—Bromo, Chloro, Chlorotodo- and Iodo Nitro Benzene

NITRO-BENZENE-AZO compounds v Azo-compounds and Diazo compounds

NITRO-BENZENE-AZOXY- compounds v. Azoxy-compounds

DI NITRO-BENZENETETRA-CARBOXYLIC ACID $C_6(NO_2)_2(CO_2H)_4$ [1 4 2 3 5 6] *Di nitro-pyromellitic acid* Formed by oxidising di nitro-tri methyl benzoic acid with alkaline $KMnO_4$ (Nef, *A* 287, 19) Silky needles (from ether)

Ethyl ether Et_4A'' [180°] Needles

NITRO-BENZENE PHOSPHONIC ACID $C_6H_5(NO_2)PO(OH)_2$ *Nitrophosphenylic acid* [132°] *S* 98 at 22°, 92 at 98° Obtained by nitrating benzene phosphonic acid (Michaelis & Benziger, *B* 8, 1810, *A* 188, 275) Deliquescent crystals exploding above 200°— $BaA''2aq$ — $Ba(HA'')_22aq$ — $CaA''4aq$ — PbA'' — AgA''

m NITRO-BENZENE-SULPHINIC ACID $C_6H_5(NO_2)SO_2H$ [1 3] [95°] Formed, together with nitrobenzene and N_2 , by boiling the compound $C_6H_5(NO_2)NHNH_2SO_2C_6H_5(NO_2)$ with baryta water Long silky needles *V* e sol ether, sl sol alcohol

Salts— KA' small e sol prisms— AgA' long silky needles, sl sol water— BaA' , $1\frac{1}{2}aq$ yellowish prisms (Limpricht, *B* 20, 1240)

p Nitro-benzene sulphonic acid $C_6H_5(NO_2)_2SO_2H$ [1 4] [120°] Formed in the same way as the preceding acid from the corresponding *p*-compound Plates Less soluble in ether than the *m* isomeride— BaA' , *aq* yellowish prisms (Limpricht, *B* 20, 1241)

o-NITRO-BENZENE SULPHONIC ACID $C_6H_5(NO_2)SO_2H$ [1 2] Formed in small quantity in the nitration of benzene sulphonic acid (Limpricht, *A* 177, 60)— NH_4A' long needles— KA' sl sol water— BaA' , *aq* v e sol water— PbA' , *3aq*

Chloride $C_6H_5(NO_2)SO_2Cl$ [67°] **Amide** $C_6H_5(NO_2)SO_2NH_2$ [186°] Reduced by HI in $HOAc$ it yields $C_6H_5<\frac{NH}{SO_2}>$ [198°] (Cleve, *B* 20, 1534)

m Nitro benzene sulphonic acid $C_6H_5(NO_2)(SO_2H)$ [1 3] Formed by sulphonating nitro-benzene, or by nitrating benzene sulphonic acid (Schmidt, *A* 120, 163, Meyer & Stuber, *A* 165, 164, Rose, *Z* 1871, 224, Limpricht, *A* 177, 60) Formed also by the action of $ClSO_3H$ on nitro benzene in CS_2 (Armstrong, *Z* 1871, 321, Limpricht, *B* 18, 2175) Deliquescent laminae— NH_4A' Prisms— KA' *S* 17 to 19 at 7°— NaA' — BaA' , *aq* *S* (of BaA'), 4 at 21° (Goslich, *A* 180, 104), 2 at 7° (*L*)— CaA' , *2aq* *S* (of CaA'), 6 at 6°— MgA' , *4aq*— ZnA' , *3aq*— PbA' , *2aq* *S* (of PbA'), 4 at 10°

Chloride $C_6H_5(NO_2)SO_2Cl$ [61°] **Amide** $C_6H_5(NO_2)SO_2NH_2$ [161°] Reduced by HI and $HOAc$ to $C_6H_5<\frac{NH}{SO_2}>$ [88°]

p-NITRO-BENZENE SULPHONIC ACID $C_6H_5(NO_2)_2SO_2H$ [1 4] Formed in small quantity

in the nitration of benzoic acid (Limpricht)— NH_4A' Plates *S* 85 at 7°— KA' Prisms *S* 37 at 7°— BaA' , *3aq* *S* (of BaA'), 46 at 6°— CaA' , *2aq*— PbA' , *2aq* *S* (of PbA'), 118 at 55°. [The solubilities of salts here given are the weights dissolved in 100 c.c. of the solution.]

Chloride $C_6H_5(NO_2)SO_2Cl$ Oil **Amide** $C_6H_5(NO_2)SO_2NH_2$ [131°]

(a) Nitro-benzene *m*-disulphonic acid $C_6H_5(NO_2)(SO_2H)_2$ Formed, together with the (8) isomeride, by nitration of benzene *m*-disulphonic acid (Heinzelmann, *A* 188, 160, 190, 222) Slender deliquescent needles (containing *2aq*)— $(NH_4)_2A'$ flat prisms— K_2A'' — $BaA''4aq$ — $BaA''6aq$ — $BaA''6aq$ — $Ba_2A''(OH)_2$, *15aq*— $PbA''4aq$ — $PbA''O2aq$ — AgA''

Chloride $C_6H_5(NO_2)(SO_2Cl)_2$ [96°] **Amide** $C_6H_5(NO_2)(SO_2NH_2)_2$ [242°]

(8) Nitro benzene *m*-disulphonic acid $C_6H_5(NO_2)(SO_2H)_2$ [4 13] Formed as above. Very hygroscopic crystals Its salts are more soluble than those of the (a) isomeride— $(NH_4)_2A''$ — $K_2A''4aq$ (?)— $BaA''5aq$ — $PbA''4aq$

Chloride $C_6H_5(NO_2)(SO_2Cl)_2$ Oil. **Nitro-benzene disulphonic acid** $C_6H_5(NO_2)(SO_2H)_2$ Formed from nitro-amido-benzene disulphonic acid by the diazo-reaction (Limpricht, *B* 8, 289)— $PbA''aq$ needles

Di nitro-benzene sulphonic acid $C_6H_5(SO_2H)(NO_2)_2$ [1 2 4] [108°] Formed from ($C_6H_5(NO_2)_2$) $_2S_2$ and fuming HNO_3 (Willgerodt & P Mohr, *J pr* [2] 34, 117) Very hygroscopic yellow prisms (containing *3aq*), v sol water, sl sol ether, insol benzene Not attacked by fuming HNO_3 at 200°

Reactions—1 *Aniline* forms, on boiling, $C_6H_5(NO_2)(NPhH)$ [156°]—2 Boiling alcoholic ammonia forms $C_6H_5(NO_2)(NH_2)$ [178°]—3 *KHS* forms, in the cold, $C_6H_5(NO_2)_2SH$ —4 Rapidly decomposed by cold *potash*, becoming di nitro phenol [114°]

Salts— KA' — NaA' , *aq*— BaA' , *aq*— CaA' , *2aq*— ZnA' , *6aq*— PbA' , *3aq*

Chloride $C_6H_5(SO_2Cl)(NO_2)_2$ [102°] **Amide** $C_6H_5(SO_2NH_2)(NO_2)_2$ [154°]

Di-nitro-benzene sulphonic acid $C_6H_5(NO_2)_2SO_2H$ [3 2 1] Formed by warming nitro-benzene *m*-sulphonic acid with H_2SO_4 (1 vol) and HNO_3 (3 vols) (Limpricht, *B* 9, 554, Sachse, *A* 188, 143) Deliquescent crystals. NH_4A' — KA' , $1\frac{1}{2}aq$ — BaA' , *aq*— PbA' , *3aq*

Chloride $C_6H_5(NO_2)_2SO_2Cl$ [89°] **Amide** $C_6H_5(NO_2)_2SO_2NH_2$ [238°]

Di-nitro-benzene disulphonic acid $C_6H_5(NO_2)_2(SO_2H)_2$ Formed by boiling nitro-benzene *m* sulphonic acid with H_2SO_4 (1 vol) and fuming HNO_3 (6 vols) (Limpricht, *B* 8, 289) Crystalline mass Its chloride and amide are crystalline and decompose without melting— $K_2A''aq$ — $NaA''3aq$ — $BaA''2aq$ — $CaA''aq$ — $PbA''8aq$ — $CuA''3aq$

Tri-nitro-benzene sulphonic acid $C_6H_5(NO_2)_3SO_2H$ [185°] Prepared by boiling chloro-tri nitro phenol (picryl chloride) with alcohol and dry H_2SO_4 (Willgerodt, *J pr* [2] 32, 117) Large crystals (containing *2aq*), melting at 100° when hydrated Decomposed by alkalis in the cold into SO_3 and picric acid— $NaA''2aq$

NITRO-BENZENYL-AMIDO-OXIM v Nitro-benzamidoxime.

NITRO-BENZIDINE v **NITRO DI-p-AMIDODIPHENYL**

NITRO-BENZIL $C_6H_5(NO_2)O_2$. [142°]. Formed by nitration of benzoin or benzil (Zinnin, *A Suppl* 3, 153, Hausmann, *B* 23, 531) Yellow crystals, sl sol alcohol

(a)-*Dioxim* $C_6H_5N_2O_2$ [225°] Formed by heating nitrobenzil with hydroxylamine hydrochloride at 100°. Crystalline body, v. sl sol. alcohol.

(B)-*Dioxim* [185°] Formed by heating the (a)-isomeride with alcohol at 165° for several hours White needles, v sol alcohol Like the (a)-isomeride it is split up by conc $HClAq$ at 100° into hydroxylamine and nitro benzil

Di-nitro-benzil $C_6H_5(NO_2)_2O_2$ [131°] S (alcohol) 78 in the cold, 24 at 78° Formed, together with the following isomeride by boiling benzil with fuming HNO_3 (Zagumenny, *J R* 4, 278). Octahedra or moss like forms

Di-nitro-benzil [147°] S (alcohol) 34 in the cold, 19 at 78° Formed as above Plates Iso-di-nitro-benzil $C_6H_5(NO_2)_2O_2$ [205°] S (95 p c alcohol) 042 in the cold, 9 on boiling Formed by oxidation of (a)- or (γ) di-nitro-deoxybenzoin by CrO_3 in $HOAc$ (Golubeff, *J. R* 13, 29, *B* 17, 581) Yellow needles

o-NITRO-BENZOIC ACID $C_6H_5(NO_2)CO_2H$ [12] Mol w 167 [147 7°] (Reissert, *B* 23, 2244) S 61 at 16.5° Formed in small quantity in the preparation of the *m* isomeride by nitration of benzoic acid (Griess, *B* 8, 526, *A*, 166, 129, L Liebermann, *B* 10, 862, Widmann, *A* 193, 204) Obtained also by saponifying its nitrile Prepared by oxidising *o* nitro cinnamic acid with chromic acid mixture (Beilstein *A*, Kuhlberg, *A* 163, 134, Widmann, *B* 8, 393) or, better, by oxidising *o* nitro toluene (Weith, *B* 7, 1058, Widmann, *A* 193, 225, Noyes, *B* 16, 53, Monnet, Reverdin, a Nolting, *B* 12, 443) or *o*-nitro-benzyl chloride (Nolting, *B* 17, 385)

Properties — Small triclinic needles (from water), v sol alcohol and ether, v sl sol water Has a sweet taste. Very slightly volatile with steam

Salts. — BaA' , 3aq Triclinic crystals, v. sol. water — CaA' , 2aq needles — PbA' , aq — AgA' Crystals, v sol hot water

Ethyl ether EtA' [30°] Triclinic crystals.

Chloride $C_6H_5(NO_2)COCl$ Oil

Cyanide $C_6H_5(NO_2)CO CN$ [54°] Formed from the chloride and $AgCy$ at 100° (Claisen a Shadwell, *B* 12, 851) White prisms

Anhydride $(C_6H_5(NO_2)CO)_2O$ [135°] Needles (Bischoff a Rach, *B* 17, 2789)

Amide $C_6H_5(NO_2)CONH_2$ [176°] (Bischoff, *A* 239, 109) With KOH and bromine it yields $C_6H_5(NO_2)(CONHBr)$ converted by boiling potash into *o*-nitro-aniline (Hoogewerf a Van Dorp, *R T C* 8, 178)

Di-nitro anilside

$C_6H_5(NO_2)CONH(C_6H_5NO_2)$. [178°] Formed by nitrating benzoyl *m*-nitro-aniline (Schwartz, *B* 10, 1708)

Nitrile $C_6H_5(NO_2)CN$ *o*-Nitro-benzonitrile Mol w 148 [110°] Obtained by heating the amide with P_2O_5 at 100° (Hubner a Bartlein, *B* 10, 1718). Formed also by the action of hot cuprous potassium cyanide on *o*-nitro diazobenzene chloride (Sandmeyer, *B* 18, 1494) and by treating the oxim of *o*-nitro-benzoic aldehyde

with Ac_2O (Gabriel a Meyer, *B* 14, 2389) Needles, v sol water and alcohol

m-Nitro-benzoic acid $C_6H_5(NO_2)CO_2H$ [13]. [141°]. S 25 at 10°, 10 at 100° (Mulder); 235 at 16.5° (Beilstein) Formed by nitration of benzoic acid (Mulder, *A* 34, 297, Gerland, *A* 91, 185, Hübner, *A* 222, 73) Formed also by boiling nitro-hippuric acid with $HClAq$ (Bertagnini, *A* 78, 104, 79, 259) and by the oxidation of *m*-nitro toluene (Beilstein, *A* 132, 137, 155, 25, 163, 136) Obtained also by the action of hot cuprous potassium cyanide solution upon *m* nitro-diazo benzene chloride, and saponification of the crude nitrile, the yield being 72 p c (Sandmeyer, *B* 18, 1494)

Properties — Colourless laminæ (from water), v e sol alcohol and ether May be sublimed Crystallises in three monoclinic modifications (Bodewig, *J* 1879, 677) Yields amido benzoic acid on reduction and azoxybenzoic acid on heating with alcoholic potash (Griess, *A* 131, 92)

Salts — NH_4A' , — NaA' 3aq colourless tables — KA' aq Needles S 14 in the cold, 200 at 100° (Sokoloff, *J* 1864, 343) — MgA' , 7aq — CaA' , 2aq S 33 in the cold, 55 at 100° — $Ca(OBz)A'$ 3aq (Salkowsky, *B* 10, 1258) — SrA' , 2aq — SrA' , 4aq — BaA' , 4aq Needles S 38 in the cold, 53 at 100° (Sokoloff) S (of BaA') 22 at 9° (Mills, *C J* 19, 363) — ZnA' , 5aq — ZnA' , 4aq S 16 in the cold 77 at 100° — CdA' , 4aq (Schiff, *A* 104, 326) — PbA' , — MnA' , 4aq — FeA' , — CuA' , aq — AgA'

Methyl ether MeA' [70°] (279°) Prisms (Chancel, *Compt Chim* 1849, 179, *A*, 72, 275)

Ethyl ether EtA' [43°] (Tafel a Enoch, *B* 23, 1551) (296°) Monoclinic prisms Decomposed by bromine at 170°-200° into nitrobenzoic acid and ethylene bromide (Naumann, *A* 133, 202)

s-Tri-chloro-phenyl ether $C_6H_2Cl_3A'$. [182°] Formed by nitrating the benzoyl derivative of (6,4,2,1) tri chloro phenol (Daccomo, *B* 18, 1165)

Di bromo phenyl ether $C_6H_3Br_2A'$ v. vol i p 607

Nitro phenyl ethers v Nitro benzoyl derivatives of NITRO PHENOLS

Chloride $C_6H_5(NO_2)COCl$ [34°] (184° at 50 mm.), (275°-278°) Prisms (Cahours, *A Ch* [3] 23, 339, Hugh, *B* 7, 1267, Claisen a Thompson, *B* 12, 1942)

Cyanide $C_6H_5(NO_2)CO CN$ (231° at 145 mm) Formed from the chloride and $AgCy$ (*C. a T*) Heavy oil

Anhydride $(C_6H_5(NO_2)CO)_2O$ Solid (Gerhardt, *A* 87, 158)

Acetic m-nitro-benzoic anhydride $C_6H_5(NO_2)CO OAc$ [45°] Formed from the Ag salt and $AcCl$ (L Liebermann, *B* 10, 863, Beilstein, *Bn* 2, 786, Greene, *Am* 11, 414)

Benzoic m-nitro-benzoic anhydride $C_6H_5(NO_2)CO OBz$ Crystalline (Gerhardt)

Amide $C_6H_5(NO_2)CONH_2$ [142°] Needles (Field, *A* 55, 45, Chancel, *Compt Chim* 1849, 180, Beilstein, *A* 132, 137, Schiff, *A* 213, 185) Its alcoholic solution mixed with $AgNO_3$ and $NaOH$ gives a gelatinous pp of $C_6H_5(NO_2)CONHAg$ (Tafel a Enoch, *B* 23, 1550) With bromine and potash it gives *m*-nitro-aniline (Hoogewerf a Van Dorp, *R. T. C.*

8, 173) With *m* nitro benzoyl chloride it is converted into its *m*-nitro benzoyl derivative [195°] (W Schulze, *A* 251, 158)

Anilide $C_6H_4(NO_2)CONHPh$ [144°]. Plates (Engler a Volkhausen, *B* 8, 34, Hubner, *B* 9, 774)

m-Nitro-anilide
 $C_6H_4(NO_2)CONHC_2H_5$ [187°] Needles (from amyl alcohol) (McHugh, *B* 7, 1268)

Di nitro-anilide
 $C_6H_3(NO_2)_2CONHC_2H_5$ [124] [165°] Formed by nitration of the benzoyl derivatives of *o* and *p* nitro-aniline (Schwartz, *B* 10, 1703
p Toluide $C_6H_4(NO_2)CONHC_2H_5$ [162°]. Needles (from alcohol) (Hubner, *A* 210, 335)

Nitro p-toluide
 $C_6H_4(NO_2)CONHC_2H_5$ Me NO_2 [142] [183 5°]. Yellow silky needles (from alcohol) Formed by nitrating the *p* toluid

Mesidide $C_6H_4(NO_2)CONHC_2H_5$ Me, [205°]
Nitro mesidide
 $C_6H_4(NO_2)CONHC_2H_5$ Me, [207°] Formed, together with the following, by nitrating the mesidide (Schack, *B* 10, 1711)

Di-nitro-mesidide
 $C_6H_3(NO_2)_2CONHC_2H_5$ Me, [307°] Needles
Nitrile $C_6H_4(NO_2)CN$ [117°] Formed by nitrating benzonitrile, or by dehydrating *m*-nitro benzoic amide (Beilstein a Kuhlberg, *A* 146, 336, Engler, *Z* [2] 4, 613, *A* 149, 297, Fricke, *B* 7, 1321) Formed also by the action of hot cuprous potassium cyanide solution upon *m* nitro-diazo benzene chloride (Sandmeyer, *B* 13, 1494) Prepared by adding benzonitrile (10 mols) to KNO_3 (11 mols) dissolved in H_2SO_4 below 25° (Schöpf, *B* 18, 1063) Needles (from water), sl sol water, v sol alcohol and ether

p Nitro benzoic acid $C_6H_4(NO_2)CO_2H$ [14] [238°] *S* 075 at 16° Formed by the action of fuming HNO_3 on toluene (Glenard a Boudault, *A* 48, 344, G Fischer, *A* 127, 137, 130, 128, Beilstein a Wilbrand, *A* 126, 255, 128, 257), and by oxidising *p* nitro toluene with chromic acid (Beilstein a Geitner, *A* 139, 335, Korner, *Z* [2] 5, 636, Rosenstiel, *Z* [2] 5, 701) Produced also by the oxidation of *p* nitro cinnamic acid, and, in small quantity, by the nitration of benzoic acid (Griess, *B* 8, 528, Ladenburg, *B* 8, 536) Obtained also by the action of a hot cuprous potassium cyanide solution upon *p* nitro-diazo-benzene chloride, and saponification of the crude nitrile thus got (Sandmeyer, *B* 13, 1492) Prepared by oxidising *p* nitro toluene (50 g) with CrO_3 (250 g) and H_2SO_4 (110 g) diluted with water (460 g) (Schlosser a Skraup, *M* 2, 519, cf Michael a Norton, *B* 10, 580)

Properties—Yellowish laminae (from water) or needles (by sublimation), v sol alcohol and ether Less soluble in water than the *m* and *o*-isomerides Reduced by tin and HCl to *p*-amido-benzoic acid, and by sodium amalgam to *p*-azo-benzoic acid

Salts— NH_4A' 2aq efflorescent laminae— NaA' 3aq triclinic crystals (Bilfinger, *A* 135, 154)— KA' 2aq *S* 83 in the cold, 200 at 100° (Sokoloff, *J* 1864, 348)— BaA' 5aq monoclinic crystals (Bücking a Haushofer, *A* 193, 212) *S* 4 in the cold, 125 at 100°— BaA' (OBz) (Salkowsky, *B* 9, 24)— CaA' 8aq efflorescent tables *S* 3 in the cold, 83 at 100°— CaA' 9aq— CaA' (OBz) 3aq (Salkowski, *B* 10, 1259)—

SrA' (OBz) aq— ZnA' 2aq *S* (of ZnA') 7 at 17° (Mills, *C J* 19, 363), 125 at 100°— PbA' , *Methyl ether* MeA' [96°]

Ethylether EtA' [57°]
Chloride $C_6H_4(NO_2)COCl$ [75°] (c. 204° at 105 mm) Slender needles (from ligroin) (Gevekoht, *A* 221, 335)

Amide $C_6H_4(NO_2)CONH_2$ [198°] Needles (Beilstein a Reichenbach, *A* 132, 143) Treatment with bromine and KOH aq converts it into *p*-nitro-aniline

Anilide $C_6H_4(NO_2)CONHPh$ [204°] (Leo, *K* 3, 552)

Nitrile $C_6H_4(NO_2)CN$ [147°] Formed from the amide and P_2O_5 (Engler, *A* 149, 298, Fricke, *B* 7, 1321), or by the action of hot cuprous potassium cyanide solution on *p* nitro diazo benzene chloride (Sandmeyer, *B* 13, 1492) Laminae (from alcohol), v sol hot alcohol

Fourth and fifth nitro-benzoic acids have been described by Fitcka (*B* 8, 252, 710, 741, 9, 788, 10, 431, *J pr* [2] 17, 183), but their existence has been contested by other chemists (Leo Liebermann, *B* 10, 1033, Widmann, *B* 10, 1159, Claus, *B* 13, 891)

(4,3,1) Di-nitro-benzoic acid $C_6H_3(NO_2)_2CO_2H$ [161°] *S* 675 at 25° Prepared by heating *p* nitro benzoic acid with nitric and sulphuric acids in sealed tubes, and separated from the (4,2,1) isomeride, simultaneously formed, by the greater solubility of the latter in water (Claus a Halberstadt, *B* 13, 915) Colourless crystals Sol alcohol, ether, and hot water, sl sol cold water Very bitter taste Sublimes undecomposed

Salts— $A'Ba$ 4aq white crystals— $A'Ca$ 3aq plates The potassium, sodium, and ammonium salts are easily soluble

Di-nitro-benzoic acid $C_6H_3(NO_2)_2(CO_2H)$ [521] [177°] Formed, together with the (4,2,1) and (6,2,1) isomerides by boiling *o*-nitro-benzoic acid with a mixture of fuming HNO_3 (1 pt) and H_2SO_4 (1 pt) for 15 minutes (Griess, *B* 7, 1223) Needles or prisms, sl sol cold water, m sol hot water The di amido benzoic acid, obtained by reduction, yields *p* phenylene diamine on distillation— BaA' 4aq six sided plates, sl sol hot water

Di nitro-benzoic acid $C_6H_3(NO_2)_2CO_2H$ [421] [179°] *S* 1849 at 25° Formed in the preparation of each of the two preceding acids (Griess, *B* 7, 1225, Claus a Halberstadt, *B* 13, 816, Hubner a Stromeyer, *B* 13, 461, *A* 222, 79) Obtained also by heating di-nitro toluene with fuming HNO_3 at 100° for a fortnight (Tiemann a Judson, *B* 3, 223) White needles or tables May be sublimed Tastes bitter On reduction with tin and HCl aq it at once yields *m* phenylenediamine— BaA' 3aq m sol cold water— CaA' 2aq— MgA' 9aq

Di-nitro-benzoic acid $C_6H_3(NO_2)_2CO_2H$ [621] [202°] Formed, together with styphnic acid $C_6H_3(NO_2)_3(OH)$, and the (4,2,1)- and (5,2,1)-di-nitro benzoic acids, by heating *o*-nitro-benzoic acid with HNO_3 and H_2SO_4 (Griess) Felted needles (from boiling water), v sol boiling water Tastes intensely bitter Split up on distillation into CO_2 and *m*-di-nitro-benzene On treatment with tin and HCl aq it yields *m*-phenylenediamine— BaA' 2aq v e sol cold water.

Di-nitro-benzoic acid $C_6H_3(NO_2)_2CO_2H$ [5 8 1] [204°]. S 19 at 100°. Formed by nitration of benzoic acid or of *m*-nitro benzoic acid with a mixture of HNO_3 and H_2SO_4 (Cahours, *A Ch* [3] 25, 30, Voit, *A* 99, 100, Tiemann a Judson, *B* 3, 223, Muretoff, *Z* [2] 6, 641, Michler, *A* 175, 152). Obtained also by heating di-nitro-toluene [93°] with fuming HNO_3 at 150°, or by oxidising it with chromic acid mixture (Staedel, *B* 14, 902, *A* 217, 194, Hubner, *A* 222, 73), and in small quantity by the oxidation of β' -di-nitro naphthalene with dilute HNO_3 (S G 1 15) at 150° (Beilstein a Kurbatov, *B* 13, 355).

Preparation—1 By heating benzoic acid (20 g) with H_2SO_4 (180 g) and fuming HNO_3 (50 g) for 4 hours nearly to boiling (Hübner) — 2 By heating *m*-nitro benzoic acid (100 g) with fuming nitric acid (500 g) and H_2SO_4 (600 g) for 12 hours (H).

Properties—Thin tables (from dilute HNO_3) or monoclinic crystals (Henniges, *J* 1882, 902), v sl sol cold water, sl sol dilute HNO_3 , v sol alcohol and HOAc. Reduced by tin and HCl to di-amido-benzoic acid, which yields *m* phenylenediamine on distillation with baryta.

Salts— NaA' — KA' — BaA' aq (Hubner) m sol hot water— BaA' , 5aq (Muretoff)— CaA' , aq— MgA' , 8aq— MnA' , 2aq— PbA' , aq— AgA' , needles (from hot water).

Ethylether EtA' [94°] (H), [91°] (B a K) S (90 p.c. alcohol) 562 at 13°.

Amide $C_6H_3(NO_2)_2CONH_2$ [183°] (Voit, *A* 99, 105), [177°] (M). Plates (from water).

Tri-nitro-benzoic acid $C_6H_2(NO_2)_3CO_2H$ [190°]. Obtained by heating tri nitro toluene with fuming HNO_3 for a fortnight at 100° (T a J). Trimetric crystals, $abc = 887.1$ 572 (Friedlander, *Z K* 1, 623). May be sublimed— AgA' plates, sl sol water.

References—Bromo, Chloro, and Iodo, NITROBENZOIC ACIDS.

***o* NITRO-BENZOIC ALDEHYDE**

$C_6H_4(NO_2)CHO$ Mol w 151 [44°] (G a M), [46°] (F a H).

Formation—1 In small quantity, together with the *m*-isomeride, by adding benzoic aldehyde to a mixture of HNO_3 and H_2SO_4 (Rudolph, *B* 13, 810, cf Fittica, *B* 10, 1630)—2 By treatment of its oxim with chromic acid mixture (Gabriel a Meyer, *B* 14, 829)—3 By oxidising *o*-nitro-cinnamic acid with dilute $KMnO_4$ (Friedlander a Henriques, *B* 14, 2801)—4 By adding $NaNO_2$ to a cold solution of *o*-nitro-cinnamic ether in fuming nitric acid (F a H)—5 In small quantity by the action of water on the compound of *o*-nitro-toluene with CrO_2Cl_2 (Richter, *B* 19, 1062).

Preparation—50 grms of crude *o*-nitro-cinnamic acid are suspended in $2\frac{1}{2}$ litres of water, neutralised with Na_2CO_3 and filtered. The clear solution is put into a large stoppered bottle, 1 litre of benzene added, and kept cold during the reaction by the addition of ice 1225 c.c. of a 6 p.c. solution of $KMnO_4$ is added by degrees, shaking continuously, in order that the nitro-benzaldehyde, as it is formed, may be removed by the benzene from the action of the oxidising agent. The emulsion which is produced is now treated with a warm solution of 150 grms of sodium sulphite and HCl added, which dissolves the MnO_2 . The benzene layer,

which contains the whole of the nitrobenzaldehyde is removed, and the latter obtained by distilling off the benzene. The yield is 50 p.c. to 60 p.c. of the theoretical (Einhorn, *B* 17, 119).

Properties—Long yellow needles (from water), sl sol water, v sol alcohol and ether. May be distilled. Volatile with steam. With $NaHSO_3$ it forms a compound crystallising in plates.

Reactions—1 On reduction with tin and HOAc it yields anthranil $C_6H_3\left(\begin{smallmatrix} CO \\ NH \end{smallmatrix}\right)$, the lactam of *o*-amido benzoic acid (Friedlander a Henriques, *B* 15, 2105, cf Rudolph, *B* 13, 310). Reduced by $FeSO_4$ and NH_3 to amido benzoic aldehyde (Friedlander, *B* 15, 2572, 17, 456)—2 Yields *o*-nitro benzoic acid on oxidation with dilute aqueous $KMnO_4$. It also undergoes this oxidation when administered to dogs (Sieber a Smirnoff, *M* 8, 88)—3 Conc $NaOHAq$ converts it into *o*-nitro benzoic acid and *o*-nitro-benzyl alcohol—4 $NaOAc$ and Ac_2O yield *o*-nitro cinnamic acid by Perkin's reaction—5 On warming with a dilute alcoholic solution of urea a compound $C_6H_3(NO_2)CH(NHCO NH_2)_2$ [200°] is formed (Ludy, *M* 10, 295). When a few drops of H_2SO_4 are added to an alcoholic solution of *o*-nitro benzoic aldehyde and urea there is formed a different compound $C_{10}H_8N_2O_2$ [170°]—6 With di-methyl-aniline it forms nitro tetra-methyl di-amido tri phenyl-methane $C_6H_3(NO_2)CH(C_6H_4NMe_2)_2$ —7 Aceto-acetic ether (2 mols) and NH_3 form *o*-nitro phenyl di-methyl pyridine dihydride dicarboxylic ether [120°] and a compound $C_{15}H_{12}N_2O_5$ [189°], forming a salt $B'HCl$, converted by nitrous acid into an indifferent isomeride [192°] (Lepetit, *B* 20, 1838)—8 Boiling conc aqueous $KClO_4$ forms *o*-azoxy benzoic acid—9 Aldehyde and a little baryta water forms $C_6H_3NO_4$ [120°], converted by further treatment with baryta water into $C_6H_3(NO_2)CH(OH)CH_2CH(OH)$ [109°], and by $NaOHAq$ into indigo (Baeyer a Drewson, *B* 15, 2861)—10 Acetone and $NaOH$ forms *o*-nitro-styryl methyl ketone, which on warming with more alkali yields indigo.

Oxim $C_6H_3(NO_2)CHNOH$ *o*-Nitro-*o*-nitroso toluene [96°]. Formed by the action of hydroxylamine on the aldehyde in alcoholic solution, or by treating (2,4,1) nitro-amidophenyl acetic acid with nitrous acid (Gabriel a Meyer, *B* 14, 826, 15, 3057, 16, 520). Slender needles (from hot water), v sol alcohol and ether. Tastes sweet. Dissolves in alkalis. Conc $HClAq$ at 160° decomposes it into *o*-nitro-benzoic acid and NH_3 . Boiling with Ac_2O and $NaOAc$ forms the nitrile of *o*-nitro benzoic acid.

Methyl derivative of the oxim
 $C_6H_3(NO_2)CHNOMe$ [58°].

Phenyl hydrazide $C_6H_3(NO_2)CHN_2HPH$ [158°] (F), [148°] (L). Formed by adding phenyl-hydrazine to an alcoholic solution of the base (Pickel, *A* 282, 232, Ludy, *M* 10, 314). Scarlet needles, sl sol hot water, forming a dark blue solution in conc H_2SO_4 .

***m*-Nitro-benzoic aldehyde** $C_6H_4(NO_2)CHO$ [8 1] [58°]. Obtained by dissolving benzoic aldehyde (1 vol) in a mixture of fuming HNO_3 (1 vol) and H_2SO_4 (10 vols) in the cold (Bertagnini, *A* 79, 259, 86, 190; Lippmann a Hawkeczek, *B* 9, 146, Friedlander a Henriques,

B 14, 2802, Ehrlich, B 15, 2010) Needles (from water), m sol hot water, v sol alcohol

Reactions—1 *Chromic acid* oxidises it to *m* nitro benzoic acid—2 *Tin* and *HOAc* reduce it to *m*-amido-benzoic aldehyde—3 *Ammonia* forms $(C_6H_4(NO_2)CH_2)_N$, which on heating with aqueous *KOH* forms tri-nitro-amarin $C_{21}H_{14}N_4O_8$ —4 *Ammonia* and *acetoacetic ether* form $C_6H_4(NO_2)C_2NH_2Me (CO_2Et)_2$ [65°] (Lepetit, B 20, 1838, 2397)—5 *Sodium succinate* and acetic anhydride form nitro phenyl paraconic acid $C_6H_4(NO_2)CHCH(CO_2H)CH_2COO$ (Salomon-

son, R T C 6, 1)—6 When its benzene solution is shaken with H_2SO_4 , there is formed $C_6H_4(NO_2)CHPh$ [90°], while toluene, by similar condensation, yields $C_6H_4(NO_2)CH(C_2H_5)_2$ [85°] (Tschacher, B 21, 188)—7 *Phenyl acetonitrile* and alcoholic *NaOEt* yields the nitrile of nitro-*a* phenyl cinnamic acid (Frost, A 250, 156)—8 After administration to a dog it is excreted as *m* nitro hippuric acid (Sieber, M 8, 88)

Combinations— $(C_6H_4NO_2)NH_2SO_3H$ 4aq colourless prisms— $(C_6H_4NO_2)NaSO_3H$ 8aq yellowish crystalline scales (from hot water)— $(C_6H_4NO_2)NPhH_2SO_3H$ needles (Schiff, A 195, 301)— $(C_6H_4NO_2)PH_3$ powder, insol alcohol Formed from the aldehyde, PH_3 , and *HCl* (Messinger a Engels, B 21, 333)

Oxim $C_6H_4(NO_2)CHNOH$ [119°] Needles (from water) (Gabriel, B 15, 3060) *PCl*, yields *m* nitro benzomitrile (Gabriel, B 16, 520)

Iso-oxim [118°] Formed by passing *HCl* into an ethereal solution of the oxim, and decomposing the hydrochloride by *NaCO*. Needles (from ether) Combines with phenyl cyanate in ethereal solution with formation of $C_6H_4(NO_2)CHNOCONPhH$ [75°], which readily changes into an isomeride [105°] By treatment with *NaOMe* and *MeI* it is converted into a mixture of two isomeric methyl derivatives $C_6H_4(NO_2)CHNOMe$ melting at 69° and at 117° (Goldschmidt a Ernst, B 23, 2170) *MeI* acting on its silver salt gives the methyl ether, melting at 69° *NaOEt* and benzyl chloride yield a benzyl ether [148°] Phenyl cyanate in ethereal solution yields $C_6H_4(NO_2)CHNOCONHPh$ [105°], crystallising in yellow needles

Phenyl-hydrazide

$C_6H_4(NO_2)CHN_2HPh$ [121°] (Pickel, A 232, 232, Ludy, M 10, 316) Orange tables Yields an acetyl derivative $C_6H_4(NO_2)CHN_2Ac$ [170°] (Schroeder, B 17, 2097) Nitrous acid converts it into $(C_6H_4(NO_2)CHN_2NPh)NOH$, which crystallises in needles, forming a deep blue pp in H_2SO_4

p Nitro-benzoic aldehyde

$C_6H_4(NO_2)CHO$ [41] [106°]

Formation—1 By boiling *p*-nitro-benzyl chloride (10 pts) with $Pb(NO_2)_2$ (14 pts), water (60 pts), and HNO_3 (8 G 13) (Fischer a Greiff, B 13, 670)—2 By oxidising *p*-nitro-cinnamic ether (Friedlander a Henriques, B 14, 2803) or *ap* di nitro cinnamic ether (Friedlander a Maly, A 229, 212, cf Baeyer, B 14, 2317)

Preparation—1 By adding *KNO*₃ to a cooled solution of *p* nitro cinnamic acid, or its methyl or ethyl ether, in strong H_2SO_4 , the yield is 60 p.c to 95 p.c (Basler, B 16, 2714)—2 45 pts of chromyl chloride (CrO_2Cl_2) are slowly added to 20 pts. *p* nitro-toluene, dissolved in 80 to 100 pts of CS_2 . After standing for two days,

the pp $(C_6H_4(NO_2)CH_2)_2Cr_2O_2Cl_2$ is separated, washed with CS_2 , and treated with water The CS_2 is driven off with steam, and the product recrystallised from water, yield, 60 p.c to 70 p.c. of the nitro toluene (Reichter, B 19, 1060)

Properties—Long colourless prisms (from hot water), sl sol water and ether, m sol alcohol May be sublimed Volatile with steam. Forms with $NaHSO_4$ a compound crystallising in plates

Reactions—1 *Chromic acid mixture* oxidises it to *p* nitro benzoic acid.—2 *Aniline* forms $C_6H_4(NO_2)CHNPh$ [93°]—3 *Dimethylaniline* condenses with it to *p* nitro tetra-methyl-di-*p*-amido-tri-phenyl-methane—4 When administered to dogs it is excreted as *p*-nitro-hippuric acid.—5 With *p* toluidine and conc *HClAq* it forms $C_6H_4(NO_2)CH(C_2H_5MeNH_2)_2$ [172°] (Bischler, B 20, 3302)—6 *Ammonia* and *acetoacetic ether* form the compound $C_6H_4(NO_2)C_2NH_2Me(CO_2Et)_2$ [118°-122°]—7 *Indoxyzyl* with *HClAq* and *HOAc* forms the indogemide $C_6H_4<\underset{NH}{C}>CCHC_6H_4NO_2$, crystallising in red needles [273°]

Oxim $C_6H_4(NO_2)CHNOH$ [129°] Reddish-yellow scales (Gabriel a Herzberg, B 16, 2000)

Iso-oxim [175°] (Behrend, A 263, 849)

Benzyl ether of the iso-oxim

$C_6H_4(NO_2)CH<\underset{O}{N(C_6H_5)}>$ [118°] Formed by oxidising (8) benzyl *p* nitro benzyl hydroxylamine with K_2FeCy_6 (Behrend a König, B 23, 2750) Needles (from alcohol), split up by hot *HClAq* into *p* nitro benzoic aldehyde and (8)-benzyl-hydroxylamine

Phenyl hydrazide $C_6H_4(NO_2)CHN_2HPh$ [125°] Red needles (Pickel, A 232, 232). Forms a red solution in conc H_2SO_4 (Ludy, M 10, 315)

NITROBENZOIC ANHYDRIDE v Anhydride of Nitrobenzoic acid

m NITRO-BENZOIC IMIDO-ETHYL ETHER $C_6H_4NO_2$, i.e. $C_6H_4(NO_2)C(NH)OEt$ Formed by treatment of $C_6H_4(NO_2)CONHAg$ with *EtI*, followed by ethereal *HCl* (Tafel a Enoch, B 23, 1550) Yellow oil—*BHCl* White mass, v sol water and alcohol, converted by hot water into *m*-nitro benzoic ether—*BHCl*, [132°]

NITRO-BENZONITRILE v Nitrile of Nitro benzoic acid

o NITRO-BENZOPHENONE $C_{12}H_8NO_2$, i.e. $C_6H_5CO C_6H_4(NO_2)$ *o*-Nitro di phenyl ketone [105°] Formed by oxidation of *o* nitro-di-phenyl methane with CrO_3 in acetic acid (Geigy a Koenigs, B 18, 2403) Colourless crystals

m Nitro-benzophenone $C_6H_5CO C_6H_4NO_2$ [95°] Formed by the action of *m* nitro-benzoyl chloride upon benzene in presence of $AlCl_3$ (Geigy a Koenigs, B 18, 2401, cf Becker, B 15, 2090) Yellowish needles

p-Nitro-benzophenone $C_6H_5CO C_6H_4$ [138°] Formed by oxidation of *p* nitro-di-phenyl-methane with CrO_3 in acetic acid (Basler, B 16, 2717) Small white plates or needles. Sol benzene, hot alcohol, and hot water, sl sol. cold water, cold alcohol, ligroin, and CS_2 .

(a) Di-nitro-benzophenone $(C_6H_4(NO_2))_2CO$ [190°] Formed, together with a larger quantity of the (8)-isomeride, by heating benzophenone with fuming HNO_3 at 60° (Prätorius, B 10,

1855, 11, 744, Stadel, A 194, 349, 218, 344) Formed also by oxidation of the corresponding di-nitro di-phenyl-methane [183°] by CrO_3 in HOAc (Stadel & Sauer, B 11, 1747) Long needles (from HOAc) Yields on reduction di-amido-benzophenone [172°]

(β)-Di-nitro-benzophenone $\text{C}_{12}\text{H}_8\text{N}_2\text{O}_5$ [149°] Formed as above Laminæ (from benzene or HOAc) Yields on reduction di-amido-benzophenone [165°]

Oasm ($\text{C}_6\text{H}_5\text{NO}_2$), CNOH [207°], small yellow needles (from hot alcohol).

Phenyl hydrazide

($\text{C}_6\text{H}_5\text{NO}_2$), $\text{C}_6\text{H}_5\text{N}_2\text{H}_3$, [220°], red powder, sol. acetic acid, sl sol. alcohol, insol water (Münchmeyer, B 20, 510)

(γ)-Di-nitro-benzophenone $\text{C}_{12}\text{H}_8\text{N}_2\text{O}_5$ [190°] Formed by oxidation of the di nitro di phenyl-methane [118°] Crystals Yields on reduction di-amido-benzophenone [181°].

Tetra-nitro-benzophenone $\text{C}_{12}\text{H}_4(\text{NO}_2)_4\text{O}$ [235°] Formed by oxidising tetra-nitro di-phenyl-methane (Stadel, A 218, 341) Needles (from HOAc).

p-NITRO-BENZOYL-ACETIC ACID $\text{C}_8\text{H}_7\text{NO}_5$, *s.* $\text{C}_6\text{H}_4(\text{NO}_2)\text{COCH}_2\text{CO}_2\text{H}$ Nitro phenyl methyl ketone carboxylic acid [135°] Formed by digesting *p* nitro phenyl propionic ether with H_2SO_4 (85 p c) at 35° till it is completely soluble in aqueous NaOH (Perkin & Bellenot, B 17, 386, 18, 952, C J 49, 444) Needles, v sol alcohol and ether, sol water FeCl_3 colours its alcoholic or hot aqueous solution reddish brown On heating by itself or with dilute acids or alkalis it splits up into CO_2 and *p* nitro-acetophenone

Methyl ether MeA' [107°] Monoclinic crystals, v sol hot alcohol NaOMe forms yellow crystals of $\text{C}_6\text{H}_4(\text{NO}_2)\text{COCHNMeCO}_2\text{Me}$, a body from which benzyl chloride produces $\text{C}_6\text{H}_4(\text{NO}_2)\text{COCH}(\text{C}_6\text{H}_5)\text{CO}_2\text{Me}$ [57°]

Ethyl ether EtA' [76°] Monoclinic crystals, $a, b, c = 358, 11, 238$, $\beta = 72^\circ 22'$ Its alcoholic solution is coloured brownish violet by FeCl_3 Nitrous acid acting on its ethereal solution forms $\text{C}_6\text{H}_4(\text{NO}_2)\text{COC}(\text{NOH})\text{CO}_2\text{Et}$ [220°] Yields $\text{C}_6\text{H}_4(\text{NO}_2)\text{COCHNMeCO}_2\text{Et}$, crystallising in orange needles, whence aqueous AgNO_3 forms an explosive amorphous Ag salt The sodium salt is converted by ethyl iodide into the ether $\text{C}_6\text{H}_4(\text{NO}_2)\text{COCHEtCO}_2\text{Et}$ [40°]

NITRO-BENZOYL-ACETOACETIC ETHER *c.* ACETOACETIC ETHER

o-NITRO-BENZOYL-ACETONE

$\text{C}_{10}\text{H}_7\text{NO}_5$, *s.* $\text{NO}_2\text{C}_6\text{H}_4\text{COCH}_2\text{COCH}_3$ [55°]. Got by boiling *o*-nitro benzoyl aceto acetic ether with H_2SO_4 (1 pt) and water (2 pts) for 4 hours (Gevekoht, A 221, 332) Crystallised from benzoline Insol. water, v sol alcohol and ether Gives with phenyl hydrazine a compound melting at 120°, crystallising from alcohol in slender needles, possibly nitro di phenyl methyl-pyrazole (Fischer & Bulow, B 18, 2136)

p-NITRO-BENZOYL-ANGELIC ETHER

$\text{C}_{12}\text{H}_9\text{NO}_5$, $\text{COCH}(\text{C}_6\text{H}_5)\text{CO}_2\text{Et}$ [46°] Formed by the action of allyl iodide upon the sodium-derivative of *p*-nitro-benzoyl-acetic ether (Perkin & Bellenot, B 18, 957) Colourless plates

(α)-DI-NITRO-BENZOYL-BENZOIC ACID $\text{C}_{14}\text{H}_8\text{N}_2\text{O}_7$, [240°] Formed from benzoyl

benzoic acid, HNO_3 , and H_2SO_4 (Plaskuda, B 7, 988). Plates — BA' , aq — CA' , 2aq

(β)-Di-nitro-benzoyl-benzoic acid

$\text{C}_{12}\text{H}_7(\text{NO}_2)_2\text{CO}_2\text{C}_6\text{H}_4(\text{NO}_2)\text{CO}_2\text{H}$ [212°] Formed by boiling di-nitro phenyl-*p* tolyl ketone with CrO_3 and HOAc (Plaskuda & Zinke, B 7, 984). Plates (from hot water)

o-NITRO-BENZOYL-BENZYL-MALONIC ETHER

$\text{C}_8\text{H}_7(\text{NO}_2)\text{OCH}(\text{C}_6\text{H}_5)(\text{CO}_2\text{Et})_2$ [94°] Formed from sodium benzyl malonic ether and *o*-nitro benzoyl chloride, or from benzyl chloride and sodium *o* nitro benzoyl malonic ether (Bischoff & Siebert, A 239, 103) Prisms (from dilute alcohol) V sol ether and hot alcohol, sl sol ligroin Gives no colour with FeCl_3 , Alcoholic KOH produces benzyl malonic acid and *o* nitro benzoic acid Ammonia forms *o*-nitro-benzamide and benzyl-malonic ether

NITRO-BENZOYL-CHLORIDE *v* Chloride of

NITRO-BENZOIC ACID

NITRO-BENZOYL CYANIDE *v* Cyanide of

NITRO-BENZOIC ACID

NITRO-BENZOYL-FORMIC ACID *v* Nitro-

PHENYL GLYOXYLIC ACID

o-NITRO-BENZOYL MALONIC ETHER

$\text{C}_8\text{H}_7(\text{NO}_2)\text{COCH}(\text{CO}_2\text{Et})_2$ [54°] The sodium-derivative $\text{C}_8\text{H}_6(\text{NO}_2)\text{COONa}(\text{CO}_2\text{Et})_2$ is formed by the action of NaOEt on di-nitro di benzoyl-malonic ether, or of *o* nitro benzoyl chloride (1 mol) on di sodio malonic ether (1 mol) (Bischoff, B 16, 1044, 17, 2791, 22, 387) Needles or prisms (from alcohol) FeCl_3 colours its alcoholic solution dark red Bromine acting on its sodium derivative forms the bromo derivative $\text{C}_8\text{H}_6(\text{NO}_2)\text{COCB}(\text{CO}_2\text{Et})_2$ [72°] On reduction by zinc-dust and HCl it yields (*Py* 1,3) di oxyquinoline (*Py* 2) carboxylic acid, and other bodies

o-Di-nitro-di-benzoyl-malonic ether

($\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CO}$), $\text{C}(\text{CO}_2\text{Et})_2$ [93°] Colourless tables or prisms Obtained by the action of *o*-nitro benzoyl chloride (2 mols) on di sodio malonic ether (1 mol) in alcoholic solution By treatment with sodium ethylate one of the nitro benzoyl groups is removed with formation of sodio nitro benzoyl malonic ether and nitro benzoic ether By alcoholic NH_3 both benzoyl groups are removed with production of malonic ether and nitro benzamide (2 mols) (Bischoff & Raab, B 17, 2789)

m-NITRO BENZOYL PEROXIDE

($\text{C}_6\text{H}_4(\text{NO}_2)\text{CO}$), O_2 , Oil, ppd by adding water to a solution of benzoyl peroxide in fuming HNO_3 (Brodie, *Pr* 12, 655)

DI - *p* - NITRO - DI - BENZOYL - SUCCINIC ETHER

$\text{C}_{12}\text{H}_9\text{NO}_5$, $\text{COCH}(\text{CO}_2\text{Et})\text{CH}(\text{CO}_2\text{Et})\text{CO}_2\text{C}_6\text{H}_4(\text{NO}_2)$ [180°] Formed by the action of iodine on an ethereal solution of the sodium derivative of *p* nitro-benzoyl acetic ether (Perkin & Bellenot, C J 49, 452, B 18, 953) Needles Forms an amorphous di-sodium derivative

o NITRO-BENZYL ALCOHOL

$\text{C}_8\text{H}_7\text{NO}_3$, CH_2OH [74°] Prepared by the action of aqueous NaOH on *o* nitrobenzoic aldehyde (Friedlander & Henriques, B 14, 2804, Gabriel & Borgmann, B 16, 2065, Geigy & Koenigs, B 18, 2403) Yellow needles, sl sol water Yields on reduction by zinc and HCl *o*-amido-benzyl alcohol [82°] Chromic acid oxidises it to *o*-nitro benzoic acid.

NITRO-BENZOPHENONE.

***m*-Nitro-benzyl alcohol** $C_6H_4(NO_2)CH_2OH$ (175°-180° at 8 mm) Formed by heating *m*-nitro-benzaldehyde with alcoholic potash, and ppg by water (Grimaux, *B* [2] 8, 433)

Preparation—2 pts of *m*-nitro benzaldehyde are mixed with a cooled solution of 1 pt of KOH in 6 pts of water. After standing overnight the product is extracted with ether, the yield is nearly the theoretical (Becker, *B* 15, 2091)

Oil, decomposed by distillation under atmospheric pressure

***p*-Nitro-benzyl alcohol** $C_6H_4(NO_2)CH_2OH$ [92°] Formed by heating its acetyl derivative with aqueous NH_3 at 100° (Beilstein a Kuhlberg, *Z* [2] 3, 467, *A* 147, 943) Formed also, together with di nitro *p*-toluidine, by warming the *p* nitro benzyl ether of di nitro-*o*-cresol with alcoholic NH_3 (Staedel, *A* 217, 183)

Preparation—1 Finely powdered *p*-nitro-benzaldehyde (1 pt) is added to 5 or 6 pts of 15 p aqueous NaOH, and allowed to stand for 12 hours, it is then diluted with a little water and extracted with ether, the yield is 80 p c to 90 p c of the theoretical—2 20 pts of *p* nitro benzyl acetate (obtained by nitration of benzyl acetate) in 40 pts of boiling alcohol is treated with 35 pts of 15 p aqueous NaOH, quickly cooled and poured into 200 pts of iced water, and the pp collected, the yield is about 75 p c (Basler, *B* 16, 2715)

Properties—Yellowish needles Yields *p* nitro benzoic acid on oxidation Concentrated nitric acid converts it into the nitrate $C_6H_4(NO_2)CH_2ONO_2$ [71°] (Staedel, *B* 14, 90)

Acetyl derivative $C_6H_4(NO_2)CH_2OAc$ [78°] Formed by dropping benzyl acetate into fuming HNO_3 Pale yellow needles

***o*-NITRO-BENZYLAMINE** $C_6H_4NO_2$ *te* $C_6H_4(NO_2)CH_2NH_2$ Formed by heating *o*-nitro benzyl phthalimide with $HClAq$ at 200° (Gabriel, *B* 20, 2228) Liquid, v sol water— $B'HCl$ Needles, v sol water— $B'H.PtCl_2$ 2aq

Formyl derivative $C_6H_4AcNO_2$ [90°] Formed by heating the hydrochloride with dry sodium formate and formic acid (Gabriel a Jansen, *B* 23, 2813) Crystals Yields quinoxaline dihydride on reduction by zinc and HCl

Acetyl derivative [99°] Needles Yields on reduction $C_6H_4(NH_2)CH_2NHAc$ [113 5°]

Benzoyl derivative [112°] Needles (from hot alcohol) Yields on reduction $C_6H_4(NH_2)CH_2NHBz$ [109°]

***m*-Nitro-benzyl-amine** $C_6H_4(NO_2)CH_2NH_2$ Formed by heating *m* nitro benzyl phthalimide with $HClAq$ (Gabriel a Hendess, *B* 20, 2869) Formed also from *m* nitro benzyl chloride and alcoholic NH_3 (Borgmann, *C* 1885, 456) Needles— $B'H.PtCl_2$ plates

Acetyl derivative $C_6H_4(NO_2)CH_2NHAc$ [91°] Needles, sol hot water

***p*-Nitro-benzyl amine** $C_6H_4(NO_2)CH_2NH_2$ Prepared from *p*-nitro benzyl chloride by treatment with potassium phthalimide and decomposition of the *p*-nitro-benzyl phthalimide (Hafner, *B* 23, 387) Strongly alkaline oil, absorbing CO_2 from the air Converted by nitrous acid into *p* nitro benzyl alcohol OS_2 forms $C_6H_4(NO_2)CH_2NHCSNH_2$, $CH_2.C_6H_4.NO_2$ [198°].— $B'HCl$ plates, m sol. water— $B'H.PtCl_2$.— $BHNO_2$, needles, sl sol water

Acetyl derivative $C_6H_4(NO_2)CH_2NHAc$ [138°] Needles, sol water Formed by heating the base with Ac_2O and $NaOAc$, or by nitration of the acetyl derivative of benzylamine (Amsel a Hofmann, *B* 19, 1286)

Benzoyl derivative [156°] Needles **Di-*p*-nitro-di-benzyl-amine** $C_{11}H_{12}N_2O_4$ *te* $(C_6H_4(NO_2)CH_2)_2NH$ [93°] Obtained, together with the following isomeride and tri nitro-tri-benzylamine, by heating nitro benzyl chloride with aqueous NH_3 at 100° (Strakosch, *B* 6, 1056) Yellow plates (from alcohol)— $B'HCl$ [212°].— $B'H.PtCl_2$

Di nitro-di-benzyl-amine [above 100°] Formed as above— $B'HCl$ [173°]

Tri-*o* nitro-tri-benzyl-amine $(C_6H_4(NO_2)CH_2)_3N$ [157°] Formed almost exclusively by heating *o* nitro-benzyl chloride with aqueous NH_3 Long yellowish needles. Sol benzene, sl sol alcohol Its salts are decomposed by water (Lellmann a Stuckel, *B* 19, 1605)

Tri *p* nitro-tri benzyl-amine $(C_6H_4(NO_2)CH_2)_3N$ [163°] Formed by heating *p*-nitro benzyl chloride with aqueous NH_3 (Strakosch) Needles (from $HOAc$)

Tri nitro-benzyl-amine $N(CH_2.C_6H_4.NO_2)_3$ [159°] Formed by nitration of tri benzyl amine with a mixture of conc HNO_3 and conc H_2SO_4 Colourless crystals Sol hot acetic acid, insol alcohol and ether (Marquardt, *B* 19, 1030)

***o*-NITRO-BENZYL-ANILINE** $C_6H_4(NO_2)CH_2.NHC_6H_5$ Prepared by heating 2 mols of aniline with 1 mol of *o*-nitro benzyl chloride in alcoholic solution for 2 hours on the water bath It crystallises in two allotropic forms, of which the unstable form changes into the stable by fusion or by long keeping The unstable form crystallises in reddish-yellow monoclinic needles, $a b c = 8585 \ 111587$, $\beta = 87^\circ 42' 52''$, and melts at [44°] The stable form crystallises in glistening brown triclinic prisms, $a b c = 1 \ 6764 \ 1^\circ$, $\alpha = 117^\circ$, $\beta = 137^\circ 40' 6''$, $\gamma = 69^\circ 38' 44''$, it melts at [57°] V sol alcohol, ether, and benzene, sparingly in petroleum-ether Reduced by tin and HCl to $C_6H_4NO_2$ [83°]— $B'HCl$ 3aq white needles, basified by much water

Acetyl derivative $C_6H_4(NO_2)CH_2.NPhAc$ [75°] (Paal a Krecke, *B* 23, 2637)

Benzoyl derivative $C_6H_4(NO_2)CH_2.NPhBz$ [101°] Well formed crystals, sol alcohol, ether, and acetic acid, v sol chloroform, sl sol petroleum-ether By reduction with tin and HCl it is converted into $C_6H_4(NH_2)CH_2.NPhBz$ [115°] (Lellmann a Stuckel, *B* 19, 1605, 24, 718, Söderbaum a Widman, *B* 23, 2198)

Formyl derivative $C_6H_4(NO_2)CH_2.NC_6H_5.CHO$ [77°] Yellow monoclinic plates, $a b c = 548 \ 1 \ 1085$, $\beta = 69^\circ 7'$ Yields phenyl-quinoxaline dihydride on reduction (Paal a Busch, *B* 22, 2683)

***m*-Nitro benzyl-aniline** $C_6H_4(NO_2)CH_2.NHC_6H_5$ [86°] From *m*-nitro-benzyl chloride and aniline (Borgmann, *C* 1885, 456) Orange-red needles— $B'HCl$ plates.

***p*-Nitro-benzyl-aniline** $C_6H_4(NO_2)CH_2.NHC_6H_5$ [68°] Formed from *p*-nitro-benzyl chloride and aniline (Strakosch,

B 6, 1062 Golden yellow needles, sol hot alcohol—B'HCl Plates

Di-nitro-di-benzyl-aniline

$(C_6H_4(NO_2)CH_2)_2NC_6H_5$ [206°] Formed in small quantity as a by product of the action of *o*-nitro benzyl chloride upon aniline Greenish-yellow needles Sparingly soluble in acetic acid (Lellmann a Stuckel, B 19, 1608)

***m* Nitro benzyl-aniline** $C_6H_4(NO_2)NHCH_2C_6H_5$ [107°] From *m* diazo nitro benzene *m* nitro-benzyl anilide by heating with conc HCl at 100° (Meldola a Streatfield, C J 51, 114)

The Nitrosamine is an oil

***p*-Nitro-benzyl-aniline** $C_6H_4(NO_2)NHCH_2C_6H_5$

[148°] From *p*-diazo-nitro benzene *p*-nitro-benzyl-anilide by heating with conc HCl at 100° (Meldola a Streatfield, C J 51, 113, Meldola a Salmon, C J 53, 779) Golden scales

Nitrosamine $C_6H_4(NO_2)N(NO)C_6H_5$ [108°]

Acetyl derivative [109°]

Benzoyl derivative

$C_6H_4(NO_2)NBzCH_2Ph$ [194°] Needles

***p*-Nitro-di-benzyl-aniline**

$C_6H_4(NO_2)N(CH_2C_6H_5)_2$ [130°] Formed by nitration of di benzyl aniline dissolved in acetic acid Yellow needles Sol hot alcohol and acetic acid, sl sol cold alcohol, v sol ether and benzene (Matzudaira, B 20, 1613)

NITRO-BENZYL-BENZENE *v* **NITRO-DIPHENYL METHANE**

Di-*m* nitro di-benzyl-benzene $C_{12}H_{10}N_2O_4$ *z* $C_6H_4(CH_2C_6H_4NO_2)_2$ [165°] Formed by the action of H_2SO_4 on a mixture of *m* nitro diphenyl methane and *m*-nitro benzyl alcohol (Becker, B 15, 2091)

Di-*p* nitro di-benzyl-benzene

$C_6H_4(NO_2)CH_2C_6H_4CH_2C_6H_4(NO_2)$ [c 146°] Formed as a by product in the preparation of *p*-nitro-di phenyl methane by the action of H_2SO_4 on a mixture of *p* nitro benzyl alcohol and benzene (Basler, B 16, 2716) Small white concentric needles Sol benzene and hot acetic acid, v sl sol other solvents

***m*-NITRO-BENZYL BROMIDE** $C_6H_4BrNO_2$ *z* $C_6H_4(NO_2)CH_2Br$ [58°] Formed by heating *m*-nitro toluene with bromine at 130° (Wachendorff, A 185, 266)

***p*-Nitro-benzyl bromide** [100°] Formed from *p*-nitro toluene and Br (W) Needles

***p*-NITRO-BENZYL-CARBAMIC ETHER**

$C_6H_4(NO_2)CH_2NHCO_2Et$ [117°] Formed from *p*-nitro benzylamine and $ClCO_2Et$ (Hafner, B 23, 840) Silky needles, v sol ether

***o*-NITRO-BENZYL CHLORIDE** $C_6H_4ClNO_2$ *z* $C_6H_4(NO_2)CH_2Cl$ [49°] Formed, together with the *p*-isomeride, by nitration of benzyl chloride in the cold (Beilstein a Gaitner, A 139, 337, Abelli, G 13, 97, Nöbling, B 17, 385, Kumpf (A 224, 100) Formed also by treating *o*-nitro benzyl alcohol with PCl_5 in the cold (Gabriel a Borgmann, B 16, 2066, Geigy a Königs, B 18, 2401) Thick crystals Reduced by $SnCl_4$ to *o*-'benzylene imide' C_6H_5N , an amorphous base (Lellmann a Stuckel, B 19, 1611) Aniline yields only $C_6H_4(NO_2)CH_2NHPh$ reduced by zinc-dust in $HOAc$ to the amido-compound $C_6H_4(NH_2)CH_2NHPh$ [82°] (Söderbaum a Widman, B 23, 2198)

***m*-Nitro benzyl-chloride** $C_6H_4(NO_2)CH_2Cl$ [47°] (c. 178° at 85 mm). Long yellow needles

V sol alcohol, ether, and benzene Formed by the action of PCl_5 on the alcohol (Gabriel a Borgmann, B 16, 2064)

***p*-Nitro-benzyl chloride**

[4 1] $C_6H_4(NO_2)CH_2Cl$ [71°] Formed by treating benzyl chloride with HNO_3 (S G 1 48) at -10° (Elbs a Bauer, J pr [2] 34, 343, cf Beilstein a Gaitner, A 139, 337, Strakosch, B 6, 1056, Grimaux, Bl [2] 8, 433) Formed also by passing chlorine into *p* nitro toluene at 190° (Wachendorff, B 8, 1101, A 185, 271) Crystals

Reactions—1 Ammonium sulphide gives

p-nitro benzyl mercaptan (Strakosch, B 5, 697)

2 Alcoholic potash gives an azo derivative of di nitro stilbene—3 An alkaline solution of SnO forms *p* dinitro dibenzyl (W Roser, A 238, 363)—4 Chromic acid mixture oxidises it to

p nitro benzene acid—5 A solution of $SnCl_4$ in $HClAq$ reduces it to *p* 'benzylene imide' C_6H_5N , an amorphous base (L a S)

NITRO BENZYL CYANIDE *v* **Nitrile of**

NITRO PHENYL ACETIC ACID

***o* - NITRO - BENZYL - ETHYL - MALONIC**

ETHER $C_{12}H_{13}NO_6$ *z* $C_6H_4(NO_2)CH_2CEt(CO_2Et)_2$

Formed from ethyl malonic ether, $NaOEt$, and *o* nitro benzylchloride (Lellmann a Schleich, B 20, 440) Reduced by zinc and $HOAc$ to $C_6H_4NO_2$ [114°]

***p* Nitro benzyl ethyl-malonic ether** [52°] Formed in like manner Needles

***o* NITRO-BENZYL ETHYL OXIDE**

$C_6H_4(NO_2)OC_2H_5$ Formed by heating *o*-nitro benzyl chloride with alcohol in a boiling brine bath (Errera, G 18, 232) Oil

***m* Nitro-benzyl ethyl oxide** Prepared by heating *m* nitro benzyl chloride with alcoholic potash (E) Oil, solidifying in a freezing mixture

***p* Nitro benzyl ethyl oxide** [24°] Prepared by heating *p* nitro benzyl chloride with alcohol (E) Trimetric crystals

NITRO BENZYLIDENE ACETONE *v* **NITRO-ETHYL METHYL KETONE**

NITRO BENZYLIDENE ACETONAMINE *v* **ACETONAMINE**

***m* DI - NITRO - DI BENZYLIDENE - DI -**

AMIDO DIPHENYL

$C_6H_4(NO_2)CHNCH_2C_6H_4NCHC_6H_4(NO_2)$ Formed by heating *op* di amido diphenyl with *m*-nitro benzoic aldehyde and alcohol at 100° (Reuland, B 22, 3011) Yellow crystalline powder, m sol alcohol The isomeric compound from *p* nitro benzoic aldehyde and *op* di amido diphenyl melts at 208°, while that from di *p*-amido-diphenyl and *m* nitro benzoic aldehyde melts at 234° (Schiff a Vanni, A 258, 375)

***m* NITRO-BENZYLIDENE ANILINE**

$C_{11}H_9N_2O_2$ *z* $[3 1]C_6H_4(NO_2)CHNPh$ [61°] Formed from *m*-nitro-benzoic aldehyde and aniline (Lazorenko, J 1870, 760) Needles

***p*-Nitro-benzylidene aniline**

[4 1] $C_6H_4(NO_2)CHNPh$ [93°] Formed by heating aniline with *p* nitro benzoic aldehyde at 100° (Fischer, B 14, 2524)

Di-*m*-nitro-benzylidene-aniline

[3 1] $C_6H_4(NO_2)CHNCH_2C_6H_4(NO_2)$ [1 3] [158°] Formed by warming the alcoholic solution of *m* nitro-benzoic aldehyde with *m* nitro aniline (Hantzsch, B 23, 2775).

***m*-NITRO-BENZYLIDENE BROMIDE**

$C_6H_4(NO_2)CHBr$ [1.3] [102°] Formed by

NITRO-BENZOPHENONE.

heating *m*-nitro toluene (1 mol) with bromine (2 mols) at 140° (Wachendorff, *A* 185, 278) Minute needles (from alcohol)

p-Nitro-benzylidene bromide

$C_6H_4(NO_2)CHBr_2$ [14] [82°] Formed by heating *p*-nitro toluene (1 mol) with bromine (2 mols) at 140° (W) Needles or rectangular plates (from alcohol) By heating with aniline it yields *p*-rosaniline (Zimmermann & Muller, *B* 17, 2936)

m-NITRO BENZYLIDENE CHLORIDE

$C_6H_4(NO_2)CHCl$ [65°] Formed from *m*-nitro benzoic aldehyde and PCl_5 (Widmann, *B* 13, 676, Erlich, *B* 15, 2010) Thin monoclinic plates or needles (from alcohol), ν sol ether

p-Nitro-benzylidene chloride

$C_6H_4(NO_2)CHCl_2$ [46°] Formed from *p*-nitro benzoic aldehyde and PCl_5 (Zimmermann & Muller, *B* 17, 2937, 18, 997), and by nitration of benzylidene chloride (Hubner & Beute, *B* 6, 803) Prisms (from alcohol) Yields *p*-rosaniline on heating with aniline

o NITRO-BENZYLIDENE HYDRAZINE

$(C_6H_4(NO_2)CH)_2N_2$ [181°] Formed from *o*-nitrobenzoic aldehyde and hydrazine salts (Curtaub & Jay, *J pr* [2] 39, 43) Yellow needles

p-NITRO-BENZYLIDENE INDOXYL *v* Indoxide of *p*-NITRO BENZOIC ALDEHYDE

o-NITRO BENZYLIDENE MALONIC ACID

$C_6H_4(NO_2)CH(CO_2H)_2$ [2] $C_6H_4(NO_2)CH(CO_2H)_2$ [161°] Formed by heating *o*-nitro benzoic aldehyde with malonic acid and HOAc at 60° (Stuart, *O J* 47, 158, 49, 365) Needles (from water), ν sol ether, ν a. sol ether, sl sol $CHCl_3$ With HBr it yields a yellow compound melting at 227° Boiling water decomposes it slowly, forming *o*-nitro benzoic aldehyde, malonic acid, and a little *o*-nitro cinnamic acid — $BaA''2'$ feathery tufts, converted by warm water into insoluble $BaA''aq$ — AgA''

Ethyl ether EtA'' [53°]

m-Nitro-benzylidene malonic acid

$[3] C_6H_4(NO_2)CH(CO_2H)_2$ [205°] Formed from *m*-nitro benzoic aldehyde, malonic acid, and HOAc (Stuart, *O J* 47, 155, 49, 361) Crystals, sl sol cold water and ether Partially decomposed by hot water into *m*-nitro benzoic aldehyde and malonic acid Split up on fusion into CO_2 and *m*-nitro-cinnamic acid [196°] HBr forms $C_6H_4(NO_2)CHBrCH(CO_2H)_2$ Bromine combines with it, yielding the dibromide $C_6H_4(NO_2)CHBrCBr(CO_2H)_2$

Ethyl ether EtA'' [73°]

p-Nitro-benzylidene-malonic acid

$C_6H_4(NO_2)CH(CO_2H)_2$ [227°] Formed from *p*-nitro-benzoic aldehyde, malonic acid, and HOAc at 60° (Stuart, *O J* 49, 408) Formed also, together with a small quantity of the *ortho*-acid from benzylidene malonic ether by nitration and saponification (Stuart, *O J* 47, 155) Crystals, split up on fusion into CO_2 and *p*-nitro-cinnamic acid Decomposed by hot water into *p*-nitro-benzoic aldehyde and malonic acid Bromine forms $C_6H_4(NO_2)CHBrCBr(CO_2H)_2$, which gives off HBr at 100°, leaving a residue [188°], and is decomposed by water yielding $C_6H_4(NO_2)CHBrCO_2H$ [208°]

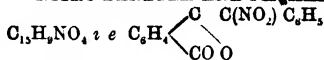
Ethyl ether EtA'' [93°] Prisms Converted by HBr into $C_6H_4(NO_2)CHBrCH(CO_2H)_2$, [99°] (Stuart, *O J* 49, 362).

m-NITRO-BENZYLIDENE DI-METHYL

DISULPHONE $C_6H_4(NO_2)CH(SO_2CH_3)_2$ [179°] Formed by oxidising *m*-nitro benzylidene di-thio di-glycolic acid (Bongartz, *B* 21, 487) Slender needles (from dilute HOAc)

p-Nitro-benzylidene di-methyl disulphone $[14] C_6H_4(NO_2)CH(SO_2CH_3)_2$ [248°] Formed by oxidising *p*-nitro benzylidene di-thio di-glycolic acid with $KMnO_4$ (B) Yellowish needles (from hot water)

NITRO-BENZYLIDENE PHTHALIDE

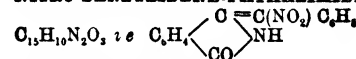


Preparation—10 pts of crude di-nitro-benzyl phthalide are dissolved in 20 pts of hot alcohol, diluted with 10 pts of hot water, and heated for half an hour on the water bath, the yield is 46 p c of theoretical, but when pure di-nitro compound is used, nearly 100 p c (Gabriel, *B* 18, 1251, 3471)

Properties—Glistening crystals Split up on dry distillation into phenyl cyanate and phthalic anhydride Alcoholic NaOH converts it into the salt $C_6H_4NO_2Na_2$ which crystallises in colourless prisms and is decomposed by acids into *o*-nitro toluene $C_6H_4(NO_2)CH_3$ and phthalic anhydride Reduced by HI and P to isobenzylidene phthalide $C_6H_4 \begin{array}{l} \diagup CHCPh \\ \diagdown COO \end{array}$

[91°] and a compound $C_6H_4NO_2$ [121°] whence KOH and MeI yield two isomeric bodies $C_6H_4NO_2$, melting at 237° and 121° (Gabriel, *B* 20, 2863)

NITRO-BENZYLIDENE-PHTHALIMIDINE



Phthalimidylnitrobenzyl Formed together with oxy-nitrobenzyl-phthalimidine by leading N_2O_4 or N_2O_5 into a benzene solution of benzyl-phthalimidine or of deoxybenzoin-*o*-carboxylamide $C_6H_4(CO NH_2)COCH_2C_6H_5$ (Gabriel, *B* 18, 2439)

Nitro-benzylidene-phthalimidic acid $C_6H_4NO_2 \text{ } \nu \text{ } C_6H_4(CO_2H)(C NH_2)C(NO_2)C_6H_5$ [145°-150°] Formed by dissolving nitro-benzylidene phthalimidine in hot dilute NaOH and precipitating with an acid (Gabriel, *B* 18, 2440) Acetyl chloride reconverts it into the anhydride Nitrous acid passed into the solution in benzene converts it into nitro-benzylidene phthalide

Salts — $A''Ag$ microcrystalline powder — $A''Ba7aq$ long yellow needles or short prisms Ethyl ether $A''Et$ [155°], yellow crystals.

Nitro-iso benzylidene-phthalimidine ν

(*Py* 142)-NITRO-OXY PHENYL ISOQUINOLINE

NITRO-BENZYLIDENE-DI-THIO-DI-GLY.

COLLIC ACID $C_6H_4NSO_2 \text{ } \nu \text{ } C_6H_4(NO_2)CH(SCH_2CO_2H)_2$ The *o*-, *m*-, and *p*-varieties of this acid are formed by the action of thioglycolic acid on the three nitro-benzoic aldehydes (Bongartz, *B* 21, 479)

o-Acid [123°] Needles (from HOAc)

m-Acid [180°] Needles (from dil HOAc)

p-Acid [162°] Needles (from dil HOAc)

o-NITRO-BENZYLIDENE-DI-UREA

$C_6H_4NO_2 \text{ } \nu \text{ } C_6H_4(NO_2)CH(NHCO NH_2)_2$ [200°] Formed by warming an alcoholic solution of urea with *o*-nitro-benzoic aldehyde (Schiff,

A. 251, 186, Ludy, M 10, 804) Needles (containing aq), sl sol water and alcohol

o-NITRO-BENZYL IODIDE $C_6H_4(NO_2)CH_2I$ [75°]

Formed from *o*-nitro benzyl chloride, KI, and alcohol (Kumpf, A 224, 103)

p Nitro-benzyl iodide $C_6H_4(NO_2)CH_2I$ [127°]

Formed in like manner Needles

p-NITRO-BENZYL-MALONIC ACID

$C_6H_4(NO_2)CH_2CH(CO_2H)_2$ Formed by saponification of the ether which is obtained in small quantity, together with di nitro di benzyl malonic ether, by the action of *p* nitro benzyl chloride upon sodio malonic ether (Lellmann a Schleich, B 20, 434) Yellow powder Carbonises at 240° without melting CaA" and BaA" yellow pps

Ethyl ether Et.A" [63°], yellowish prisms, sol ordinary solvents

Di o nitro di benzyl malonic ether

$(C_6H_4(NO_2)CH_2)_2C(CO_2Et)_2$ [97°] Obtained by the action of *o* nitro benzyl chloride upon sodio-malonic ether Yellow crystals Sol alcohol and ether (Lellmann a Schleich, B 20, 438)

Di p nitro di benzyl malonic ether

$(C_6H_4(NO_2)CH_2)_2C(CO_2Et)_2$ [170°] Formed by the action of *p* nitro benzyl chloride upon sodio malonic ether Colourless silky needles Sol acetic acid, sl sol alcohol, and chloroform

p-NITRO BENZYL MERCAPTAN $C_6H_4NSO_2$

$\epsilon \epsilon C_6H_4(NO_2)CH_2SH$ [140°] Formed from *p*-nitro-benzyl chloride and alcoholic ammonium sulphide (Strakosch, B 5, 698) Lamine

TRI-NITRO-BENZYL-MESITYLENE

$C_6H_3(NO_2)_3$, [185°] Formed by nitration of benzyl-mesitylene at 0° (Louise, A Ch [6] 6, 182) Prisms (from alcohol chloroform)

TRI-NITRO-BENZYL METHYL KETONE

$C_6H_4(NO_2)_3CH_2COCH_3$ [89°] Obtained by boiling tri nitro phenyl acetoacetic ether (10 g) dissolved in HOAc (100 g) with H_2SO_4 (20 g) and water (30 g) for 8 hours (Dittreich, B 23, 2723) Long yellowish white needles, sol alkalis

Phenyl hydrazide $C_6H_3N_2O_5$ [125°]

NITRO p-BENZYL-PHENOL $C_6H_4NO_2$, $\epsilon \epsilon$

$PhCH_2C_6H_4(NO_2)OH$ [134] [75°] From benzyl phenol and HNO_3 (S G 14) Prisms Volatile with steam—KA' brick-red needles (Rennie, C J 41, 221)

Di nitro p benzyl phenol

$PhCH_2C_6H_3(NO_2)_2OH$ [135? 4] [88°] From the above, HOAc, and HNO_3 Formed also by the action of HNO_3 on *p* benzyl phenol sulphonic acid (Rennie, C J 49, 408) Oxidised by CrO_3 it gives benzoic acid—KA', orange needles—BaA'.

Di-nitro o benzyl phenol [82°] Obtained by warming *o*-benzyl phenol sulphonic acid with dilute HNO_3 (Rennie)—KA' aq—BaA'.

Tri - nitro - benzyl phenol $C_6H_3(NO_2)_3OH$ [148°] Got by dissolving potassic benzyl-phenol sulphonic acid in HNO_3 (S G 14) and evaporating (Rennie, C J 41, 36, 223) Silky pale yellow needles Yields *p* nitro-benzoic acid on oxidation with chromic acid mixture— $C_{12}H_7(NO_2)_3OK$

NITRO - p - BENZYL - PHENOL SULPHONIC

ACID $C_6H_4C_6H_4(OH)(NO_2)SO_3H$ —KA' Formed by action of HNO_3 (S G 12) and potassic benzyl-phenol sulphate (Rennie, C J 41, 85)

Nitro - o - benzyl - phenol sulphonic acid Formed in like manner—KA' yellow scales

DI-NITRO BENZYL PHTHALIDE $C_{12}H_8N_2O_5$

$\epsilon \epsilon C_6H_4 \begin{matrix} \diagup C(NO_2)CH(NO_2)C_6H_4 \\ \diagdown COO \end{matrix} (?)$ [113°]

Formed by passing nitrous acid gas into a solution of benzylidene phthalide in benzene (Gabriel, B 18, 1251) Colourless crystals Readily converted into nitro-benzylidene phthalide (q v)

o-NITRO-BENZYL-PHTHALIMIDE

$C_6H_4 \begin{matrix} \diagup CO \\ \diagdown NCH_2C_6H_4(NO_2) \end{matrix}$ [12] [219°]

Formed from potassium phthalimide and *o* nitro-benzyl chloride at 100°-130° (Gabriel, B 20, 2227) Prisms

m Nitro benzyl phthalimide [155°] Formed in like manner Needles

p Nitro benzyl-phthalimide [175°] (S), [172°] (Hafner, C C 1889, 671) Prisms (from HOAc) (Salkowski B 22, 2142)

p NITRO BENZYL-PIPERIDEIN

$C_6H_4(NO_2)CH_2NC_4H_8$ [35°-40°] Formed by dissolving its polymeride in HClAq and ppg by ammonia (Lellmann a Schwaderer, B 22, 1333) Readily polymerises to $C_6H_4(NO_2)CH_2NC_4H_8$, [120 5°], which is also obtained by treating di-piperidein with *p* nitro benzyl chloride and NaOHaq

NITRO BENZYL-PIPERIDINE

$C_6H_4(NO_2)CH_2NC_4H_8$. The *o*, *m*, and *p*-isomerides are formed by the action of *o*, *m*, and *p*-nitro benzyl chloride on piperidine in hot alcoholic solution (Lellmann a Pekrun, A 259, 40)

o Isomeride Oil Reduced by $SnCl_2$ to *o*-amido benzyl piperidine [82 5°]—B'HCl—B'H₂PtCl₆

m Isomeride Oil Reduced by $SnCl_2$ to *m*-amido benzyl piperidine [112°]—B'HCl

p Isomeride [34°] B'HCl—B'H₂PtCl₆

NITRO - BENZYL QUINOLINE TETRA

HYDRIDE $C_6H_4(NO_2)CH_2NC_8H_6N_2$. The three isomerides are formed by heating *o*, *m*, and *p*-nitro benzyl chloride (1 mol) with quinoline tetrahydride (2 mols) in hot alcoholic solution (Lellmann a Pekrun, A 259, 50)

o Isomeride [111°] Brownish red tables—B'H₂PtCl₆, yellow amorphous pp

m Isomeride [99°] Red prisms, reduced by $SnCl_2$ to *m*-amido benzyl-quinoline tetrahydride [82°]

p Isomeride [102°] Red prisms—B'H₂PtCl₆

NITRO BENZYL SELENOCYANIDE

$C_6H_4N_3SeO_4$, $\epsilon \epsilon C_6H_4(NO_2)CH_2SeCN$ [122 5°]

Formed by nitration of benzyl selenocyanide at -4° (C L Jackson B 8, 321, A 179, 16)

Needles (from alcohol)

o NITRO BENZYL SULPHIDE

$(C_6H_4(NO_2)CH_2)_2S$ [124°] Formed, together with a little of the disulphide, by passing NH_3 and H_2S into an alcoholic solution of *o* nitro-benzyl chloride (Jahoda, M 10, 880) White plates Yields on oxidation by HNO_3 the sulphoxide $(C_6H_4(NO_2)CH_2)_2SO$ [163°] and the sulphone $(C_6H_4(NO_2)CH_2)_2SO_2$ [200°].

o Nitro-benzyl disulphide

$(C_6H_4(NO_2)CH_2)_2S_2$ [47°] Formed as above (J)

p-Nitro-benzyl disulphide [89°] Yellow crystals (Strakosch, B 5, 698)

p-NITRO-BENZYL SULPHOCYANIDE

$C_6H_4(NO_2)CH_2SCN$. Formed from *p* nitro-

NITRO-BENZOPHENONE.

benzyl chloride and alcoholic potassium sulphocyanide (Henry, *B* 2, 638) Small needles

***p* NITRO-BENZYL-DI-THIO-CARBAMIC ACID** $\text{NO}_2\text{C}_6\text{H}_4\text{CH}_2\text{NHCS}_2\text{SH}$ The *p* nitro benzyl ammonium salt [193°] of this acid is formed by treating *p* nitro benzylamine with CS_2 in ether (Hafner, *B* 23, 339)

***Di-p*-NITRO-DI-BENZYL-THIO-UREA** $\text{OS}(\text{NHCH}_2\text{C}_6\text{H}_4\text{NO}_2)_2$ [202°] Formed by boiling *p*-nitro benzyl ammonium *p* nitro-benzyl di thio carbamate with HgO and alcohol (Hafner, *B* 23, 340) Needles, sl sol alcohol

***o* NITRO-BENZYL-*p* TOLUIDINE** $\text{C}_6\text{H}_4(\text{NO}_2)\text{CH}_2\text{NHCH}_2\text{Me}$ [72°] Obtained by heating 4 pts of *p* toluidine with 1 pt of *o*-nitro benzyl chloride for $\frac{1}{2}$ hr on the water bath Yellow crystals V sol most ordinary solvents Salts — B^+HCl colourless needles — $\text{B}^+\text{H}_2\text{PtCl}_6$ sparingly soluble pp — $\text{B}^+\text{H}_2\text{SO}_4$ thin glistening plates

Acetyl derivative

$\text{C}_6\text{H}_4(\text{NO}_2)\text{CH}_2\text{N}(\text{Ac})\text{CH}_2\text{Me}$ [65°], stout crystals, on reduction with tin and HCl it is converted into *o* amido benzyl toluidine [79°] (Lellmann a Stuckel, *B* 19, 1609, 24, 718)

Formyl derivative [79°] Formed by heating a solution of formyl *p* toluidine in benzene with sodium till dissolved, and then adding *p* nitro benzyl chloride (Paal a Busch, *B* 22, 2695) Needles

***o* Nitro benzyl *o* toluidine** **Formyl derivative** $\text{C}_6\text{H}_4(\text{NO}_2)\text{CH}_2\text{N}(\text{C}_6\text{H}_4)\text{CHO}$ [76°] Formed from *o* nitro benzyl chloride and sodium formyl *o*-toluidine (Paal a Busch, *B* 22, 2701) Yellow needles grouped in spheres

***p* NITRO BENZYL UREA** $\text{C}_6\text{H}_4\text{N}_2\text{O}_3$ $\text{NH}_2\text{CO NHCH}_2\text{C}_6\text{H}_4\text{NO}_2$ [197°] Formed by heating *p* nitro benzylamine hydrochloride with silver cyanate at 100° (Hafner, *B* 23, 339) Pale yellow needles, v sol HOAc and alcohol

***Di-p*-NITRO-DI-BENZYL UREA** $\text{CO}(\text{NHCH}_2\text{C}_6\text{H}_4\text{NO}_2)_2$ [234°] Formed by boiling the corresponding thio urea with H_2O , or by adding a solution of COCl_2 in benzene to one of *p* nitro benzylamine in ether (Hafner, *B* 23, 340) Silvery needles (from HOAc)

NITRO BROMO compounds v BROMO NITRO COMPOUNDS

NITRO BRUCINE v BRUCINE

***o* NITRO-*n* BUTANE** $\text{C}_4\text{H}_9\text{NO}_2$ $\text{CH}_3\text{CH}_2\text{CH}_2\text{CH}_2\text{NO}_2$ (152° cor) *S* G Σ 9945 Formed from *n* butyl iodide and AgNO_2 (Zublin, *B* 10, 2083, Pribram a Handl, *M* 2, 656) Reduced by tin and HCl to butylamine HClAq at 140° forms hydroxylamine and *n* butyric acid

***o* Nitro butane** $\text{CH}_3\text{CH}_2\text{CH}(\text{NO}_2)\text{CH}_3$ (139°) Formed from *sec* butyl iodide (120 pts) and silver nitrite (150 pts) (V Meyer a Locher, *B* 7, 1506, *A* 180, 134) Formed also from $\text{CH}_3\text{CHBrNO}_2$ and ZnEt_2 (Bevad, *J* R 20, 125)

***o*-NITRO ISOBUTANE** $(\text{CH}_3)_2\text{CHCH}_2\text{NO}_2$ (137°-144°) *S* G Σ 1 0088 Formed from *sec* isobutyl iodide and AgNO_2 (Demole, *B* 7, 709, 790, *A* 175, 142, *P* a *H*) Unlike *o* nitro *n* butane, it does not give a crystalline pp with NaOEt , although it dissolves in alkalis

Nitro *tert*-butane $(\text{CH}_3)_3\text{CNO}_2$ (110°-180°) Formed, in small quantity, together with *tert* butyl nitrite, by the action of AgNO_2 on *tert* butyl iodide (Tscherniak, *A* 180, 155) Oil, smelling like peppermint Does not dissolve in

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alkalis May be reduced to *tert* butylamine (*cf* V Meyer, *A* 244, 222)

***Di*-nitro-butane** $\text{C}_4\text{H}_8\text{N}_2\text{O}_4$ Formed from bromo *o* nitro *n* butane, aqueous KNO_2 and dilute H_2SO_4 (Zublin, *B* 10, 2085) Oil, decomposed at 190° by distillation — $\text{KC}_4\text{H}_7\text{N}_2\text{O}_4$ golden scales, sol water and alcohol — $\text{AgC}_4\text{H}_7\text{N}_2\text{O}_4$ yellow scales with blue reflex

***Di* nitro butane** $\text{C}_4\text{H}_8(\text{NO}_2)_2$ (197°) *S* G Σ 1 205 Formed by the action of HNO_3 on di isoamyl ketone or on propyl acetoacetic ether (Chancel, *C* R 94, 399, 96, 1466) Heavy oil, forming crystalline *K* and *Ag* salts Decomposed on distillation

***Di*-(*o*)-nitro-butane** $\text{CH}_3\text{CH}_2\text{C}(\text{NO}_2)_2\text{CH}_3$ (199° cor) Formed by oxidising butyl ψ nitrole with nitric acid (V Meyer, *B* 9, 701), or by boiling isovaleric acid with HNO_3 (Bredt, *B* 15, 2324) Oil, not soluble in alkalis Tin and HCl convert it into hydroxylamine and methyl ethyl ketone

***Di*-nitro-isobutane** $(\text{CH}_3)_2\text{CHCH}(\text{NO}_2)_2$ Formed from bromo nitro-isobutane, KNO_2 and dilute H_2SO_4 (*Z*) Oil — $\text{KC}_4\text{H}_7(\text{NO}_2)_2$ — $\text{AgC}_4\text{H}_7(\text{NO}_2)_2$ *saq*

***Di*-NITRO-ISOBUTYL-ANILINE** $\text{C}_6\text{H}_4\text{N}_2\text{O}_4$ $\text{C}_6\text{H}_4\text{NHCH}_2\text{C}_4\text{H}_8(\text{NO}_2)_2$ [80°] Formed from bromo *m* di nitro benzene and isobutylamine (Romburgh, *R* T C 4, 192) Yellow needles

***Tri* nitro isobutyl aniline** $\text{C}_6\text{H}_4(\text{NO}_2)_3\text{NHCH}_2\text{C}_4\text{H}_8$ [95°] Formed from chloro tri nitro benzene (picryl chloride) and isobutylamine (*li*) Converted by fuming HNO_3 into the nitramine $\text{C}_6\text{H}_4(\text{NO}_2)_3\text{N}(\text{NO}_2)\text{C}_4\text{H}_8$ [110°]

***m* NITRO ISOBUTYL BENZENE** $\text{C}_6\text{H}_4\text{H}_2\text{NO}_2$ $\text{C}_6\text{H}_4\text{CH}_2\text{CH}_2\text{CH}_2\text{NO}_2$ (251°) at 740 mm Formed from nitro amido isobutyl benzene by elimination of NH_2 (Gelzer, *B* 21, 2941) Oil Yields *m* nitro benzoic acid on oxidation

***o* Nitro-*tert* butyl benzene** $\text{C}_6\text{H}_4(\text{NO})\text{CMe}_3$ [12] (249°) *S* G Σ 1 074 Formed from *tert* butyl benzene and fuming HNO_3 (Senkowsky, *B* 23, 2416) Yellow oil, smelling like cymene Reduced by tin and HCl to $\text{C}_6\text{H}_4(\text{NH}_2)\text{CMe}_3$ (*c* 235°), *S* G Σ 977, which yields an acetyl derivative [159°]

***p*-Nitro-*tert*-butyl benzene** $\text{C}_6\text{H}_4(\text{NO}_2)\text{CMe}_3$ [14] (30°) (276°) Formed at the same time as the preceding (*S*) Yellow needles (from alcohol) Yields, on reduction, $\text{C}_6\text{H}_4(\text{NH}_2)\text{CMe}_3$, *S* G Σ 953

NITRO *m* ISOBUTYL BENZOIC ACID $\text{C}_6\text{H}_4(\text{C}_4\text{H}_8)(\text{NO})\text{CO}_2\text{H}$ [140°] Formed by nitration of *m* isobutyl benzoic acid (Kelbe a Pfeiffer, *B* 19, 1727) Small needles (from petroleum ether) — AgA' somewhat soluble pp — MeA' liquid

Nitro *p* isobutyl benzoic acid $\text{C}_6\text{H}_4(\text{C}_4\text{H}_8)(\text{NO}_2)\text{CO}_2\text{H}$ [161°] Long fine needles (from water) Formed by nitration of *p* isobutyl benzoic acid — AgA' white pp

Methyl ether MeA' fluid (Kelbe a Pfeiffer, *B* 19, 1726)

NITRO-BUTYLENE $\text{C}_4\text{H}_7\text{NO}_2$ (154°-158°) Formed by allowing HNO_3 (*S* G 1 52) to drop into *tert* butyl alcohol (Hastinger, *Sitz* W 77 [2] 428, *A* 193, 366, *M* 2, 286) Formed also in small quantity by saturating HNO_3 with iso butylene (*H*) Pale yellow oil, heavier than water Dissolves in alkalis, and is reppd by acids, Excess of water at 100° splits it up into

N N

acetone and nitro methane Br unites, forming only $C_2H_5Br_2NO_2$ — $NaC_2H_5NO_2$ powder, v sol water

NITRO ISOBUTYL-PHENOL

$C_6H_5C_3H_7(NO_2)(OH)$ [1 3 4] [95°] (290°) at 711 mm Formed by boiling nitro amido isobutyl benzene with dilute potash (Gelzer, *B* 21, 2947) Red needles (from alcohol), v sol hot water

Di-nitro isobutyl-phenol $C_6H_5C_3H_7(NO_2)_2OH$ [98°] Formed from isobutyl phenol [99°], HOAc, and HNO_3 (Studer, *A* 211, 244, *B* 14, 1474, Liebmann, *B* 14, 1842) Sulphur yellow needles (from alcohol) Yields di nitro amido isobutyl benzene on heating with NH_3Aq

TRI-NITRO-ISOBUTYL-TOLUENE

$C_9H_7Me_2(NO_2)_3$ Artificial musk [97°] Formed by heating isobutyl toluene with HNO_3 and H_2SO_4 for 24 hours on a water-bath (Baur, *J* R 111, 238) White needles, sol alcohol and ether Its solutions smell like musk Forms a crystalline compound with naphthalene [90°]

TRI NITRO ISOBUTYL XYLENE

$C_{10}Me_2(C_6H_4)(NO_2)_3$ [110°] Formed by nitration of isobutyl xylene (Baur, *C* R 111, 238) White needles Its alcoholic solution smells like musk

NITRO-CAMPHOLENIC ACID v CAMPHOLENIC ACID

NITRO-CAMPHOR v CAMPHOR

TETRA NITRO CARBAZOLE

$C_{12}H_6(NO_2)_4NH$ Formed by nitrating carbazole (Graebe, *A* 202, 26) Lemon yellow crystals (from HOAc), insoluble in alcohol and ether— $C_{12}H_6(NO_2)_4NK$ insol water

Four tetra-nitro carbazoles have been described by Ciamician and Silber (*G* 12, 277) as formed by the nitration of carbazole The melting points of three of them are 308°, above 320°, and about 285°, while the fourth decomposes before melting

NITRO CARBOXY CINNAMIC ACID

$C_9H_7(OH\ CH\ CO_2H)(NO_2)(CO_2H)$ [2 4] [287°] (Löw, *A* 231, 371) Formed by nitrating carboxy cinnamic acid Hemispherical aggregates or regular tablets (from water) That the NO_2 is in the *o* position is shown by the fact that, by Baeyer's synthesis, it gives rise to indigo di carboxylic acid Heated with conc H_2SO_4 it does not turn blue It yields a dibromide, which is converted by aqueous NaOH into nitro carboxy phenyl-propionic acid

NITRO-CARBOXY-PHENYL-PROPIONIC

ACID $C_9H_7(NO_2)(CO_2H)CH\ CH\ CO_2H$ [3 1 4] ? [192°] Got by nitration (Widman, *B* 22, 2273)

NITRO-CARVACROL

$C_{10}H_{14}(C_6H_5)Me(NO_2)(OH)$ [78°] Formed from nitroso carvacrol, KOH, and K_2FeC_4 (Paterno a Canzoneri, *G* 10, 235) Needles, almost insol water

NITRO-CHLORO-derivatives v CHLORO-NITRO-derivatives

NITRO-CHLOROFORM v TRI-CHLORO-NITRO-METHANE

DI-NITRO-CHOLESTERIN $C_{26}H_{42}(NO_2)_2O$ [121°] Obtained by nitration of cholesterol (Preis a Raymann, *B* 12, 224) Colourless needles, sl sol cold alcohol. By boiling a hot saturated solution of cholesterol in HOAc with HNO_3 (S G 1 54), Reinitzer (*M* 9, 440) obtained a nitro compound melting at 94° By adding cholesterol (1 pt) to a mixture of HOAc (10

pts) and fuming HNO_3 (3 pts), Latschnoff (*J* R 10, 360) obtained a compound crystallising in plates, decomposing at 180° without melting

NITRO CHOLESTERYL CHLORIDE

$C_{26}H_{42}(NO_2)Cl$ [148°–149°] Colourless needles Prepared by nitration of cholesteryl chloride (Preis a Raymann, *B* 12, 225)

NITRO CHRYSENE v CHRYSENE

NITRO-CHRYSOQUINONE v CHRYSOQUINONE

NITRO-CINCHONAMINE v CINCHONA BASES

o-NITRO CINNAMIC ACID $C_9H_7NO_2$

ve [2 1] $C_9H_7(NO_2)CH\ CH\ CO_2H$ Mol w 193 [232°] (M), [237°] (T a O), [240°] (Baeyer, *B* 13, 2257) Formed, together with the *p* isomeride, by nitrating cinnamic acid (Beilstein a Kuhlberg, *A* 163, 126, Muller, *A* 212, 124) Formed also from *o* nitro benzaldehyde, Ac_2O , and NaOAc (Gabriel a Meyer, *B* 14, 830)

Preparation—1 Cinnamic acid (1 pt) is dissolved in nitric acid (5 pts), the mixture is poured upon snow, and the *ppd* acids separated by means of alcohol (B a K)—2 Cinnamic ether (10 g) is dropped into fuming HNO_3 at 0° The solution is poured at once into water at 0° The solid *pp* is washed with water, and treated with alcohol, which dissolves the *o*, leaving the *p* compound (Stuart, *C* J 43, 408) Fair yield (4 g)—3 The product of nitration of cinnamic acid is etherified, and the *o* nitro cinnamic ether is separated from the *p* isomeride by means of its greater solubility in ether (Tiemann a Oppermann, *B* 13, 2060) The acid may then be obtained from its ether by hydrolysis with H_2SO_4 (Fischer a Kuzel, *A* 221, 265)

Properties—Needles, insol water, sl sol cold alcohol Yields *o* nitro benzoic acid on oxidation by chromic acid mixture Its solution in H_2SO_4 becomes blue on warming or on standing

Salts— CaA' , 2aq yellow needles— BaA' , 2aq minute yellow needles

Methyl ether MeA' [73°]

Ethyl ether EtA' [42°] (M), [44°] (B)

Trimetric crystals, $a\ b\ c = 927\ 1\ 517$ Readily combines with bromine Reduced by aqueous ammonium sulphide to carbostyryl In hot alcoholic solution tin and HCl reduce it to *o* amido cinnamic ether

Chloride $C_9H_7(NO_2)CH\ CH\ COCl$ [65°]

Crystalline solid (Fischer a Kuzel, *B* 16, 34)

m Nitro cinnamic acid

[3 1] $C_9H_7(NO_2)CH\ CH\ CO_2H$ [197°] Prepared by heating *m*-nitro benzoic aldehyde with Ac_2O and NaOAc (Schiff, *B* 11, 1783, Tiemann a Oppermann, *B* 13, 2060) Yellow needles Yields *m* nitro benzoic acid on oxidation A mixture of HNO_3 (2 pts of S G 1 5) and conc H_2SO_4 (5 pts) converts it at 0° into *mm* di nitro styrene (Friedlander a Lazarus, *A* 229, 233) Reduced by tin and HCl to *m* amido cinnamic acid— AgA' insoluble *pp*

Ethyl ether EtA' [79°]

p-Nitro cinnamic acid

[4 1] $C_9H_7(NO_2)CH\ CH\ CO_2H$ [286°] (T a O), [288°] (D) Formed by nitration of cinnamic acid (Mitscherlich, *A* Oh. [3] 4, 78, E Kopp, *C* R 53, 634, Tiemann a. Oppermann, *B* 13, 2059)

Preparation—From cinnamic acid (1 pt) and cold HNO_3 (5 pts) The acids produced are etherified, the *p* ether, [189°], crystallising from

alcohol It is saponified by H_2SO_4 (1 pt), HOAc (1 pt), and water (1 pt) (Drewson, *A* 212, 150)

Properties—Prisms, ν sl sol boiling alcohol Yields *p*-oxy-benzoic acid on oxidation

Salts— KA' very soluble crystals— CaA' , 8aq— CaA' , 2aq— SrA' , 5aq— BaA' , 3aq minute needles— MgA' , 6aq nodules— HgA' , ν — $\text{Hg}_2\text{A'}$, Cl_2 , 8aq— AgA' insoluble pp

Methyl ether MeA' [161°] (286°)

Ethyl ether EtA' [139°] (B & K), [137°] (Muller, *A* 212, 125) Combines with bromine forming a dibromide whence alcoholic potash produces two bromo nitro cinnamic ethers, melting at 63° and 93° Reduced by tin and HCl to *p* amido cinnamic acid and *p*-amido-styrene (Bender, *B* 14, 2359)

Anhydride ($\text{C}_6\text{H}_4(\text{NO}_2)\text{CH}(\text{CH})\text{CO}_2\text{O}$)

Formed from the K salt and POCl_3 (Chiozza, *A* *Ch* [2] 39, 231) Melts under boiling water

Amide $\text{C}_6\text{H}_4(\text{NO}_2)\text{CH}(\text{CH})\text{CONH}_2$ [155°–160°] (Cahours, *A* *Ch* [3] 27, 452)

***a m* Di-nitro-cinnamic ether**

$\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CH}(\text{C}(\text{NO}_2)\text{CO}_2\text{Et})$ Formed from *m* nitro cinnamic ether, HNO_3 (1 pt) and H_2SO_4 (2 pts) below 20° (Friedlander & Lazarus, *A* 229, 235) Thick plates (from ether) Insol water and light petroleum

Reactions—1 With alcohol, on warming, it forms $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CH}(\text{OEt})\text{CH}(\text{NO}_2)\text{CO}_2\text{Et}$, a colourless oil When this is dissolved in ether and ammonia is passed in, a pp of $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CH}(\text{OEt})\text{C}(\text{NH}_2)(\text{NO}_2)\text{CO}_2\text{Et}$ is formed This is soluble in water and gives pps with many metallic salts The compound $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CH}(\text{OEt})\text{CH}(\text{NO}_2)\text{CO}_2\text{Et}$ is converted by the simultaneous action of bromine and NaOH into the ethyl derivative of di bromo di nitro *a* phenyl methyl carbinol—2 Boiled with water it forms alcohol, nitro methane, CO_2 , and *m* nitro benzoic aldehyde

***ap* Di nitro cinnamic acid**

[4] $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CH}(\text{C}(\text{NO}_2)\text{CO}_2\text{H})$ Formed from *p* nitro cinnamic acid, HNO_3 , and H_2SO_4 at –10° to –20° White plates* At 0° it decomposes into CO_2 and *ap* di nitro styrene

Methyl ether MeA' [127°]

Ethyl ether EtA' [110°] Formed from *p* nitro cinnamic ether (1 pt), HNO_3 (2 pts of $\text{S.G. } 1.5$), and H_2SO_4 (4 pts) (Friedlander & Mahly, *A* 229, 210) Thin plates (from benzene ligroun), ν sol benzene, nearly insol ligroun

Reactions—1 CrO_3 in HOAc oxidises it to *p* nitro benzoic acid—2 $\text{K}_2\text{Cr}_2\text{O}_7$ and HOAc forms *p* nitro benzoic aldehyde—3 Boiling water decomposes it into alcohol, CO_2 , *p* nitro benzoic aldehyde, and nitro methane Boiling dilute acids behave in the same way, only the nitro-methane appears as hydroxylamine—4 A solution in conc H_2SO_4 at 100° poured into water gives a pp of *p* nitro benzaloxim—5 Boiling dilute (5 p c) Na_2CO_3 forms bright yellow plates of $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CH}(\text{OEt})\text{CH}(\text{NO}_2)\text{CO}_2\text{Et}$ [52°], whence the alcohol cannot be removed by heating at 110°, or even with dilute HCl at 90° Methyl alcohol forms the corresponding $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CH}(\text{OMe})\text{CH}(\text{NO}_2)\text{CO}_2\text{Et}$ [77°]—7 Tin and HCl reduce it to *ap*-diamido-phenyl-propionic ether and *p*-amido phenyl-acetomitrile

***o* NITRO CINNAMIC ALDEHYDE** $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CH}(\text{CH})\text{CHO}$ [127°] Formed by boiling *o* nitro oxy phenyl propionic aldehyde with Ac_2O (Baeyer & Drewson, *B* 16, 2207) Formed also by condensation of *o* nitro benzoic aldehyde with aldehyde by means of dilute NaOH , the yield being 40 p c of the theoretical (Diehl & Einhorn, *B* 20, 2335)

Preparation—25 g of cinnamic aldehyde are slowly added to 500 g of conc H_2SO_4 containing 20 g of KNO_3 . The solution is precipitated in water, and the mixture of *o*- and *p* nitro cinnamic aldehydes is crystallised from alcohol with animal charcoal The product is dissolved in absolute alcohol and the boiling solution mixed with an equal volume of sodium bisulphite solution, the solution is quickly cooled and separates the greater part of the bisulphite compound of the *p* isomeride, the remaining portion being salted out by the addition of NaCl , the filtrate contains the bisulphite of the *o*-isomeride The two isomerides are obtained from their bisulphites by decomposing the latter in aqueous solution with H_2SO_4 (Diehl & Einhorn)

Properties—Colourless needles, ν sol boiling water and CHCl_3 , sl sol alcohol and ether Combines with bisulphites Yields quinoline on reduction On heating with malonic acid in HOAc it yields $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CH}(\text{CH})\text{CH}(\text{OH})\text{CH}(\text{CO}_2\text{H})_2$ [269°] and *o* nitro phenyl butinene dicarboxylic acid $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CH}(\text{CH})\text{CH}(\text{CO}_2\text{H})_2$ [213°] (Einhorn & Gehrenbeck, *A* 253, 374)

Phenyl hydrazide

$\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CH}(\text{CH})\text{CH}(\text{N}(\text{HPh}))\text{CONHPh}$ [158°] Needles

Di phenyl hydrazide

$\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CH}(\text{CH})\text{CH}(\text{N}(\text{Ph}))\text{CONHPh}$ [69°] Yellow crystals (Cornelius & Homolka, *B* 19, 2240)

***m* Nitro cinnamic aldehyde**

$\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CH}(\text{CH})\text{COH}$ [116°] Formed by eliminating H_2O from *m* nitro β oxy β phenyl-propionic aldehyde (Göhring, *B* 18, 720) Prepared by dissolving 100 pts of *m*-nitro benzaldehyde in 2,000 pts of alcohol, diluting with 4,000 pts of water, and adding at once to the milky liquid 35 pts of conc acetic aldehyde and 70 pts of 10 p c aqueous NaOH After 12 hours' standing the pp is separated, pressed, washed, and crystallised, the yield is 50 p c of the theoretical (Kunkeln, *B* 18, 483) Long thin prisms, ν sol benzene and acetic acid, sl sol cold alcohol, ether, and hot water Unites with Br forming $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CHBrCHBrCHO}$ [c 90°]

Phenyl hydrazide

$\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CH}(\text{CH})\text{CH}(\text{N}(\text{HPh}))\text{CONHPh}$ [160°], red tables.

***p*-Nitro-cinnamic aldehyde**

[4] $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CH}(\text{CH})\text{CHO}$ [142°] Formed from *p*-nitro benzoic aldehyde, aldehyde, and dilute NaOH , the resulting *p* nitro β oxy- β -phenyl propionic acid being boiled with HOAc (Göhring, *B* 18, 372, Einhorn, *A* 253, 348) Formed also, together with the *o* isomeride, by nitration of cinnamic aldehyde (*supra*) Colourless needles Combines with bisulphites Condenses with acetone in presence of NaOH to $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CH}(\text{CH})\text{CH}(\text{CH})\text{CO}$ [218°] and $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CH}(\text{CH})\text{CH}(\text{CH})\text{COCH}_3$ [132°], the last compound forming a phenyl-hydrazide melting at 210°

Oxim $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CH}(\text{CH})\text{CH}(\text{NOH})$ [179°]

Phenyl-hydrazide

$\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CH}(\text{CH})\text{CH}(\text{N}(\text{HPh}))\text{CONHPh}$ [181°] Orange

red crystals, forming a red solution in conc H_2SO_4 .

Anhydride $\text{C}_6\text{H}_5(\text{NO}_2)\text{C}_6\text{H}_4\text{NPh}$ [183°]

***o*-NITRO-CINNAMOYL-ACETO ACETIC ETHER** $\text{C}_6\text{H}_5(\text{NO}_2)\text{CH}(\text{CH}_3)\text{COCH}_3$ [120°] Prepared by the action of *o* nitro cinnamoyl chloride on sodio-acetacetic ether (Fischer & Kuzel, *B* 16, 84) Yellow prisms Sol chloroform, sl sol alcohol and ether It forms stable salts with alkalis, soluble in water with a reddish-yellow colour Gives a dark red colouration in alcoholic solution with Fe_2Cl_6 Boiled with 30 p.c dilute H_2SO_4 , it gives nitro cinnamoyl acetone

***o*-NITRO-CINNAMOYL ACETONE**

$\text{C}_6\text{H}_5(\text{NO}_2)\text{CH}(\text{CH}_3)\text{COCH}_3$ [118°] Prepared by boiling *o* nitro cinnamoyl-aceto acetic ether with dilute H_2SO_4 (30 p.c) By longer boiling with dilute H_2SO_4 , it converted into *o* nitro-styryl-methyl ketone (*o*-nitro benzylidene acetone) (Fischer & Kuzel, *B* 16, 85) Fine yellow prisms Sol hot alcohol, sl sol cold alcohol, ether and CS_2 Gives a red colouration with Fe_2Cl_6 Dissolves in alkalis with a yellow colour

NITRO-COCCUSIC ACID *v* **TRI-NITRO OXY-M-TOLUIC ACID**

NITRO-CODEINE *v* **CODEINE**

NITRO-COMENIC ACID *v* **COMENIC ACID**

NITRO-COMPOUNDS Compounds containing the group nitroxyl (NO_2) directly united, by means of its nitrogen, to carbon Their most general characteristic is that they yield amido compounds on reduction (*v* **AMINES** and **AMIDO ACIDS**)

Formation—1 Fatty nitro compounds are formed by the action of silver nitrite on alkyl iodides The isomeric nitrites are also formed in this reaction—2 Aromatic nitro compounds are formed by the direct action of nitric acid The nitric acid must usually be concentrated, and its action is intensified by mixture with H_2SO_4 Phenols and oxy acids may be nitrated by dilute nitric acid Various anilides, boiled with dilute nitric acid (S.G. 1.029), are converted into di-nitro derivatives of the base Thus acetyl methyl-aniline becomes di-nitro methyl aniline (Norton & Allen, *B* 18, 1995)—3 Aromatic amido compounds may be converted into the corresponding nitro compounds by the cuprous reaction (Sandmeyer, *B* 20, 1495) For this purpose cupric sulphate (50 g) is dissolved, together with glucose (15 g), in boiling water (100 c.c.), and at once treated with soda (20 g) dissolved in water (60 c.c.) After cooling, the mixture is neutralised with acetic acid To this mixture the nitrate of the diazotised base (prepared from the base, HNO_3 , and NaNO_2) is added in the cold—4 Fatty compounds of the form X CHBrNO_2 are converted by KNO_3 in alkaline solution into salts of di-nitro compounds of the form $\text{X CK}(\text{NO}_2)_2$.

Reactions—1 Acid reducing agents yield amido compounds directly, but alkaline reducing agents (*eg* sodium amalgam, zinc-dust and NaOH) acting upon aromatic nitro compounds yield in the first place intermediate bodies, *v* **Azo-** and **Azoxy-** compounds In some cases where reduction is effected by tin and HClAq chlorination may take place, thus *m*-nitro-toluene gives a chloro-toluidine The best

general reducing agent is a solution of SnCl_2 in HClAq —2 In the groups CH_2NO_2 and CHNO_2 hydrogen is displaceable by metals, and hence bodies containing these groups dissolve in alkalis Such solutions yield, on addition of bromine, compounds containing the groups CHBrNO_2 and CBrNO_2 The compounds X CHBrNO_2 can further give rise to X CNaBrNO_2 and $\text{X CBr}_2\text{NO}_2$ —3 Primary fatty nitro compounds yield hydroxylamine on heating with HClAq at 140° , *eg* $\text{CH}_3\text{CH}_2\text{NO}_2 + \text{H}_2\text{O} = \text{CH}_3\text{COOH} + \text{H}_2\text{NOH}$ (V Meyer & Locker, *A* 180, 163)—4 Nitrous acid converts primary fatty nitro compounds into nitrolic acids, containing the group $\text{CH}(\text{NO})(\text{NO}_2)$ or $\text{C}(\text{NOH})(\text{NO}_2)$, which form red solutions with alkalis Nitrous acid converts secondary fatty nitro compounds into nitroles, containing the group $\text{C}(\text{NO})(\text{NO}_2)$, which are blue when in the liquid state or in solution—5 Aromatic nitro compounds may be reduced by heating with halogen acids Thus nitro benzene is reduced to aniline by heating with HIAq at 104° , by HBrAq at 185° , and by HClAq at 245° (Baumhauser, *A Suppl* 7, 212)—6 Boiling aqueous alkalis can in some cases displace NO_2 by hydroxyl In this way *o*-di-nitro benzene is converted into *o* nitro phenol—7 When aromatic compounds containing two or more nitroxyls in one benzene nucleus are treated with alcoholic NH_3 and H_2S it is usual for one nitroxyl only to be reduced to amidogen—8 Aromatic di and tri nitro compounds frequently form molecular compounds with one another and with aromatic hydrocarbons—9 The acetyl derivatives of nitrated aromatic amines, in which the NO_2 group is in the *o*- or *p*-position to the NHAc group (*eg* acetyl *o* and *p* nitro aniline, diacetyl nitro *p* phenylene diamine, diacetyl nitro *p* naphthylene diamine, acetyl *o* nitro β naphthylamine, &c.), are readily soluble in cold aqueous KOH (1.2) with a deep-yellow colour On standing saponification takes place, and the nitro compound crystallises out When the NO_2 group is in the *m*-position to the NHAc group, *eg* acetyl *m* nitraniline, acetyl *m* nitro *p* toluidine, &c., the substance is not soluble in aqueous KOH (Kleemann, *B* 19, 836)

NITRO-*o*-COUMARIC ACID *Methyl derivative* $\text{C}_6\text{H}_5(\text{NO}_2)\text{CH}(\text{OMe})\text{CHCO}_2\text{H}$ [298°] Formed by heating $\text{C}_6\text{H}_5(\text{NO}_2)(\text{OMe})\text{CHO}$ with NaOAc and Ac_2O (Schnell, *B* 17, 1833) White needles, sl sol cold water, *v* sol alcohol and ether— $-\text{CaA}_2$,— BaA_2 ,— AgA

Anhydride $\text{C}_6\text{H}_5(\text{NO}_2)\text{CH}(\text{CH})\text{COCH}(\text{CH})\text{CO}$ **Nitro-coumarin** [188°] Formed by dissolving coumarin in cold fuming HNO_3 (Delalande, *A Ch* [3] 6, 343, Bleibtreu, *A* 59, 191) Formed also by heating nitro salicylic aldehyde [125°] with Ac_2O and NaOAc (Taegge, *B* 20, 2110) Needles Oxidised by KMnO_4 to nitro *o*-oxy benzoic acid [228°] Reduced by FeSO_4 and NH_3 to amido coumarin [161°] Bromine vapour gives a di-bromide [271°]

Nitro-*o*-coumaric acid $\text{C}_6\text{H}_5(\text{NO}_2)\text{CH}(\text{OH})\text{CHCO}_2\text{H}$ [242°] Formed by heating its methyl derivative with NaOHAq Yellow crystals (from alcohol) Not converted into nitro-coumarin by boiling water or by HBrAq .

Methyl derivative

$C_6H_3(NO_2)(OMe)CHCHCO_2H$ [198°] Formed from the methyl derivative of (3,2,1)-nitro salicylic aldehyde by Perkin's reaction (Miller & Kinkelin, *B* 22, 1709) Prisms (from alcohol) With methyl iodide it gives the ether $C_6H_3(NO_2)(OMe)CHCHCO_2H$ [89°]

Nitro-o coumarinic acid

[3 2 1] $C_6H_3(NO_2)(OH)CHCHCO_2H$ [150°] Formed by dissolving nitro coumarin in boiling aqueous alkalis, and ppg the cold solution by HCl (Miller & Kinkelin, *B* 22, 1706) Yellow prisms (from warm alcohol) On warming with water or alcohol it changes to its anhydride, nitro coumarin Its salts are explosive — $Na_2C_6H_3NO_2$, — $BaC_6H_3NO_2$, 3:1aq — $AgC_6H_3NO_2$

Methyl derivative

$C_6H_3(NO_2)(OMe)CHCHCO_2H$ [186°] Tables
Methyl derivative of the methyl ether $MeC_6H_3NO_2$ [89°] Formed from the Ag salt and MeI Prisms Readily reconverted into the acid, even by aqueous Na_2CO_3

Anhydride $C_6H_3(NO_2) < \begin{smallmatrix} CH \\ CO \end{smallmatrix}$ *Nitro*

coumarin [191°] Formed from (3,2,1) nitro salicylic aldehyde (60 g), NaOAc (90 g), and Ac_2O (130 g) Prisms (from benzene), or needles (from alcohol)

Nitro m coumaric acid

[2 3 1] $C_6H_3(NO_2)(OH)CHCHCO_2H$ [218°] Formed from *m* amido cinnamic acid by nitration and displacement of NH by OH through the diazo reaction (Luff, *B* 22, 293) Needles (from water or alcohol)

Nitro m coumaric acid

[4 3 1] $C_6H_3(NO_2)(OH)CHCHCO_2H$ [248°] Formed by nitrating *m* coumaric acid (Luff, *B* 22, 296) Golden yellow needles (from alcohol)

Methyl derivative

$C_6H_3(NO_2)(OMe)CHCHCO_2H$ [218°] Formed from [4,3,1] $C_6H_3(NO_2)(OMe)CHO$ by Perkin's reaction (Landsberg, *D P J* 262, 139)

Methyl ether of the methyl derivative $C_6H_3(NO_2)(OMe)CHCHCO_2Me$ [143°] Plates (Rieche, *B* 22, 2359) Formed by nitration of $C_6H_3(OMe)CHCHCO_2Me$

Ethyl ether of the methyl derivative $C_6H_3(NO_2)(OMe)CHCHCO_2Et$ [163°] Needles (from alcohol) (Ulrich, *B* 18, 2572)

s-Nitro m coumaric acid

[5 3 1] $C_6H_3(NO_2)(OH)CHCHCO_2H$ Accompanies the (2,3,1) acid (*supra*) Crystals

Nitro m coumaric acid

[6 3 1] $C_6H_3(NO_2)(OH)CHCHCO_2H$ [216°] Formed from acetyl *m* amido cinnamic acid by nitration and displacement of $NHAc$ by hydroxyl (Luff, *B* 22, 292) Yellow powder

Nitro p coumaric acid

[3 4 1] $C_6H_3(NO_2)(OH)CHCHCO_2H$ [198°] Formed from its methyl derivative by heating with $HOAc$ saturated with HBr (Einhorn & Grabfield, *A* 243, 374) Yellow needles (from alcohol) Forms a dibromide [72°]

Ethyl ether EtA' [109°]*Methyl derivative*

$C_6H_3(NO_2)(OMe)CHCHCO_2H$ [140°] Formed from [3 4 1] $C_6H_3(NO_2)(OMe)CHO$, NaOAc, and Ac_2O White needles Yields a dibromide [178°], from which alkalis form the acid $C_6H_3(NO_2)(OMe)C_2HBr.CO_2H$ [205°]

Methyl ether of the methyl derivative $C_6H_3(NO_2)(OMe)CHCHCO_2Me$ [125°]*Ethyl ether of the methyl derivative* $C_6H_3(NO_2)(OMe)CHCHCO_2Et$ [100°]*Di-nitro-o-coumaric acid* *Methyl derivative* $C_6H_3(NO_2)_2(OMe)CHCHCO_2H$ [198°]

Formed by nitrating the methyl derivative of *o*-coumaric acid (Perkin, *O J* 89, 416) Orange-brown needles (from alcohol)

Di nitro-m-coumaric acid *Methyl derivative of the methyl ether*

$C_6H_3(NO_2)_2(OMe)CHCHCO_2Me$ [178°] Formed by nitration (Rieche, *B* 22, 2359) Yellowish needles (from alcohol) Yields on saponification $C_6H_3(NO_2)_2(OMe)CHCHCO_2H$, which decomposes at 215°

NITRO-o COUMARIC ALDEHYDE

$C_6H_3(NO_2)(OH)CHCHCHO$ [200°] Formed from $C_6H_3(NO_2)(OH)CHO$ [126°], aldehyde, and aqueous NaOH (Von Miller & Kinkelin, *B* 20, 1931, 22, 1716) Yellow needles Form a sodium derivative, crystallising in red tables, and a phenyl hydrazide [235°]

Nitro-o coumaric aldehyde

[3 2 1] $C_6H_3(NO_2)(OH)CHCHCHO$ [133°] Formed in the same way from the aldehyde $C_6H_3(NO_2)(OH)CHO$ [109°] (M & K) Golden needles, v sol alcohol Forms a sodium derivative, crystallising in red needles, a phenyl hydrazide [157°], and a methyl derivative [115°]

*NITRO COUMARIN v Anhydride of Nitro coumaric acid**NITRO-o-CRESOL* $C_6H_3NO_2$, 1:6

$C_6H_3Me(OH)(NO_2)$ [1 2 3] [70°] Obtained, together with the (1,2,5) isomeride, by nitrating *o* cresol (Hofmann & Miller, *B* 14, 567, Staedel, *A* 217, 50, 203, Rapp, *A* 224, 175, Hirsch, *B* 18, 1512) Formed also by boiling a dilute acid solution of *o* diazotoluene sulphate (1 mol) with HNO_3 (1 mol) (Nöling & Wild, *B* 18, 1839, Deninger, *J pr* [2] 40, 299) Yellow prisms (from dilute alcohol), insol water, v sol alcohol and ether, volatile with steam — $KC_6H_3NO_2$, garnet red trimetric tables — AgA' red needles

Methyl ether MeA' Oil*Ethyl ether EtA'* Oil

Nitro-o cresol $C_6H_3Me(OH)(NO_2)$ [1 2 4] [108°] Formed by boiling diazotised nitro *o*-toluidine [107°] with water (Nöling & Collin, *B* 17, 269) Yellow needles (from ligroin) Its K , Ag , and NH_4 salts form yellow crystals

Methyl ether [74°] (Witt, *B* 23, 8638)

Nitro-o-cresol $C_6H_3Me(OH)(NO_2)$ [1 2 5] [95°] (N & W), [80°–85°] (H) Formed from nitro *o*-toluidine [128°] by the diazo-reaction, or by merely boiling with conc $NaOHAq$ (Neville & Winther, *O J* 41, 423) Formed also by nitrating *o* cresol (Hirsch, *B* 18, 1512) Needles (from water), not volatile with steam When crystallised from water it melts at 30°–34°, but it melts at 95° after crystallisation from ether

Ethyl ether EtA' [71°]

Formed by nitrating the ethyl derivative of *o* cresol (Staedel, *A* 217, 155, 203, Kayser, *B* 15, 1133) Needles

Nitro-o-cresol $C_6H_3Me(OH)(NO_2)$ [1 2 6]

[148°] Formed from nitro *o*-toluidine [92°] by the diazo reaction (Ullmann, *B* 17, 1961) Yellow needles (from water) Has an intensely sweet taste

o-Nitro-m-cresol $C_6H_3Me(OH)(NO_2)$ [1 8 4]

[56°] Formed, together with the (1,3,6) iso-

meride from *m* cresol, HOAc, and HNO₃ (Staedel, *A* 217, 51, *A* 259, 223, Claus, *J pr* [2] 39, 63) Yellow monoclinic plates (from benzene), volatile with steam Gives di bromo nitro cresol [93°] Its K salt forms red plates
Ethyl ether EtA' [51°] White needles (from dilute alcohol)

s-Nitro-*m*-cresol C₆H₄Me(OH)(NO₂) [1 3 5] [91°] Formed from nitro *m* toluidine by the diazo-reaction (Neville a Winther, *C J* 41, 417) Yellow crystals, not volatile with steam Crystal lises from water in a hydrated condition, and then melts at 62°

Nitro *m* cresol C₆H₄Me(OH)(NO₂) [1 3 6] [129°] Formed, together with the (1,3,4) iso meride from *m* cresol, HNO₃, and HOAc below 0° (Staedel, *A* 259, 210, Claus, *J pr* [2] 39, 63) Formed also by oxidising nitroso *m* cresol with alkaline K₂FeCy₄ (Bertoni, *G* 12, 304) Colourless crystals, not volatile with steam Gives di-bromo nitro cresol [143°] Reduces to amido cresol [174°]—KA' 2aq yellow plates —NaA' 2aq

Ethyl ether EtA' [54°]

Nitro *p* cresol C₆H₄Me(OH)(NO₂) [1 4 3] [84°] Formed from acetyl *p* toluidine by nitrating and boiling the product with conc NaOHaq (Wagner, *B* 7, 537, Neville a Winther, *C J* 41, 426), by the action of nascent nitrous acid on *p* toluidine (Deninger, *J pr* [2] 40, 299), or by boiling *p* diazotoluene sulphate (1 mol) with HNO₃ (1 mol) (Nölting a Wild, *B* 18, 1339) Formed also by nitrating *p* cresol (Armstrong a Thorpe, *B* *A* 1875, 112, Hofmann a Miller, *B* 14, 572, Staedel, *A* 217, 54) Yellow crystals (from benzene), volatile with steam —NaA'—AgA'

Methyl ether MeA' (274°)

Ethyl ether EtA' (275°-285°)

Benzylether C₆H₅A' [54°] Formed from benzyl chloride and the Ag salt (Frische, *A* 224, 142)

o Nitro-*benzyl ether* [163°]

Nitro-*p* cresol C₆H₄Me(OH)(NO₂) [1 4 2] [77°] Formed from the corresponding nitro *p*-toluidine by the diazo reaction (Neville a Winther, *C J* 41, 422, Knecht, *A* 215, 87) Yellow needles (from ligroin)

Methyl ether MeA' (267°) Oil

Di nitro *o* cresol C₆H₃Me(OH)(NO₂)₂ [1 2 3 5] [86°] S (alcohol) 7.8 at 15° Formed by heating (1,2,5)-*o*-cresol sulphonic acid with dilute HNO₃ (Neville a Winther, *C J* 37, 631, 41, 422) Formed also by boiling *o* diazo toluene disulphonic acid with dilute HNO₃ (N a W), or from *o* diazo toluene nitrate and HNO₃ (Nölting a De Salis, *B* 14, 987, *A Ch* [6] 4, 105) Obtained also from the corresponding di nitro toluidine (N a S), and by nitrating *o* cresol and the (3,1,2)- and (5,1,2) nitro *o* cresols (Hirsch, *B* 13, 1512, Barr, *B* 21, 1543) Formed also from C₆H₄Me(OH)Br₂ and fuming HNO₃ (Claus, *J pr* [2] 38, 327) Yellow needles, slightly volatile with steam. Yellow dye —KA' 2aq. yellow crystals

Salts —BaA'₂—BaA'₂·3½aq —AgA'

Ethyl ether EtA' [51°] Obtained by nitrating C₆H₄Me(OEt) [1 2] (Staedel, *B* 14, 899, *A* 217, 153, 259, 219), or from the Ag salt and EtBr (N a S). Yields di nitro-*o* toluidine

[210°] on heating with alcoholic NH₃ at 130° (Van Romburgh, *R T O* 8, 397)

p-Nitro-*benzylether* C₆H₄(NO₂)CH₂A' [145°] Obtained by nitrating the benzyl ether of *o* cresol (Staedel) Needles

Di-nitro-*m*-cresol

C₆H₃Me(OH)(NO₂)₂ [1 3 4 5or2] [99°] Formed from di-nitro-amido cresol by elimination of NH₃ (Nietzki a Ruppert, *B* 23, 3479) Needles.

Ethyl ether EtA' [22°]

Di-nitro-*m*-cresol

C₆H₃Me(OH)(NO₂)₂ [1 3 4 6]

Ethyl ether EtA' [97°] Formed by nitrating ethyl-nitro-cresol [51°] (Staedel, *A* 259, 226)

Di-nitro *p* cresol C₆H₄Me(OH)(NO₂)₂ [1 4 3 5] [85°] A product of the action of nitrous acid on *p* toluidine nitrate, on amido toluic acid [167°], and on *p* toluidine disulphonic acid (Beilstein a Kreusler, *A* 144, 183, Martius a Wichelhaus, *Z* [2] 5, 440, *B* 2, 207, Richter, *A* 230, 323) Formed also by boiling di-nitro *p* toluidine with aqueous NaOH (Wagner, *B* 7, 536), by nitrating *p* cresol and (3,1,4) nitro *p* cresol (Armstrong a Field, *B* 6, 974, Frische, *A* 224, 139), or boiling diazo *p* toluene sulphonate with dilute HNO₃ (Neville a Winther, *C J* 37, 631) Yellow needles (from dilute alcohol)—NaA' red needles S 2.3 at 17°—KA' S 1.5 at 16° (M a W), 99 at 17° (Staedel) Used as a dye (gold-yellow) A sample of 'Victoria yellow' was found by Martius and Wichelhaus to consist of a salt of a dimnitrocresol [110°] Victoria yellow is poisonous (Weyl, *B* 20, 2835)—BaA'₂—AgA' S 29 at 17°

Methyl ether MeA' [122°]

Ethyl ether EtA' [73°] Formed by nitrating ethyl *p* cresol (Staedel, *A* 217, 161) When heated with alcoholic NH₃ it yields di nitro *p* toluidine [168°] (Romburgh, *R T O* 3, 405)

Benzoyl ether C₆H₅A' [109°]

p Nitro-*benzyl ether* [186°]

Di nitro *p* cresol C₆H₃Me(OH)(NO₂)₂ Obtained by the action of excess of nitrous acid on C₆H₄Me(OH)(NO₂) [1 4 2] (Knecht, *A* 215, 90) Yellow needles (from water) Does not melt when heated but forms a violet sublimate

Tri nitro *o* cresol C₆H₂Me(OH)(NO₂)₃ [102°] Formed by heating nitro *o* diazo toluene nitrate with nitric acid (S G 133) (Nölting a Collin, *B* 17, 270) Orange prisms (from acetone) Yields NaNO₂ when heated with NaOHaq With naphthalene it forms (C₆H₃N₃O₂)C₁₀H₆ [106°]

Tri nitro *m*-cresol

C₆H₃Me(OH)(NO₂)₃ [1 3 2 4 6] [106°] S 2.2 at 20°, 8 at 100° (Duclos) Formed by nitration of *m* cresol or its sulphonic acid (Duclos, *A* 109, 141, Nölting a De Salis, *B* 14, 987, 15, 1861, *A Ch* [6] 4, 118, Beilstein a Kellner, *A* 128, 165) Formed also by the action of HNO₃ on nitroso *m*-cresol (Wurster a Riedel, *B* 12, 1799), by heating nitro coccine acid with water at 180° (Luebermann a Dorp, *A* 163, 101, Kosta-necki a Niemientowski, *B* 18, 251), and by the action of cold conc HNO₃ on (2,6,3,4,1) di-iodo-toluquinone (Kehrmann, *J pr* [2] 39, 392) Yellow needles (from water) Forms with naphthalene a compound (C₆H₃N₃O₂)C₁₀H₆ [127°]—NH₃A' Converted by warm aqueous KOy into

purple crystals of potassium 'eresyl purpurate' $\text{KC}_6\text{H}_4\text{N}_2\text{O}_6$ (Sommaruga, *Z* 1870, 557) — KA' yellow needles — $\text{Pb}(\text{OH})\text{A}'$ — AgA' prisms

Ethyl ether EtA' [72°] (N & S), [75°] (Staedel, *A* 259, 221, 227) Converted by cold alcoholic NH_3 into tri nitro toluidine [126°]

NITRO ψ CRESOL SULPHONIC ACID

$\text{C}_6\text{H}_4\text{NSO}_3$, *ee* $\text{C}_6\text{H}_4\text{Me}(\text{OH})(\text{NO}_2)\text{SO}_3\text{H}$ Formed from (1,2,5) *o* toluidine sulphonic acid by dissolving in fuming HNO_3 and boiling the resulting nitro diazo toluene sulphonic anhydride with water (Hayduck, *A* 172, 218) Deliquescent — $\text{BaC}_6\text{H}_4\text{NSO}_3 \cdot 3\frac{1}{2}\text{aq}$ — $\text{Ba}(\text{C}_6\text{H}_4\text{NSO}_3)_2 \cdot 5\text{aq}$

Nitro ψ cresol sulphonic acid *Ethyl*

derivative $\text{C}_6\text{H}_4\text{Me}(\text{OH})(\text{NO}_2)\text{SO}_3\text{H}$ [142.5] Formed by heating nitro diazo toluene sulphonic acid with NaOEt (Foth, *A* 230, 306) Needles — BaA' , 4aq yellow plates

NITRO CRYPTOPINE ψ CRYPTOPINE

NITRO CUMENE $\text{C}_6\text{H}_4\text{NO}_2$, *ee* $\text{C}_6\text{H}_4\text{Pr}(\text{NO}_2)$ [—35°] Formed from cumene and fuming HNO_3 at 0° (Pospeckhoff, *J R* 18, 52, *Bl* [2] 45, 178, *cf* Cahours, *C R* 25, 552, 26, 315, Nicholson, *C J* 1, 2, Rutthausen, *J pr* 61, 79) Oil, volatile with steam

Nitro ψ cumene $\text{C}_6\text{H}_4\text{Me}_3(\text{NO}_2)$ [134.5] [20°] Formed by eliminating NH_3 from nitro ψ -cumidine (Edler, *B* 18, 629) Large thick prisms, volatile with steam

Nitro ψ cumene $\text{C}_6\text{H}_4\text{Me}_3(\text{NO}_2)$ [71°] (265°) Formed from ψ cumene and cold fuming HNO_3 (Schaper, *Z* [2] 3, 12, Fittig & Laubinger, *Z* [2] 4, 577) Colourless needles (from alcohol), volatile with steam Yields on oxidation $\text{C}_6\text{H}_4\text{Me}(\text{NO}_2)\text{CO}_2\text{H}$ [195°]

Tri nitro cumene $\text{C}_6\text{H}_3\text{N}_3\text{O}_6$, *ee* $\text{C}_6\text{H}_3\text{Pr}(\text{NO}_2)_3$ [124.6] [109°] Formed from cumene, HNO_3 , and H_2SO_4 (Fittig, *A* 149, 328) Needles, sl sol cold alcohol

Tri nitro ψ -cumene $\text{C}_6\text{Me}_3(\text{NO}_2)_3$ [185°] Formed by nitrating ψ cumene (Fittig & Laubinger, *A* 151, 261) Prisms (from benzene), almost insol boiling alcohol By passing hydrogen sulphide into its boiling ammoniacal alcoholic solution, nitro ψ cumidine sulphonic acid $\text{C}_6\text{Me}_3(\text{NO}_2)(\text{NH}_2)\text{SO}_3\text{H}$ [134.2.6.5] is formed (Mayer, *B* 19, 212, 20, 966)

NITRO ψ CUMENOL

$\text{C}_6\text{HMe}_3(\text{OH})(\text{NO}_2)$ [134.6.2] [48°] Obtained by evaporating an alcoholic solution of the nitrate to dryness, and distilling the residue with steam (Auwers, *B* 17, 2979, 18, 2658) Long reddish yellow crystals (from alcohol), insol hot water Reconverted into the nitrate by warming with dilute HNO_3

Nitrate $\text{C}_6\text{HMe}_3(\text{NO}_2)\text{ONO}$ [84°] Formed from ψ cumenol and cold fuming HNO_3 Trimetric tables or prisms, insol water, sl sol cold alcohol

Methyl ether $\text{C}_6\text{HMe}_3(\text{NO}_2)\text{OMe}$ [42°]

Nitro-cumenol $\text{C}_6\text{H}_3\text{Pr}(\text{NO}_2)(\text{OH})$ Oil, formed, together with an isomeride [86°], by nitrating $\text{C}_6\text{H}_3\text{Pr}(\text{OH})$ [12] (Filet, *G* 16, 120)

Di-nitro- ψ -cumenol

$\text{C}_6\text{Me}_3(\text{OH})(\text{NO}_2)_2$ [134.6.2.5] [112°] Formed by passing NH_3 into an alcoholic solution of the nitrate of nitro ψ cumenol [84°] Yellow crystals, insol water Forms a red solution in alkalis

NITRO CUMIDINE $\text{C}_6\text{H}_4\text{N}_2\text{O}_2$, *ee*

$\text{C}_6\text{H}_2(\text{C}_6\text{H}_4)(\text{NO}_2)(\text{NH}_2)$ [below 100°]. Formed

by reducing di nitro cumene with alcoholic ammonium sulphide (Cahours, *C R* 24, 557, 26, 315) Yellow scales Forms a crystalline benzoyl derivative — $\text{B}'\text{HCl aq}$ — $\text{B}'_2\text{H}_2\text{SO}_4\text{aq}$ needles

Nitro ψ cumidine

$\text{C}_6\text{HMe}_3(\text{NO}_2)(\text{NH}_2)$ [134.5.6] [47°] Formed from acetyl ψ cumidine by nitration and saponification (Edler, *B* 18, 629) Red needles (from dilute alcohol)

Acetyl derivative [194°] (E), [204°] (Auwers, *B* 18, 2661) Prisms (from alcohol)

Nitro- ψ -cumidine $\text{C}_6\text{HMe}_3(\text{NO}_2)(\text{NH}_2)$ [137°] Formed by treating tri nitro ψ -cumene with alcoholic ammonium sulphide (Fittig & Laubinger, *A* 151, 262) Yellow needles — $\text{B}'\text{HCl}$ — $\text{B}'_2\text{H}_2\text{SO}_4\text{aq}$

Nitro- ψ cumidine *Acetyl derivative* $\text{C}_6\text{HMe}_3(\text{NO}_2)\text{NHAc}$ [131°] Formed by nitration of acetyl ψ cumidine [112°] (Engel, *B* 18, 2231) Yellow needles

Di-nitro ψ cumidine $\text{C}_6\text{Me}_3(\text{NO}_2)_2(\text{NH}_2)$ [78°] Formed from acetyl ψ cumidine [112°] by nitration and saponification (Engel, *B* 18, 2232) Yellow needles

Acetyl derivative [204°] Needles

Di nitro ψ cumidine

$\text{C}_6\text{Me}_3(\text{NO}_2)_2(\text{NH}_2)$ [134.2.5.6] [183°] Obtained from acetyl ψ cumidine [164°] by nitration and saponification (Auwers, *B* 18, 2661) Orange needles (from alcohol)

Acetyl derivative [280°] Sl sol alcohol

NITRO ψ -CUMIDINE SULPHONIC ACID

$\text{C}_6\text{H}_4\text{N SO}_3$, *ee* $\text{C}_6\text{Me}_3(\text{NO}_2)(\text{NH}_2)(\text{SO}_3\text{H})$ [134.2.6.5] Formed by passing H_2S into a boiling solution of tri nitro ψ cumene in alcoholic NH_3 , and also by heating nitro ψ cumidine with ClSO_3H at 165° (Mayer, *B* 19, 2312, 20, 966) Colourless plates Melts, with decomposition at 240°–260° Forms an amorphous acetyl derivative $\text{C}_6\text{H}_4\text{AcN}_2\text{SO}_3$ [o 230°]

NITRO CUMINIC ACID $\text{C}_6\text{H}_4\text{NO}_2$, *ee* $\text{C}_6\text{H}_3\text{Pr}(\text{NO}_2)\text{CO}_2\text{H}$ [42.1] [99°] Formed by boiling nitro isopropyl cinnamic acid with CrO_3 in HOAc (Widman, *B* 19, 269) Tables or monoclinic prisms

Nitro-cuminic acid $\text{C}_6\text{H}_3\text{Pr}(\text{NO}_2)\text{CO}_2\text{H}$ [43.1] [159°] Obtained by nitration of cuminic acid (Gerhardt & Cahours, *A Ch* [3] 1, 73, 25, 36, Filet, *G* 11, 15, Alexejeff, *J R* 17, 112, *Bl* [3] 2, 727) Formed also by oxidation of its aldehyde (nitro cuminol) [54°] (Lippmann & Strecker, *B* 12, 77, Widman, *B* 15, 2547) and of nitro cumyl methyl ketone (Widman, *B* 21, 2232) Yellowish scales (from alcohol), turned red by sunlight, especially when dissolved in benzene (Alexejeff, *Bl* [2] 45, 178) — CaA' , — PbA' , — AgA'

Ethyl ether EtA' (290°) Oil

Nitrile $\text{C}_6\text{H}_3\text{Pr}(\text{NO}_2)\text{CN}$ [71°] Formed by nitrating cumonitrile (Czumpelik, *B* 2, 183)

Nitro ψ cuminic acid

$\text{C}_6\text{H}_3\text{Pr}(\text{NO}_2)\text{CO}_2\text{H}$ [43.1] [113°] Formed by oxidation of nitro ψ cumyl methyl ketone by KMnO_4 (Widman, *B* 21, 2231) Formed also by nitration of ψ -cuminic acid (Korner, *A* 216, 230) Colourless needles (from hot water), turned brown by light — BaA' , 4aq — SrA' , 5aq

Methyl ether MeA' [64°] Crystals (from alcohol) (Abenius, *J pr* [2] 40, 438).

Nitro-*n*-cuminic acid $C_9H_7Pr(NO_2)CO_2H$ [4 2 1] [157°] Formed by oxidation of nitro-propyl cinnamic acid (Widman, *B* 19, 276) Tables (from dilute alcohol), sl sol water

Di-nitro-cuminic acid $C_9H_5Pr(NO_2)_2CO_2H$ [221°] Formed by nitration of cuminic acid (Cahours, *A* 69, 243, Lippmann a Strecker, *B* 12, 78) Reddish crystals— BaA'_2 — CaA'_2 — AgA'_2 aq (Kraut, *C O* 1859, 85)

Ethyl ether EtA' [77 5°]

Amide $C_9H_5(NO_2)_2CONH_2$ Crystals

Di-nitro- ψ -cuminic acid $C_9Me_2(NO_2)_2CO_2H$ [1 8 4 2 5 6] [205°] *Di-nitro-durylic acid* Formed by nitrating ψ -cuminic acid (Gissmann, *A* 216, 207, Nef, *A* 237, 8) Prisms (from dilute alcohol)— CaA'_2 3aq— BaA'_2 3aq

NITRO CUMINIC ALDEHYDE $C_{10}H_{11}NO_3$ \pm $C_9H_7Pr(NO_2)CHO$ [4 2 1] *Nitro-cuminol* Formed by oxidising nitro isopropyl cinnamic acid with $KMnO_4$ (Einhorn a Hess, *B* 17, 2019) Oil, volatile with steam With acetone and $NaOH$ aq it produces di-isopropyl indigo

Nitro cuminic aldehyde $C_9H_7Pr(NO_2)CHO$ [4 3 1] [54°] Formed by nitration of cuminic aldehyde (Lippmann a Strecker, *B* 12, 76, Widman, *B* 15, 166) Trichino prisms Com bines with bisulphites

NITRO- ψ -CUMOQUINONE $C_9H_5NO_4$ \pm $C_9Me_2O_2(NO_2)$ [1 3 4 2 5 6] [113°] Formed by heating ψ -cumoquinone carboxylic acid with HNO_3 (S G 14) at 100° for half an hour (Nef, *C J* 53, 428, *A* 237, 17) Yellow plates, which may be sublimed Heated with alcoholic SO_2 in sealed tubes it yields $C_9Me_2(OH)(NO_2)$ [106°]

NITRO CUMYL ACRYLIC ACID ν **NITRO-PROPYL CINNAMIC ACID**

NITRO CUMYL METHYL KETONE $C_{11}H_{15}NO_3$ \pm [1 2 4] $C_9H_7Pr(NO_2)COCH_3$ *Nitro-aceto cumene* [49°] Formed by nitrating cumyl methyl ketone in the cold (Widman, *B* 21, 2227) Prisms, ν sol benzene, sl sol hgroun

Oxim $C_9H_7Pr(NO_2)C(OH)CH_3$ [117°]

Phenyl-hydrazide [138°]

Nitro-n-cumyl methyl ketone

[1 2 4] $C_9H_7Pr(NO_2)COCH_3$ Formed by nitrating *n*-cumyl methyl ketone (W) Oil

Oxim $C_9H_{10}(NO_2)C(OH)CH_3$ [86°]

Phenyl hydrazide [139°]

NITRO-*n*-CUMYL-PROPIONIC ACID

$C_9H_7Pr(NO_2)CH_2CH_2CO_2H$ [4 3 1] [99°] Formed by nitration (Widman, *B* 19, 2776) Crystals (from dilute $HOAc$)

NITRO-CYME NE $C_{10}H_{13}NO_2$ \pm $C_9H_7MePr(NO_2)$ [1 4 2] SG^{12} 1 085 Formed by nitrating cymene (Barlow, *A* 98, 245, Landolph, *B* 6, 987, Fittica, *A* 172, 314, Widman, *B* 19, 584, Söderbaum a Widman, *B* 21, 2126) Yellow oil Oxidised by $KMnO_4$ to nitro oxy-isopropyl benzoic acid and terephthalic acid

Nitro-isocymene $C_9H_7MePr(NO_2)$ [1 3 α] Formed from *m*-isocymene and fuming HNO_3 (Kelbe a Warth, *A* 221, 161) Oil, volatile with steam, but decomposed on distillation Yields nitro toluol acid [214°] on oxidation

Di-nitro cymene $C_9H_7MePr(NO_2)_2$ [54°] Formed by nitrating cymene (Kraut, *A* 92, 70) Got also from di-nitro amido cymene (Mazzara, *G* 19, 160) Iridescent tablets (from alcohol)

Di-nitro-cymene [78°] Formed from di-nitroso cymene [72°] and HNO_3 (S G 135)

(Kehrmann a Messinger, *B* 23, 3562) Crystals, ν sol alcohol

Di-nitro-cymene $C_{10}H_{13}(NO_2)_2$ SG^{12} 1 206 Formed by nitrating cymene from pyrochotis oil (Landolph) Oil, volatile with steam

Di-nitro cymene $C_{10}H_{13}(NO_2)_2$ [250°] Got from a coal tar cymene (Rommier, *Bl* [2] 19, 434)

Tri-nitro-cymene $C_9HMePr(NO_2)_3$ [119°] Formed by nitrating cymene (from camphor) (Fitting, *A* 145, 142) Thin plates

Tri-nitro isocymene $C_9HMePr(NO_2)_3$ [73°] Formed by nitration of *m*-isocymene (Kelbe, *A* 210, 54) Yellow leaflets, smelling like musk

NITRO CYME NE SULPHONIC ACID $C_9H_7MePr(NO_2)SO_3H$ [1 4 6 2] Formed from cymene by sulphonation and nitration (Errera, *G* 19, 533)— BaA'_2 aq— MgA'_2 5aq

Amide [139°] Scales

Nitro-cymene disulphonic acid $C_{10}H_{13}NS_2O_6$ \pm $C_9HMePr(NO_2)(SO_3H)_2$ Formed from nitro cymene and $ClSO_3H$ (Leone, *G* 11, 512) Not obtained pure— BaA' 3 $\frac{1}{2}$ aq— PbA' 4 $\frac{1}{2}$ aq needles

NITRO ISOCYIMIDINE $C_{10}H_{11}N_2O_3$ \pm $C_9H_7Me(C_2H_5)(NO_2)(NH_2)$ Formed by heating its phthalyl derivative with conc HCl aq at 180° (Kelbe a Warth, *A* 221, 176) Oil, volatile with steam

Benzoyl derivative [177°] Formed by nitrating benzoyl isocymidine (K a W) Needles (from alcohol)

Phthalyl derivative $C_{10}H_{13}(NO_2)N(C_2O_2C_6H_5)_2$ [167°] Formed by nitrating phthalyl isocymidine Needles

NITRODECOIC ACID $C_9H_5(NO_2)CO_2H$ A product of the action of boiling HNO_3 on the acids of cocoanut oil (Wirtz, *A* 104, 291)

NITRO DRACYLIC ACID is *p* **NITRO BENZOIC ACID**

NITRO-DULCITE ν **DULCITE**

NITRO-*c*-DURENE $C_{10}H_{13}NO_2$ \pm $C_9HMe_2(NO_2)$ [1 2 3 4 5] *Nitro prehnitene* [61°] (295° ν V) Formed by the action of HNO_3 on *c*-durene (Töhl, *B* 21, 905) Needles Yields on reduction *c*-duridine [70°]

Di-nitro *c*-durene $C_9Me_2(NO_2)_2$ [1 2 3 4 5 6] [178°] Formed from *c*-durene, HNO_3 , and H_2SO_4 in the cold (Jacobsen, *B* 19, 1214) and also from penta-methyl benzene and fuming HNO_3 (Gottschalk, *B* 20, 3287) Yellowish needles or prisms (from alcohol)

Di-nitro durene $C_9Me_2(NO_2)_2$ [1 2 4 5 3 6] [205°] Formed from durene and conc HNO_3 at 0° (Fitting a Jannasch, *Z* 1870, 162, Nef, *A* 237, 3, *C J* 53, 428) Colourless prisms, sl sol alcohol

Di-nitro isodurene $C_9Me_2(NO_2)_2$ [1 2 3 5 4 6] [156°] Prepared from isodurene, HNO_3 , and H_2SO_4 (Jacobsen, *B* 15, 1853) Prisms, sl sol cold alcohol

NITRO-DURENOL $C_9Me_2(NO_2)OH$ [180°] Formed by nitration of durenol with ordinary HNO_3 at 0° Yellow crystals ν e sol alcohol, nearly insol water Dissolves in alkalis with a dark-yellow colour (Jacobsen a Schnapauff, *B* 18, 2844)

NITRO-*c*-DURIDINE

$C_9Me_2(NO_2)(NH_2)$ [1 2 3 4 5 6] [131°] Formed by reducing di-nitro *c*-durene with alcoholic ammonium sulphide (Töhl, *B* 21, 904). Red

needles, sol alcohol Yields, on reduction, $C_6Me_3(NH_2)_2$ [140°]

NITRO-ERYTHRITE v ERYTHRITE TETRA-NITRATE

NITRO ETHANE $C_2H_5NO_2$, *ie* $CH_3CH_2NO_2$. Mol w 75 (114°) $\bar{S}G$ $\frac{1}{2}$ 10561, $\frac{25}{25}$ 10461 (Perkin, *C J* 55, 689) $\bar{M}M$ 2837 $\bar{S}V$ 803 (Schiff, Lossen, *A* 254, 73) $\bar{H}Fp$ 26,880 $\bar{H}Fv$ 25,140 (Thomsen, *Th*) Formed by adding EtI to cold silver nitrite and subsequently distilling from a water bath (*V Meyer*, *B* 5, 399, *A* 171, 1, 175, 88, Götting, *A* 243, 115, Kissel, *J R* 1882, 226) Formed also by distilling $KEtSO_3$ with $NaNO_2$, the yield being 6 p c of the theoretical (Lauterbach, *B* 11, 1225), and by the action of $AgNO_3$ on potassium chloro propionate (Kolotoff, *Bl* [2] 47, 169) Oil, with pleasant odour With alcoholic soda it gives an amorphous pp of C_2H_5ONa , a salt which is very soluble in water, forming a solution in which $HgCl_2$ ppts crystalline $C_2H_5(NO_2)HgCl$ The solution of sodium nitro ethane gives with $FeCl_3$ a blood red colour, with $CuSO_4$ a green solution, and with $AgNO_3$ a white pp rapidly turning black

Reactions —1 Iron and acetic acid reduce it to ethylamine —2 When mixed with potash and KNO_3 , and H_2SO_4 is slowly added, there is formed ethyl nitrolic acid $CH_3C(OH)NO_2$, [81°], the alkaline salts of which form deep red solutions (*V Meyer*, *B* 7, 425) —3 Fuming H_2SO_4 yields ethane s di sulphonic acid —4 $HClAq$ ($\bar{S}G$ 114) at 140° splits it up into hydroxylamine and $HOAc$ (*Meyer* & *Locher*, *A* 180, 163) —5 $NaOEt$ and EtI form only C_2H_5NO (168°) (*Götting*) $NaOMe$ and MeI form C_2H_5NO (c 155°) According to Socoloff (*J R* 20, 579) alcoholic soda forms C_2H_5NO (175°) and the presence of alkyl iodides does not affect the product —6 $ZnEt_2$, followed by water forms di ethyl hydroxylamine and other products (*Kissel*, *J R* 1887, 109) —7 Benzoyl chloride forms di benzoyl hydroxylamine and di acetyl hydroxyl amine (*Kissel*, *J R* 1882, 40)

Constitution —The constitution of nitro-ethane has been discussed by *Victor Meyer* (*B* 5, 404, 8, 30, *A* 244, 222), *Geuther* (*B* 7, 1620), *Alexejeff* (*Bl* [2] 46, 266), *Socoloff* (*Bl* [2] 47, 166), *Bevad* (*J R* 20, 125), and others

Di nitro methane $CH_2CH(NO_2)_2$ (186° cor) $\bar{S}G$ $\frac{25}{25}$ 13503 Formed by the action of KNO_2 and alcoholic potash on bromo nitro ethane (*Ter Meer*, *A* 181, 1) Formed also by the action of conc HNO_3 on di ethyl ketone and on methyl acetoacetic ether (*Chancel*, *Bl* [2] 31, 504, *C R* 96, 1466) Oil, with sweet taste Reduced by tin and $HClAq$ to hydroxylamine, NH_3 , and $HOAc$ Reduced by sodium amalgam to ethyl azaurolic acid — $CH_2CK(NO_2)_2$ yellow monoclinic crystals, m sol cold water, insol alcohol Explodes when struck Its aqueous solution gives a reddish-brown pp with $FeCl_3$, a pale blue pp with $CuSO_4$, and a light-brown pp with $HgCl_2$ — AgA lustrous yellow plates

Tri - nitro - ethane (?) $CH_3C(NO_2)_3$ [55°]. Formed from methyl-malonic acid and HNO_3 (*Franchimont*, *R T C* 5, 281) Crystals

Tetra-nitro-ethane (?) *Potassium derivative* $C_2K_4(NO_2)_6$ Formed from di-bromo-tetra-nitro-ethane, potash, and ammonium sulphide (*Vilhers*, *C R* 97, 258, 98, 431). Crystals,

which decrepitate below 100° and detonate at 200°, or even when treated with dilute acids

NITRO ETHENYL-TRI-AMIDO BENZENE $C_6H_3N_3O_3$, *ie*

[1 2 3 4] $C_6H_2(NO_2)_3(NH_2) < \begin{smallmatrix} NH \\ N \end{smallmatrix} > CCH_3$ [295°-300°] Formed by heating di acetyl di nitro *p*-phenylene diamine with alcoholic NH_3 at 150° (*Nietzki* & *Hagenbach*, *B* 20, 331, cf *Biedermann* & *Ledoux*, *B* 7, 1532) Red needles

Nitro - di - ethenyl - tetra - amido - benzene $C_6H_2O_2N_4$, *ie*

[5 1 2 3 4] $C_6H_2(NO_2)_4 < \begin{smallmatrix} NH \\ N \end{smallmatrix} > CMe_2$ [276°]

Formed by nitration of di ethenyl tetra amido-benzene (*Nietzki* & *Hagenbach*, *B* 20, 331) Orange red needles (containing aq) By reduction it is reconverted into di ethenyl tetra amido-benzene — $B''H_2Cl_2$, $\frac{1}{3}$ aq long golden yellow needles

DI NITRO-DI-ETHENYL-TETRA AMIDO-DITOLYL

$OMe < \begin{smallmatrix} NH \\ N \end{smallmatrix} > C_6HMe(NO_2)C_6HMe(NO_2) < \begin{smallmatrix} NH \\ N \end{smallmatrix} > OMe$ [1 3 4]

[242°] Formed from acetyl di nitro toluidine by reduction with ammonium sulphide, and treatment of the resulting hydrazo-compound with boiling dilute $HClAq$ (*Bankievitch*, *B* 21, 2407) Prisms, v e sol alcohol — $B''H_2Cl_2$ — $B'2HNO_3$ [214°] Yellow needles

NITRO ETHENYL-PHENYLENE DIAMINE

$C_6H_3N_2O_2$, *ie* [1 3 4] $C_6H_2(NO_2) < \begin{smallmatrix} NH \\ N \end{smallmatrix} > CMe$

[216°] Formed by heating nitro *p*-phenylene diamine with Ac_2O at 190°, cooling, and boiling with dilute H_2SO_4 (*Heim*, *B* 21, 2307) Yellowish-brown needles (from water), v sol hot alcohol

NITRO-ETHENYL-TOLYLENE-DIAMINE

$C_6H_2Me(NO_2) < \begin{smallmatrix} N \\ NH \end{smallmatrix} > CMe$ [1 3 4] [246°].

Formed from acetyl *p*-toluidine by nitration and reduction (*Bankievitch*, *B* 21, 2402) Needles Yields on reduction ethenyl-tetra amido toluene [100°] — $B''H_2Cl_2$ — $B'HNO_3$ [207°] Yellowish plates

NITRO ETHENYL-TOLYLENE-DIAMINE

[185°] (*L*), [202°] (*N*) Formed by nitrating ethenyl tolylene *o* diamine (*Ladenburg*, *B* 8, 677, *Niementowski*, *B* 19, 723) Needles

NITRO-ETHYL-ALCOHOL C_2H_5NO , *ie*

$CH_2(NO_2)CH_2OH$ $\bar{S}G$ $\frac{194}{194}$ 11691 Formed from glycol iodhydrin and $AgNO_3$ (*Demuth* & *V Meyer*, *B* 21, 3529, *A* 256, 29) Yellowish liquid, sol water, decomposed on distillation. Reacts with diazo salts yielding azo- dyes (e.g. $C_6H_5N_2CH(NO_2)CH_2OH$) Nitrous acid converts it into methyl nitrolic and glycollic acids — $CH_2(NO_2)CH_2ONa$ white granular powder

NITRO-ETHYL-AMIDO-BENZOIC ACID

$C_6H_3N_2O_4$, *ie* [1 3 5] $C_6H_2(NO_2)(NHEt)CO_2H$ [208°] Formed from nitro amido benzoic acid and $EtBr$ (*Rollwage*, *B* 10, 1704) Yellow needles (from water) — BaA , $\frac{1}{2}$ aq red needles

Nitro di-ethyl-m amido-benzoic acid

$C_6H_2(NO_2)(NEt_2)CO_2H$ Monoclinic crystals; $\alpha b c = 893 \ 1 \ 1095$ $\beta = 74^\circ 57'$ (*Heintze*, *J* 1885, 1454, cf. *Lehmann*, *Dessert*, Göttingen, 1884)

NITRO-ETHYL-AMIDO-PHENOL *Nitrosamine of the ethyl ether* $C_6H_3N_2O_4$, *ie*

$C_6H_5(NO_2)(OEt)NEt(NO)$ Formed from $C_6H_5(OEt)NHEt$ and nitrous acid (Förster, *B* 21, 364) Yellowish prisms Does not form salts

***o*-NITRO TETRA ETHYL-DI-*p*-AMIDO TRI-PHENYL-METHANE** $C_{24}H_{23}N_5O_2$ ϵ $C_6H_5(NO_2)CH(C_6H_5NEt)_2$ [110°] Formed by heating *o*-nitro benzene aldehyde with di ethyl-aniline and dehydrated oxalic acid (Fischer a Schmidt, *B* 17, 1898) Orange triclinic prisms. ***p*-Nitro-tetra ethyl di *p*-amido tri-phenyl-methane** [113°] Obtained from *p*-nitro benzene aldehyde and di ethyl aniline (Kaeswurm, *B* 19, 744) Thick needles or monoclinic plates

NITRO ETHYLAMINE C_2H_5NHNO *Ethyl-nitramine* [3°] Formed from ethylamine by treatment with $ClCO_2Me$ and decomposition of the resulting $C_2H_5NHCO_2Me$ by ammonia (Franchimont a Klobbie, *R T C* 7, 356)

***o*-NITRO ETHYL-ANILINE** $C_6H_4(NO_2)NEt$ ϵ $C_6H_4(NO_2)NHEt$ [12°] Formed by heating *o*-nitro phenol with alcoholic ethylamine for 12 hours at 175° Formed also by heating the ethylene ether of *o* nitro phenol with alcoholic ethylamine at 140° (Hempel, *J pr* [2] 39, 199, 41, 162) Red oil, sol acids, but reppd by water Yields on reduction *o* phenylene ethyl diamine (249°) Nitrous acid converts it, in ethereal solution, into di nitro ethyl aniline [114°]

Nitrosamine $C_6H_4(NO_2)NEt(NO)$ [30°] Formed from the hydrochloride of the base and $NaNO_2$ in aqueous solution Yellow needles (from dilute alcohol or HOAc)

***m*-Nitro-ethyl aniline** $C_6H_4(NO_2)NHEt$ [13°] [60°] Formed by heating *m* nitro aniline (16 g) with $EtBr$ (14 g) and aqueous $NaOH$ (6 g) Formed also by adding HNO_3 (41.5 g of SG 1.39) to a cooled solution of ethyl aniline (50 g) in H_2SO_4 (1,000 g) (Nolting a Stricker, *B* 19, 546) Reddish yellow needles, volatile with steam With diazotised *p* bromo aniline it yields $C_6H_4BrN_2NEtC_6H_4NO_2$ [136°] (Meldola a Streatfield, *C J* 55, 429)

Nitrosamine $C_6H_4(NO_2)NEt(NO)$ [47°] **Acetyl derivative** $C_6H_4(NO_2)NEtAc$ [89°]

***p*-Nitro ethyl-aniline** $C_6H_4(NO_2)NHEt$ [14°] [95°] Obtained by nitration of acetyl ethyl aniline dissolved in H_2SO_4 (5 pts), the product being saponified (Weller, *B* 16, 31, Nölting a Collin, *B* 17, 267) Formed also by heating *p* nitro aniline with $EtBr$ and alcoholic potash at 110° (Schweitzer, *B* 19, 142) Yellow prisms with violet reflex (from alcohol) Somewhat volatile with steam With diazotised *p*-bromo aniline it yields $C_6H_4BrN_2NEtC_6H_4NO_2$ [125°]

Nitrosamine $C_6H_4(NO_2)NEt(NO)$ [120°] Yellow needles (from alcohol) (Meldola a Streatfield, *C J* 49, 61)

Acetyl derivative $C_6H_4(NO_2)NEtAc$ [118°]

Benzoyl derivative $C_6H_4(NO_2)NEtBz$ [98°] Needles, v sl sol hot water (Meldola a Salmon, *C J* 53, 774)

***m*-Nitro-di-ethyl-aniline** $C_6H_4(NO_2)NEt_2$ [13°] (289°) Formed, together with a small quantity of the *p*-isomeric, by nitration of di-ethyl-aniline dissolved in H_2SO_4 (20 pts) (Groll, *B* 19, 199) Obtained also by heating *m*-nitro-aniline (20 g) with EtI (46 g) and $NaOH$ (12 g.)

in alcoholic solution for 8 hours at 100° (Nölting a Stricker, *B* 19, 550) Dark yellow oil

***p* Nitro di ethyl aniline** $C_6H_4(NO_2)NEt_2$ [14°] [78°] Formed by oxidation of nitroso di ethyl-aniline with $KMnO_4$ and H_2SO_4 , and also by the action of nitrous acid on di ethyl amido benzene-azo di ethyl aniline (Lippmann a Fleissner, *B* 16, 1422, Groll) Yellow monoclinic needles with blue reflex— $B'H_2PtCl_4$ thin prisms

Di nitro ethyl aniline

$C_6H_4(NO_2)_2NHEt$ [4 2 1] [114°] Formed from bromo di nitro benzene and alcoholic ethylamine (Van Romburgh, *R T C* 2, 104) Formed also by boiling C_6H_5NEtAc with dilute HNO_3 (SG 1.029) (Norton a Allen, *B* 18, 1997), and by the action of nitrous acid on an ethereal solution of *o* nitro ethyl aniline (Hempel, *J pr* [2] 39, 199, 41, 168) Yellow needles (from alcohol) Decomposed by boiling conc $KOHAq$ into ethyl amine and di nitro phenol

Di-nitro-di-ethyl-aniline $C_6H_4(NO_2)_2NEt_2$ [4 2 1] [80°] Prepared by nitrating di ethyl aniline, and also by treating bromo di nitro benzene with diethylamine (Van Romburgh, *R T C* 2, 35, 8, 251) Yellow needles Decomposed by boiling aqueous KOH into di ethylamine and di nitro phenol When gently oxidised by CrO_3 it yields di nitro aniline [175°]

Tri-nitro-ethyl-aniline $C_6H_3(NO_2)_3NHEt$ *Ethylpicramide* [84°] Formed from chloro-tri nitro benzene (picryl chloride) and NH_4Et in alcohol (Van Romburgh, *R T C* 2, 107) Crystals (from alcohol) which turn brown in air **Tri nitro-di-ethyl aniline** $C_6H_3(NO_2)_3NEt_2$ [164°] Prepared by adding a hot alcoholic solution of $NHEt_2$ to (1,2,4,6) chloro-tri nitro benzene Orange crystals (from benzene) Decomposed by potash into picric acid and di-ethylamine

Tetra-nitro-ethyl-aniline

$C_6H_3(NO_2)_4NEt(NO_2)$ *Nitramine of tri nitro-ethyl aniline* [96°] Obtained by the action of HNO_3 and H_2SO_4 on ethyl aniline and on di ethyl aniline (Van Romburgh, *R T C* 2, 31, 114) Yellow plates (from alcohol) Decomposed by 10 pc aqueous Na_2CO_3 into picric acid and ethylamine Reduced by tin and $HClAq$ to tri amido phenol

NITRO-ETHYL-ANTHRONE $C_{14}H_9NO_2$ ϵ

$C_6H_5 \begin{matrix} CO \\ \diagup \\ CET(NO) \end{matrix} C_6H_5$ [102°] Formed, as a by-product, in the preparation of ethyl anthracene hydride nitrite by the action of HNO_3 on ethyl anthracene hydride dissolved in $HOAc$ (Liebermann a Landshoff, *B* 14, 474)

***o*-NITRO-ETHYL BENZENE** $C_6H_4(NO_2)C_6H_5$ [228°] SG 25.1 126 Formed, together with the *p*-isomeric, by dissolving ethyl benzene in HNO_3 (Beilstein a Kuhlberg, *A* 156, 206, *Z* [2] 5, 524) Oil

***p*-Nitro-ethyl-benzene** $C_6H_4(NO_2)C_6H_5$ (246°) SG 25.1 124 Oil

Nitro di-ethyl benzene $C_6H_4Et_2(NO_2)$ (155° at 23 mm) Formed from di ethyl-benzene and fuming HNO_3 at 0° (Voswinkel, *B* 22, 316) Oil, boils with partial decomposition at 280°–285°

Di-nitro-tetra-ethyl-benzene $C_4Et_4(NO_2)_2$ [115°] Pale yellow prisms (Galle, *B* 16, 1745)

Tri-nitro-di-ethyl-benzene $C_6H_3(NO_2)_3Et_2$ [62°] Yellow prisms (Voswinkel, *B* 21, 2680).

o-NITRO-ETHYL-BENZENE SULPHONIC ACID $C_6H_4NSO_3$, $\pm C_6H_4(NO_2)EtSO_3H$ Formed by sulphonation (Beilstein a Kuhlberg, *A* 156, 207) — BaA'_2 *S* 54 at 17.5°

p-Nitro-ethyl-benzene sulphonic acid — BaA'_2 4aq *S* 261 at 17.5° Needles

NITRO-p-ETHYL-BENZOIC ACID

$C_6H_4Et(NO_2)CO_2H$ [156°] Formed from *p* ethyl benzoic acid and cold fuming HNO_3 (Aschenbrandt, *B* 12, 1304, *A* 216, 220) Needles (from water) — NaA'_2 2aq — CaA'_2 2aq — BaA'_2 4aq — BaA'_2 4aq leaflets, sl sol water

p-NITRO- α -ETHYL BENZOYL-ACETIC ETHER $C_6H_4(NO_2)COCH_2CO_2Et$ [40°] Formed from $C_6H_4(NO_2)COCH_2NaCO_2Et$ and EtI (Perkin a Bellenot, *C J* 49, 451) Plates

DI-NITRO-ETHYLENE-UREA

$CO \langle N(NO_2)CH_2 \rangle$ [210°] Obtained from ethylene urea and HNO_3 (Franchumont a Klobbe, *R T C* 7, 17) Prisms On boiling with water it loses CO_2 and forms ethylene di nitramine $C_2H_4(NHNO_2)_2$ [174°]

DI-NITRO-ETHYLIC ACID $C_6H_4N_2O_4$ *Ethyl-nitramine*? Formed from $ZnEt_2$ and NO Prepared by passing nitric oxide into a benzene solution of $ZnEt_2NaEt$, obtained by adding sodium (12.7 g) to cold zinc ethyl (100 g) The product is successively treated with ether, alcohol, and water, zinc is pptd by CO_2 , the filtrate evaporated, and the sodium salt extracted by alcohol (Frankland a Graham, *C J* 37, 570, cf Frankland, *C J* 9, 89, Zuckschwerdt, *B* 7, 291, *A* 174, 302) The free acid is unstable Its salts yield ethylamine on reduction by sodium amalgam Alcoholic potash forms ethylamine and nitric acid (Zorn, *B* 15, 1008) — $NaC_6H_4N_2O_4$ — CaA'_2 3aq — BaA'_2 — MgA'_2 — ZnA'_2 — ZnA'_2 4aq — CuA'_2 3aq flat dark blue needles (from alcohol) — AgA'_2 — AgA'_2AgNO_3

NITRO-o-ETHYL-PHENOL $C_6H_4Et(NO_2)OH$ (212°–215°) Formed in small quantity by the action of nitrous acid on $C_6H_4EtNH_2$ (Suida a Plohn, *Sitz W* [2] 81, 245) — BaA'_2 4aq orange plates

Di-nitro-o-ethyl-phenol $C_6H_4Et(NO_2)_2OH$ Formed from *o* ethyl phenol and cold HNO_3 (*S* a *P*) Heavy oil — BaA'_2 (at 100°) Yellow plates (from alcohol)

NITRO-p-ETHYL-ISOPROPYL-BENZENE $C_6H_4EtPrNO_2$ (265°) Obtained from [4.1] C_6H_4EtPr and HNO_3 (Von der Becke, *B* 23, 3194)

DI-NITRO-(β)-ETHYL-THIOPHENE $C_6H_4(C_2H_5)(NO_2)_2S$ Formed by nitration of (β) ethyl thiophene by passing air charged with its vapour into fuming HNO_3 (Bonz, *B* 18, 552) Crystalline solid With alcoholic KOH it yields a blue colouration, becoming red on longer exposure to the air, or by addition of more KOH

NITRO-o-ETHYL-TOLUENE

$C_6H_4MeEt(NO_2)$ Oil (Claus a Pieszcsek, *B* 19, 3087)

Di-nitro o-ethyl-toluene $C_6H_4MeEt(NO_2)_2$ Oil, not solid at 0°

Di-nitro-p-ethyl-toluene [52°] Obtained, with an oily isomeride, by nitrating *p* ethyl-toluene (Jannasch a Dieckmann, *B* 7, 1513)

Tri nitro-p ethyl-toluene $C_6H_4MeEt(NO_2)_3$ [92°] Obtained by nitration (Glinzer a Fittig, *A* 186, 303) Prisms (from alcohol)

Nitro-ethyl-p toluidine

$C_6H_4Me(NO_2)NH_2$ [4.3.1] [48°] Formed from ethyl *p* toluidine (1 pt), H_2SO_4 (20 pts), and HNO_3 (Nöling a Stricker, *B* 19, 549). Flat red prisms, v sol alcohol

Nitro-ethyl-p toluidine

$C_6H_4Me(NO_2)NH_2$ [4.2.1] [59°] Formed by heating nitro-*p* toluidine with EtI (Gattermann, *B* 18, 1483, Niementowski, *B* 20, 1883) Red crystals (from alcohol), v sol ether

Acetyl derivative (245°–250° at 150 mm)

Di-nitro-ethyl-o-toluidine *Nitramine* [1.3.5.6] $C_6H_4Me(NO_2)_2NH_2$ [72°] Formed, in small quantity, by the action of HNO_3 on di-ethyl *o* toluidine (Van Romburgh, *R T C* 3, 402) Yellow crystals (from alcohol)

Di nitro ethyl p toluidine

$C_6H_4Me(NO_2)_2NH_2$ [1.3.5.4] [126°] Formed by nitration of nitro ethyl *p* toluidine (Gattermann, *B* 18, 1485) Orange crystals

Nitrosamine $C_6H_4(NO_2)_2NH_2(NO)$ [78°]

Nitramine $C_6H_4Me(NO_2)_2NH_2(NO_2)$ [116°] (*R*, *G*), [106°] (*N* a *L*) Formed from di ethyl *p* toluidine and fuming HNO_3 (Van Romburgh, *R T C* 3, 408) Formed also from $C_6H_4Me(NH_2)Ac$ and dilute (10 p c) HNO_3 (Norton a Livermore, *B* 20, 2271) Converted by boiling $NaOH$ aq into di nitro *p* cresol [83°]

TRI-NITRO-ETHYL-o-XYLENE

$C_6Me_2Et(NO_2)_3$ [121°] Needles (from alcohol) (Fittig a Ernst, *A* 139, 193, Stahl, *B* 23, 992)

Tri-nitro-ethyl-m-xylene [127°] Formed from ethyl *m* xylene, HNO_3 , and H_2SO_4 (Stahl) White needles, m sol alcohol

Tri-nitro s ethyl m xylene [138°] Needles, v sl sol alcohol (Jacobsen, *B* 7, 1434)

Tri-nitro-ethyl-p xylene [129°] Prisms (from hot alcohol) (Jacobsen, *B* 19, 2516)

NITRO EUGENOL $C_{10}H_7NO_2$, \pm

$C_6H_4(C_2H_5)(NO_2)(OMe)(OH)$ [1.5.3.4] [44°] Formed by nitration of eugenol (Weselsky a Benedikt, *M* 3, 387) Trichmic crystals, sl sol water Volatile with steam

Acetyl derivative [61°] Tables

NITRO EUXANTHIC ACID v **EUXANTHIC ACID**

NITRO PSEUDO-FLAVENOL v FLAVENOL

NITRO-FLUORANTHENE v FLUORANTHENE

p NITRO-FLUORENE $C_{13}H_9NO_2$, \pm

$CH_2 \langle C_6H_4(NO_2) \rangle$ [151°] (*H*), [154°] (*S*) Formed from fluorene, $HOAc$, and HNO_3 (Hodgkinson, *C J Proc* 1, 37, Strasburger, *B* 17, 107) Prisms, v sl sol alcohol

Di nitro fluorene $CH_2 \langle C_6H_4(NO_2)_2 \rangle$ [201°]

Formed by nitrating fluorene (Fittig a Schmitz, *A* 193, 134) Needles (from $HOAc$)

DI NITRO FLUORESCÉIN $C_{20}H_9(NO_2)_2O_2$

Formed from fluorescein (1 pt), H_2SO_4 (20 pts), and HNO_3 (2 pts) at 0° (Baeyer, *A* 183, 1) Amorphous yellow powder

Di acetyl derivative Pale-yellow needles (from alcohol) On boiling for some minutes with dilute (15 p c) KOH it forms a blue solution

Tetra-nitro-fluorescein $C_{20}H_9(NO_2)_4O_2$

Formed from fluorescein (1 pt) and fuming HNO_3 (5 pts) Colourless crystals (from $HOAc$) Its alcoholic solution is yellowish red, and, on

adding an acid, becomes first reddish violet and then colourless

NITROFORM *v.* Tri-NITRO-METHANE

NITRO-FURFURYL-ETHYLENE

$C_4H_2O_2CH_2CH(NO_2)$ [184°] From furfuraldehyde and an alkaline solution of nitro ethane (P) Yellow prisms

Nitro-furfuryl-nitro-ethylene

$C_4H_2(NO_2)O_2CH_2CH(NO_2)$ [144°] Yellow felted needles Formed by nitration of furfuryl nitro ethylene It is oxidised by CrO_3 to nitro pyromucic acid

Dibromide [111°], yellow prisms (Priebs, B 18, 1862)

NITROGEN N (Azote) At w 14 01 Mol w 28 02 Boils at -194.4° (Olszewski, W 31, 58) According to Sarrau (*C R* 94, 639, 718, 845) the critical temperature of N is -128.8° , and the critical pressure is 42.1 atmos SG (gas) 97247 (air=1) SG (liquid) 885 (water=1) at b.p. (O, *l.c.*, cf Wroblewski, C R 102, 1011). SG (gas) at 3000 atmos (water=1) 823 (Amagat, C R 107, 522) VD 14 (von Jolly, W 6, 536) SH p (equal vol of water=1) 2868, (equal volume of air=1) 2377 (Regnault, Acad 26, 302) CE 0036677 (von Jolly, P Jubeldd 82) S 01843 at 4° , 01751 at 6.2° , 0152 at 12.6° , 01436 at 17.7° , 01392 at 23.7° , the absorption coefficient

= 020346 - 00053887t + 000011156t² (Bunsen, Gasom Methoden [2nd ed] 209) S (alcohol) 12561 at 1° , 12384 at 3° , 12241 at 11.2° , 12148 at 14.6° , 12053 at 19° , 11973 at 23.8° , absorption coefficient

= 126338 - 000418t + 000006t² (Carius, A 94, 136, Bunsen, Gasom Methoden [2nd ed] 209)

Refraction-equivalent $\left(\frac{\mu-1}{d} \text{ At } w\right) = 4.1 \text{ to } 5.3$

(Gladstone, Pr 18, 49) Mean value of μ for white light = 1.0003019, dispersion power = 2086 (Croullebois, A Ch [4] 26, 236, v also Mascart, P 153, 149) MM N v c 114, Nⁱⁱⁱ c 611 (Perkin, C J 55, 736) TC (air=1) 98 (Nair, P 142, 123), 993 (Plank, Carl Rep 13, 164) HC [NⁱO] = -17,740, [NⁱO] = -21,575, [NⁱO] = -2,005, [NⁱOⁱAg] = -6,820, [NⁱOⁱAg] = -29,820 (Th 2, 198), [NⁱO] = -22,200, [NⁱO] = -1,200, [NⁱOⁱAg] = -8,400, [NⁱOⁱAg] = -28,600 (Berthelot, A Ch [5] 20, 255) Coefficient of compressibility 750-1,000 atmos 000407, 1000-1500 atmos 000265, 1500-2000 atmos 00017, 2000-2500 atmos 000123, 2500-3000 atmos 000091 (Amagat, C R 107, 522) Transpiration coefficient (O=1) 873 (Meyer a Springmuhl, P 148, 526), 885 (von Obermayer, W A B 73 [2nd part], 433) Friction coefficient at 0° = 000184 (M a S, *l.c.*), 0001659 (von O, *l.c.*) The spectrum of N varies much, there are two distinct spectra, known as the elementary line spectrum and the band-spectrum (for measurements of lines, v B A 1884 429, also Ames, P M [5] 30, 48)

Occurrence—In the atmosphere, forming c^{ths} by volume In the fluid-cavities of some specimens of rock crystals (Davy, T 1822 367) In the air-bladders of fishes, and in other cavities of the bodies of animals and vegetables In the gases from some fumaroles In certain wells (v L Smith, Am S [2] 12, 866) Probably in the sun (Young, Am S [3] 4, 856, Draper,

ibid [3] 14, 89) Compounds of N occur in very large quantities throughout the animal, vegetable, and mineral kingdom

In 1772 Rutherford (*De aere mephitico*, Edinburgh, 1772) showed that the expired breath of animals contained a gas which extinguished flame, but which was not carbonic acid, as it was not absorbed by potash A little later Lavoisier proved that this gas was present in air As this gas did not support animal life, Lavoisier called it *azote* (*à* and *wt*) Chaptal afterwards gave it the name *nitrogen*, because it was present in nitre

Formation—1 From air, by removing CO_2 by KOHAq, moisture and NH_3 by conc H_2SO_4 , and O by passing over red hot Cu (*v. Preparation*, No 1)—2 By passing air through a mixture of sawdust and Fe sulphide (obtained by saturating ppd Fe_2O_3 with H_2S), and then through alkaline pyrogallate solution, and finally through conc H_2SO_4 When the process is completed, passage of H_2S re forms Fe sulphide, which may be used again—3 By burning P in an inclosed quantity of air, over water, and allowing the P_2O_5 formed to dissolve in the water—4 By passing air through conc NH_4Aq , and then sending the mixture of air and NH_3 over Cu heated to redness, the CuO formed is reduced by the NH_3 (Lupton, C N 33, 90)—5 Berthelot (*Bl* [2] 13, 314) partly covers with NH_4Aq c 200 grams pure Cu turnings in a 10-14 litre flask, closes the flask by a cork carrying a safety funnel tube, and a delivery tube which is stopped by a caoutchouc cap, and shakes from time to time The O is thus completely removed from the air in the flask, the N may be obtained by pouring into the flask water previously freed from O by shaking with NH_4Aq and Cu, the gas should be passed through KOHAq, conc H_2SO_4 , and then through $CrCl_3Aq$ (B, *Bl* [3] 2, 643)—6 By shaking $FeO.H_2O$ or $MnO.H_2O$ with air, the hydroxides are obtained by adding NaOHAq to conc $FeSO_4Aq$ or $MnSO_4Aq$, and at once stopping the ingress of air—7 By placing pyrogallic acid in a flask, adding NaOHAq, corking, and shaking for some time (v Liebig, A 77, 107)—8 By passing over Pt black a mixture of 100 vols air (from which CO_2 has been removed) with 42 vols H (Dumoulin, *L'Institut*, 1851 11)—9 By passing Cl into rather dilute NH_4Aq , keeping the NH_3 always in large excess, $8NH_3 + 3Cl_2 = 6NH_4Cl + N_2$ (the experiment is dangerous, as NCl_3 may be formed and explode)—10 By warming conc NH_4NO_3Aq , or more easily conc KNO_3Aq mixed with 3 vols conc NH_4ClAq , whereby KCl and NH_4NO_3 are formed and the NH_4NO_3 is decomposed ($NH_4NO_3 = 2H_2O + N_2$, Cornwinder, A 72, 225) Addition of conc $K_2Cr_2O_7Aq$ oxidises N oxides (which are generally produced) to HNO_3 (*v. Preparation*, No 2) Loew (B 23, 3018) has found that a 4-5 p c solution of NH_4NO_3 is decomposed at the ordinary temperature by Pt black, with evolution of N—11 By heating in a retort an intimate mixture of equal parts dry NH_4Cl and $K_2Cr_2O_7$, and passing the gas through $FeSO_4Aq$ to absorb NO which is generally formed The chief change is represented approximately thus $2NH_4Cl + K_2Cr_2O_7 = 2KCl + 4H_2O + Cr_2O_3 + N_2$ —12 By heating powdered $(NH_4)_2Cr_2O_7$, which decomposes to Cr_2O_3 , H_2O , and N.—13. By

adding conc NH_4Cl aq to a strongly alkaline conc solution of NaBrO (prepared by adding Br to cold NaOH aq), N is evolved rapidly ($3\text{NaBrO} + 2\text{NH}_4\text{Cl} = 3\text{NaBr} + 3\text{H}_2\text{O} + 2\text{HCl} + \text{N}_2$). Solution of bleaching powder may be used, but there is danger of formation and explosion of NCl_3 . — 14 By heating a mixture of NH_4NO_3 and MnO_2 to c 200° (not over 215°) ($4\text{NH}_4\text{NO}_3 + \text{MnO}_2 = \text{Mn}(\text{NO}_3)_2 + 8\text{H}_2\text{O} + 3\text{N}_2$, Gatehouse, *B* 10, 1007) — 15 N is evolved in the reactions of several metals with HNO_3 , the gas evolved by the action of Zn on HNO_3 aq in presence of much NH_4NO_3 consists of c 90 p c N , with N_2O and NO (Acworth, *C J* 28, 839).

Preparation. — 1 A very slow stream of air is passed from a gasholder through U tubes containing slightly moistened KOH , to absorb CO_2 , then through U tubes containing CaCl_2 , to absorb H_2O , and then through a long hard glass tube, containing Cu turnings, or better Cu obtained by reducing CuO in H (Carius, *A* 94, 126), and heated to bright redness in a furnace, the gas which issues is allowed to bubble through a solution of CrCl_3 to remove any traces of O which may remain, and is then dried by passing through CaCl_2 in several U tubes, and then over P_2O_5 . Before the air stream is started, the tube containing the Cu should be heated and a stream of H passed through it, to remove traces of CuO (this is not necessary, of course, if the Cu has been prepared by reducing CuO by H), after cooling, one end of the tube should be sealed and the other connected with a Sprengel pump, and the Cu should be heated in a vacuum for some time. If this precaution is not taken, the N will contain H (von Jolly, *W* 6, 536). The solution of CrCl_3 is prepared before use by digesting CrCl_3 aq with scrap Zn and HCl aq until a clear blue liquid is obtained, which is poured into Na acetate solution, in an atmosphere of CO_2 , the red pp of chromium acetate is washed with H_2O containing CO_2 , and is then placed in a flask closed by a cork with entrance tube (to be attached to the N apparatus), exit tube, and a funnel through which HCl aq is dropped on to the acetate, which is thereby changed to CrCl_3 , the whole of the acetate is not dissolved, to avoid free HCl , the N apparatus is at once attached to the flask (O von der Pfordten, *A* 228, 112). — 2 Solid NH_4Cl is added to an almost saturated cold solution of NaNO_2 , when no more NH_4Cl dissolves the liquid is poured into a capacious flask, cold conc $\text{K}_2\text{Cr}_2\text{O}_7$ aq is added, about 1 pt $\text{K}_2\text{Cr}_2\text{O}_7$, for each 1 pt NaNO_2 used, and the mixture is gently warmed. The $\text{K}_2\text{Cr}_2\text{O}_7$ oxidises any oxides of N to HNO_3 (*v* Gibbs, *B* 10, 1387). It is advisable to pass the N through KOH aq to absorb any traces of Cl compounds coming from impurities in the salts used (Gibbs, *l.c.*) — 3 A solution of 1 pt $\text{K}_2\text{Cr}_2\text{O}_7$, 1 pt NH_4NO_3 , and 1 pt NaNO_2 in 3 pts water is warmed in a fair sized flask (Böttger, *Jahr des phys Vereins zu Frankfurt*, 1876-77 24).

Properties. — A tasteless, colourless, odourless gas, which does not burn, nor support combustion, nor form a pp with CaO aq. Liquid N is obtained by cooling the gas to -186° under a pressure of some hundred atmos., and then reducing pressure, not too suddenly, to not less than 50 atmos., the N does not remain liquid for more

than a few seconds. Liquid N is colourless, transparent, and shows a very sharp meniscus (Wroblewski & Olaszewski, *A Ch* [6] 1, 112). According to Cailletet (*A Ch* [5] 5, 132), N is liquefied, for a second or two, by subjecting the gas to 200 atmos pressure at 13° , and then suddenly reducing the pressure. It has not been solidified. N is slightly lighter than air, 1 litre at the sea level, lat 45° , weighs 1.2574614 grams (von Jolly, *W* 6, 536). N is very slightly sol water (for S v beginning of this art). Small quantities of N are absorbed by molten pig iron, cast iron, and steel (Parry, *Am Ch* 6, 107, Troost & Hautefeuille, *C R* 76, 482, 562, 80, 909, Ledebur, *C C* 1873 810). Wood charcoal also absorbs N , according to R A Smith, charcoal which has absorbed N and O , when exposed to the air for a time gives off O only (*Pr* 12, 424, cf Montmagon & de Laire, *Bl* [2] 11, 261). N is chemically inert, it combines slowly with O when electric sparks are sent through a mixture of the gases for some time, HNO_3 is formed when electrolytic gas ($\text{H}_2 + \text{O}$) is strongly compressed and then exploded in compressed air in presence of a little KOH aq, and also when C is burnt in a mixture of strongly compressed air and O (in presence of a little KOH aq) (Hempel, *B* 23, 1455). N and H combine under the influence of the electric discharge. At or towards white heat N combines with B , Cr , Mg , Si , and V , probably also with Al , Fe , and Zn . The compounds of N are extremely numerous and exhibit great differences of properties. N is related chemically to P , V , As , Nb , Sb , Bi , Er , Ta , and Br , these elements form Group VI. Most of the oxides of N are acidic, none is basic, NH_3 , however, is markedly basic and alkaline (*v* Nitrogen group of elements, p 571).

The influence exerted on the molecular volumes of N compounds by the N atoms has not yet been measured satisfactorily (for a synopsis of data *v* Kopp, *A* 250, 1). The molecular rotatory power of N compounds varies according as the N atom is in direct union with 3 or 5 other atoms, but the exact numerical value to be assigned to N^{III} and N^V has not yet been finally determined (*v* Perkin, *C J* 55, 680). Neither have final values been yet determined for the atomic refractions of N^{III} and N^V .

The atomic wt of N has been determined (1) by finding the ratio of Ag to AgNO_3 , the at wts of Ag and O being known (Marignac, *A* 59, 289, Stas, *Rech* 50, *Nouv R* 281), (2) by finding the ratio of NH_4Cl to Ag needed to ppt the Cl (*M, l.c.*, Stas, *Rech* 87, *Nouv R* 57).

Supposed allotropic form of nitrogen. — By passing a succession of powerful electric sparks through N at not more than 20 mm pressure, Thomson & Threlfall (*Pr* 40, 329) observed a diminution in the volume of the N , at 8 mm the diminution amounted to 8 to 10 p c of the original volume, after long warming to 100° the gas attained its original volume. T \& T supposed that an allotropic form of N is produced under these conditions. According to Johnson (*C J* 39, 130), when N , obtained from KNO_3 aq and NH_4Cl aq, is mixed with H , and the gases are passed over spongy Pt , NH_3 is formed, but NH_3 is not produced if the gases are passed through a hot tube before coming in contact with the spongy Pt . Johnson concluded that N can exist

in two forms one active and the other inactive, the latter being formed by the action of heat on the former (*v* also Johnson's pamphlet, *Elementary Nitrogen, and on the Synthesis of Ammonia* (Churchill, 1885))

Fixation of nitrogen by growing plants—Experiments conducted in recent years tend to show that certain plants, notably *Leguminosae*, are able to absorb N from the air and build up nitrogenous material therewith. The absorption of N seems to occur in nodules which grow on the roots of the plants. For an account of the more important experiments up to the early part of 1890 *v* Lawes & Gilbert, *Pr* 46, 85, abstract in *N* 42, 41 *v* also Atwater & Woods, *Am* 12, 526, also Schloesing & Laurent, *C R* 111, 750, abstracts in *C J* 60, 353

Reactions and Combinations—1 Combines with oxygen to form NO_2 when electric sparks are sent through the gases for some time. When air is strongly compressed and mixed with compressed O and some electrolytic gas ($\text{H}_2 + \text{O}$), an open tube containing KOHAq is placed in the vessel, and a spark is passed, a considerable quantity of KNO_3 is found in the KOHAq (Hempel, *B* 23, 1455). HNO_3 is also formed by exploding $\text{H}_2 + \text{O}$ in air at the ordinary pressure standing over Hg (Bunsen, *Gasom Methoden* [2nd ed.], 71). Nitrites, or HNO_2 , are formed in very small quantities in certain cases of combustion in air, *e g* when P, H, or ether is slowly burnt (Schönbein, *J pr* 84, 193, 86, 129, Berthelot, *A Ch* [5] 12, 440, *C R* 108, 543, Kolbe, *A* 119, 176, Zöller & Grete, *B* 10, 2145, Ilosva, *Bl* [3] 2, 734). It is doubtful whether the nitrites are produced by the oxidation of N or of NH_3 in the air, the experiments of L T Wright (*C J* 35, 42) tended to show that nitrites are not formed by burning H in air from which NH_3 has been carefully removed. According to Ilosva (*Bl* [3] 2, 734), nitrites are formed by passing air over Pt black heated to c 250°. Loew (*B* 23, 1443) showed that small quantities of nitrites are produced when pure Pt black is treated with pure NaOHAq in the air. Neither of these sets of experiments proves conclusively that the N, and not the NH_3 , of the air was the source of the N of the nitrites produced. Schönbein's statement that N combines with ozone has been disproved by Carius (*A* 174, 31).—2 N combines with hydrogen to form NH_3 , under the influence of the electric discharge (Chabrier, *C R* 75, 484, Donkin, *Pr* 21, 281, Morren, *C R* 48, 482, Perrot, *C R* 49, 204, *c* Johnson, *C J* 89, 130, and Wright, *C J* 39, 359). Ramsay & Young assert that a trace of NH_3 is formed when a mixture of moist N and H is passed through a red hot tube containing iron filings (*C J* 45, 93).—3 At a very high temperature N combines with boron, chromium, magnesium, and silicon, and probably also with aluminium, iron, and zinc, to form nitrides (*v* these elements).—4 N combines with carbon to form C_2N_2 , when induction-sparks are passed between C poles in an atmosphere of N (Morren, *C R* 48, 842). Cyanides are formed when a mixture of C with oxide of an alkali, or alkaline earth, metal is heated in N, Hempel (*B* 23, 8890) has shown that considerable quantities of cyanides are thus formed if the reaction occurs at pressures from 10 to 60 atmospheres.

Detection of nitre nitrogen, i.e. N in combination as nitrite or nitrate One part of N existing as a nitrite or nitrate in 20,000,000 parts of water suffices to give a violet-blue colour with a drop of diphenylamine sulphate in H_2SO_4 , followed by 2 cc conc H_2SO_4 , and stirring (*v* Warrington, *C J* 45, 644).

Nitrogen, acids of The compound N_2H_4 , known as *hydrazoic acid*, is described under *Nitrogen, hydrides of*, p 559, for the *Oxyacids of nitrogen v* p 587

Nitrogen, boride of, v BORON NITRIDE, vol 1, p 527

Nitrogen, bromide of ? NBr_2 . According to Millon (*A Ch* [2] 69, 75) the red, very explosive, oily, liquid formed by adding KBrAq to N chloride covered with a little water is a bromide of N

Nitrogen, chloride of NCl_3 . This compound is frightfully explosive. Experiments must be conducted with small quantities and with the greatest care. V Meyer (*B* 21, 26) describes a glass case in which experiments with NCl_3 may be conducted

Preparation—1 A stick of NH_4Cl is suspended in as conc HClO_4aq as can be obtained, an oily liquid slowly collects in a small leaden basin placed at the bottom of the vessel in which the reaction proceeds (*cf* Troost & Hautefeuille, *C R* 69, 152).—2 NH_4ClAq saturated at 35° is poured into a glass basin, and a glass cylinder, closed at one end by parchment, and partly filled with the same NH_4ClAq , is placed upright in the liquid in the basin, a Pt plate, forming the positive pole of a battery of at least 6 to 8 Grove or Bunsen cells, is immersed in the NH_4ClAq in the cylinder, and the negative pole—also a Pt plate—is placed obliquely under the parchment which closes the lower end of the cylinder. A very thin layer of turpentine is spread on the surface of the NH_4ClAq in the cylinder. When the current is sent through the liquid very small oily drops form at the positive pole and float to the surface, where they explode on coming into contact with the turpentine (Böttger & Kolbe, *A* 64, 236, Böttger, *J pr* 68, 374).—3 About 30 grams pure NH_4Cl are dissolved in hot water, the solution is filtered if necessary, diluted to 1½ litres, and placed in a perfectly clean leaden basin, a small leaden basin with a handle is placed in the centre of the larger basin, a fair-sized, very clean flask is filled with Cl, and this flask is immersed in the NH_4ClAq , so that the mouth of the flask covers the small leaden dish. The apparatus is placed out of direct sunlight, in a glass case with double walls, having an open door at one end (*v* V Meyer, *B* 21, 26). The Cl is slowly absorbed by the NH_4ClAq which rises in the jar, when about ½ of the Cl has disappeared, oily drops begin to be formed in the liquid, these drops increase in quantity and size, and at last sink into the small leaden dish. The leaden dish is very carefully removed, and its contents are poured into a small separating funnel made of very thin glass, the NH_4ClAq is removed by a pipette, the greatest care being taken that the liquid does not come into contact with any kind of organic matter, the oil in the funnel is repeatedly washed in the cold water till the washings are free from Cl, and a gentle stream of air is sent through the oil to remove

the last traces of Cl. The oil is now allowed to drop from the funnel into a very small glass vessel, where it is dried by contact with a little bit of dry CaCl_2 , the oil is then poured into a little weighed tube holding about 1 c.c., and closed by a loosely fitting stopper. The very greatest care is required in conducting these operations, especially the removal of the oil from the separating funnel, as the rubbing of the glass tap against the funnel is very apt to cause explosion, when a little of the oil has been dropped into the vessel in which it is to be dried, another clean glass dish must at once be placed beneath the funnel, as explosion would occur if a trace of the oil should drop on to the table (For more details *v* Gattermann, *B* 21, 751.) The oil thus obtained is a mixture of chlorinated ammonias $\text{NH}_{3-2}\text{Cl}_x$ (*G*, *l* c), the composition of portions of the oil varies. To prepare pure NCl_3 , after washing the oil in a separating funnel till free from Cl, and separating the water as completely as possible, Gattermann passes a fairly rapid stream of pure Cl over the oil, which is in the narrow part of the funnel, for about $\frac{1}{2}$ an hour, he then washes and dries the oil in the way described. The analysis was made by decomposing the oil by NH_4Aq , when N and HCl are formed (the HCl combining with excess of NH_3 to form NH_4Cl), and estimating Cl. The process is carried out by Gattermann (*l* c) by dropping the little weighing tube and the stopper (which is removed from the tube) into water in a flask, closed by a cork carrying a small dropping funnel and a tube passing downwards into a beaker of water, allowing about 20 c.c. conc. NH_4Aq to flow very slowly into the flask, when the decomposition is complete (about 4 hours are required) adding the water in the beaker to the contents of the flask, and boiling for a short time, adding HNO_3Aq and AgNO_3Aq , and weighing the AgCl formed.

Properties and Reactions.—A dark yellow oil, *S* *G* c 1 b (determined by finding that the oil very slowly sank in $\text{Fe}_2(\text{SO}_4)_3\text{Aq}$, *S* *G* 1 578, Porret, Wilson, a Kirk, *G* 47, 56). Explodes when exposed to direct sunlight or the light of burning Mg. Explodes at c. $90^\circ\text{--}95^\circ$ when heated in a perfectly clean tube (*v* *G*, *l* c), explodes on contact with wood, grease, oil, or almost any kind of organic matter. The explosion of NCl_3 is frightfully violent. The older observers said that explosions occurred under most curious and apparently contradictory conditions, *eg* contact with P, As, or Se caused explosion, but no explosion occurred by contact with C, S, gum, starch, or wax. Gattermann (*l* c) thinks that light was the cause of many of these explosions. The vapour of NCl_3 acts on the eyes and mucous membrane of the nostrils. NCl_3 is decomposed by conc. HClAq , giving NH_4Cl and Cl, NH_4Aq produces NH_4Cl and N, Hg forms HgCl_2 , and N, SO_3Aq produces NH_3 , H_2SO_4 , and HCl, $\text{As}_2\text{O}_3\text{Aq}$ and Se_2Aq also set free N.

The formation of NCl_3 from N and Cl would be accompanied by the disappearance of much heat. Deville a Hautefeuille give $[\text{N}, \text{Cl}] = -38,000$ (*C* *R* 69, 152, *cf* Oger, *A* *Ch* [5] 20, 5).

References.—Dulong, *G* 47, 48; Porret, Wilson, a Kirk, *G* 47, 56, H. Davy, *T* 1813 1, 242, Serullas, *P* 17, 304, Milon, *A* *Ch* [2] 69, 76, Bineau, *A* *Ch* [3] 15, 52; Gladstone,

C *J* 7, 51, Deville a Hautefeuille, *C* *R* 69, 152, Böttger a Kolbe, *A* 64, 236, Böttger, *J* *pr* 68, 374, Gattermann, *B* 21, 751.

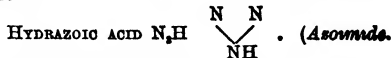
Nitrogen, chlorophosphide, *v* Nitrogen, phosphochloride of, p 570

Nitrogen, chlorosulphide of, *v* Nitrogen, sulphochloride of, p 571

Nitrogen, fluoride of Warren (*C* *N* 55, 289) says that a yellow oil, probably a fluoride of N, is obtained by electrolysis NH_4FAq , the oil explodes by contact with a gold wire.

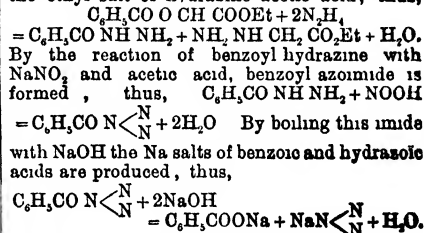
Nitrogen, hydrides of

Three compounds of N and H have been isolated, *viz* ammonia, NH_3 , hydrazine, N_2H_4 , and hydrazoic acid N_3H . Ammonia is described in vol 1 p 196, and hydrazine in vol 1 p 706. As hydrazoic acid has been isolated since the publication of vol 11 this compound is described here—



Hydrogen nitride (*B* 23, 3023) This acid was discovered by Carius in 1890 (*B* 23, 3023).

Formation.—Ethyl benzoylglycolate reacts with N_2H_4 to form benzoyl-hydrazine and the ethyl salt of hydrazine acetic acid, thus,



By adding dilute $\text{H}_2\text{SO}_4\text{Aq}$ and warming, hydrazoic acid gas is evolved.

Preparation.—Ethyl hippurate is dissolved in as small a quantity as possible of boiling alcohol, $\text{N}_2\text{H}_4\text{H}_2\text{O}$ is added in the ratio $\text{C}_6\text{H}_5\text{CO} \text{NH} \text{CH}_2 \text{COOEt} : \text{N}_2\text{H}_4\text{H}_2\text{O}$, hippuryl hydrazine, $\text{C}_6\text{H}_5\text{CO} \text{NH} \text{CH}_2 \text{CO} \text{NH} \text{NH}_2$, separates on cooling. The crystals are recrystallised from alcohol, and dissolved in much warm water, with addition of rather more than a molecular proportion of NaNO_2 , the solution is cooled to 0° , and mixed with excess of acetic acid, when lustrous tablets of a nitroso compound (probably $\text{C}_6\text{H}_5\text{CO} \text{NH} \text{CH}_2 \text{CO} \text{N} \text{N} \text{O}$) separate, the

crystals are collected by help of a filter pump, washed with cold water, and dissolved in very dilute NaOHAq . This solution is gently warmed for a short time on the water bath, and is then placed in a flask connected with a condenser and furnished with a dropping funnel. A flask containing AgNO_3Aq is used as a receiver, dilute $\text{H}_2\text{SO}_4\text{Aq}$ is allowed to drop very slowly into the boiling liquid in the flask, N_2H_4 distils over with steam, and, reacting with the AgNO_3 in the receiver, produces AgN_3 , the operation is continued so long as a pp is produced in the receiver. The AgN_3 is collected by the help of the pump, and well washed with cold water, it must not be heated above 60° , else there is danger of a severe explosion. The AgN_3 is decomposed by boiling with dilute HClAq , the dis-

tillate is fractionated, and the portion which distils over in the early stages is collected separately from the rest. In this way a solution of N_2H containing 27 p.c. N_2H is obtained. By repeatedly fractionating with great care, N_2HAq containing over 90 p.c. N_2H is obtained, and all water can be removed from this solution by fused $CaCl_2$ (Curtius a. Badenhause, *J pr* [2] 43, 207). The process of fractionation is often attended with explosions.

Properties and Reactions.—A 27 p.c. solution of N_2H is a thickish liquid, which sinks in water, it possesses an extremely offensive odour, with NH_3 it gives white clouds. Pure N_2H is a clear, colourless, very foully smelling, liquid. It boils at 87° . It is very explosive. When touched with a hot substance it explodes with violence, it also explodes when placed in a barometric vacuum. The solution of N_2H corrodes the skin and causes headache and giddiness. N_2HAq is a strong monobasic acid, it dissolves Fe, Zn, Cu, Al, and Mg with rapid evolution of H , it appears to dissolve slightly Au and Ag. With $AgNO_3$ and $HgNO_3$ white pps of AgN and HgN are obtained. The affinity of N_2HAq is a little greater than that of acetic acid. The salts of N_2H are also very explosive, with the exception of those of the alkali and alkaline earth metals.

The following salts are described by Curtius (*B* 23, 3032). $N(NH_2)_2$, $(N_2)_2Ba$, N_2Hg , N_2Ag , salts of Cu, Fe, and Na were also prepared.

In connexion with N_2H v. Mendelejeff, *B* 23, 3464.

Nitrogen, iodides of (Iodamines). Very explosive compounds containing N and I are obtained by rubbing I with conc. NH_3 or by pouring an alcoholic solution of I into NH_3 or alcoholic NH_3 , by pouring NH_3 into alcoholic I, by pouring a solution of I in *aqua regia* into NH_3 or NH_4Cl , by adding bleaching powder solution (neutralised by acetic acid) to NH_4I , by adding NH_3 to a mixed solution of HCl and HIO_3 , by the action of N chloride on $KIAq$, and by adding I and alcohol to 'white precipitate'. The products of these reactions are very dark-coloured powders, which explode, more or less readily and violently, by rubbing or striking. Analyses of these substances seemed to show that at least three different explosive compounds existed, viz. NH_2I , NH_2I , and $N_2H_2I_2$ ($=NH_2NI$). For details and analyses v. Serullas, *P* 17, 304, Millon, *A Ch* [2] 69, 78, Marchand, *J pr* 19, 1, Bineau, *A Ch* [8] 16, 71, Gladstone, *C J* 4, 84, 7, 61, Bunsen, *A* 84, 1, Stahlschmidt, *P* 119, 421, Champion a. Pellet, *Bl* [3] 24, 447, Mallet, *Am* 1, 4, Guyard, *A Ch* [6] 1, 358, Guthrie (*C J* [2] 1, 239), by adding I to conc. NH_4NO_3 or $(NH_4)_2CO_3$ containing KOH , obtained a brown black liquid, which, he said, had the composition NH_2I (*v post*).

Raschig (*A* 230, 212) has re-examined the various methods of preparing and analysing N iodides. According to B, three compounds exist, NI , NH_2I , and NH_2I , but only NI and NH_2I have been isolated by B.

Tri-iodamine NI_3 . NH_4Cl and I in the ratio NH_4Cl 61 (1.14 24) were dissolved in $KIAq$, and $NaOH$ was added in the ratio NH_4Cl 4NaOH, v.e. 2.99 parts NaOH for each part NH_4Cl used, the black pp which forms was collected immediately (with the help of a filter-pump) and

washed 6-8 times with cold water, as rapidly as possible, then dissolved in HCl and analysed. Raschig (*lc*) expresses the reaction thus, $NH_4Cl + 6IAq + 4NaOH$

$= NI_3 + NaCl + 3NaI + 4H_2O$. NI_3 is rapidly decomposed by water to NH_2I . Mallet (*Am* 1, 4) obtained NI_3 by triturating I with a large excess of the most conc. NH_3 , keeping temperature at or below 0° , pouring off the liquid and triturating with more NH_3 , repeating this several times, then agitating 2 or 3 times in a cooled flask with absolute alcohol, then with dry ether, and allowing the ether to evaporate. NI_3 is a heavy, nearly black, powder. According to Raschig (*lc*), the NI_3 prepared by the action of NH_3 on I is much more explosive than the iodide formed by the action of NH_4Cl and $NaOH$ on I. NI_3 dissolves in $KOCy$, forming IOy , KOH , and NH_3 (Millon, *A Ch* [2] 69, 78, Raschig, *A* 230, 212). $KSCy$ reacts similarly, producing IOy , HI , KOH , NH_3 , and H_2SO_4 (B, *lc*).

Di-iodamine NHI . Prepared similarly to NI_3 , using the materials in the ratio NH_4Cl 4I 8NaOH (Raschig, *lc*). Mallet (*Am* 1, 4) obtained this compound by triturating I with not very conc. NH_3 at the ordinary temperature, washing with water till NH_3 could not be detected in the washings, keeping under water for three days, washing with alcohol and ether, and allowing to dry. Gladstone obtained NHI by acting on an alcoholic solution of I with NH_3 (*C J* 4, 34, 7, 51), with H_2S it gave HI and NH_3 , with SO_4 the products were NH_3 , HI , and H_2SO_4 . Stahlschmidt obtained NHI by adding alcoholic NH_3 to an alcoholic solution of I (*P* 119, 421).

Mono-iodamine NH_2I . Raschig (*A* 230, 212) expected to obtain this compound by the reaction of NH_4Cl and $NaOH$ with I, using these materials in the ratio NH_4Cl 2I 2NaOH, but the pp was very quickly decomposed by water to NH_2NI . Millon (*A Ch* [2] 69, 78) gave the formula NH_2I to the iodide prepared by him, and Marchand (*J pr* 19, 1) confirmed this composition. Guyard (*A Ch* [6] 1, 358) describes a light-brown explosive compound, decomposed by light, exploding in contact with water, prepared by NH_3 reacting with I in an iodide solution, to this compound he gives the formula NH_2I , with an ammoniacal solution of a Cu salt it forms $Cu_2I_2 \cdot 2NH_3$.

Tri-iod diamine $N_2H_2I_2$ ($=NH_2NI_2$). Obtained by mixing cold nearly saturated alcoholic solution of I and NH_3 , decomposed by HCl giving NH_3 and IOI in the ratio $2NH_3$, 3IOI. This compound seems to have been obtained by Raschig (*A* 230, 212) by mixing NH_4Cl , I, and $NaOH$ in the ratio NH_4Cl 2I 2NaOH, and washing the pp with water.

OTHER COMPOUNDS OF NITROGEN, IODINE, AND HYDROGEN. (1) *Iodammonium iodide*, NH_4II , a mobile brownish red liquid, formed by adding finely powdered I to saturated NH_4NO_3 or $(NH_4)_2CO_3$ mixed with about $\frac{1}{2}$ of an equivalent of KOH (Guthrie, *C J* [2] 1, 239). Soluble alcohol, ether, $CHCl_3$, CS_2 , and $KIAq$, decomposes by heat, giving I and probably NH_2I , decomposes in the air to NH_3 and I, water forms NH_2I , HI , and NH_4I , which explodes producing N , I, and H_2O . According to Seamon (*C N* 44, 189), this compound is formed by act-

ing on dry I with dry NH_3 , and absorbing the excess of NH_3 by standing near H_2SO_4 . S describes the compound as a nearly black liquid, S G 2 46 at 15° , solidifying at -2° , decomposing slowly at 15° , quickly at 70°

(2) *Compounds of ammonia with iodine*— NH_3I , obtained by the action of NH_3 on I at 10° (Millon, *A Ch* [2] 69, 78), formed at 80° according to Raschig (*A* 241, 253) β $(\text{NH}_3)_2\text{I}$ formed at 20° (Bineau, *A Ch* [3] 15, 71, Raschig, *lc*) γ $(\text{NH}_3)_3\text{I}$, formed at 0° δ $(\text{NH}_3)_4\text{I}_2$ formed at -10° (R, *lc*) It is doubtful whether these bodies are true compounds or not

Nitrogen, oxides of N forms five oxides N_2O , NO , N_2O_3 , NO_2 , N_2O_5 N_2O_3 and N_2O_5 are the anhydrides of HNO_2 and HNO_3 , respectively, NO , reacts with water to produce both HNO and HNO_2 , NO is obtained by the decomposition of HNO_3Aq , but the acid has not been obtained from the oxide, NO is a neutral oxide Whether N_2O_3 exists in the gaseous state is not yet finally settled, the other oxides, with the exception of N_2O_5 , are gases under ordinary conditions NO_2 exhibits polymerism, at low temperature the molecular weight corresponds with the formula N_2O_4 , and at higher temperatures with the formula NO_2 Besides these five oxides, there is said to exist a pernitric oxide NO_3 , or N_3O_4

NITROUS OXIDE N_2O (Nitrogen monoxide Laughing gas) Mol w 43.98 Melts at -99° , and boils at -92° (Wills, *C J* [2] 12, 21) S G 1 527 S G liquid N_2O 9756 at -5° , 937 at 0° , 8964 at 10° , 8365 at 20° (Andr  eff, *A Ch* [3] 56, 317, cf Wills, *C N* 28, 170, Wroblewski, *C R* 97, 166, Caill  tet a Mathias, *C R* 102, 1202) V D 22.1 S H p (equal wt of $\text{H}_2\text{O}=1$) 16° to $207^\circ=22616$ (Regnault, *Acad* 26, 1), 26° to $103^\circ=2126$, 27° to $206^\circ=2241$ (Wiedemann,

P M [5] 2, 81) S H p 1 3106 at 0° , 1 27238 at S H v

100° (Clausius, *Mechan W  rmetheorie*, 1 62) CE (22° to 98°) 0037067 (von Jolly, *P Jubelbd* 82) S 1305 at 0° , 1095 at 5° , 92 at 10° , 778 at 15° , 67 at 20° , absorption coefficient = 1 30521—0 45362t + 0 006843t (Carius, *A* 94, 139) Absorption coefficient in alcohol = 4 17805—0 69816t + 0 00609t (Carius, *lc*) H F [N , O] = -17,740, [NO , N] = 3835 (*Th* 2, 198) For vapour pressures from -25° to 40° v Regnault, *J* 1863 66

Nitrous oxide was discovered by Priestley in 1776, and carefully studied by Davy

Formation—1 By dissolving Zn in HNO_3Aq (S G 1 2 diluted with an equal vol of water)—2 By decomposition of NH_4OHAq by AgNO_3 , K_2CO_3 , &c. v HYDROXYLAMINE, *Reactions*, No 1 (vol u p 735)—3 By the gradual decomposition of $\text{H}_2\text{N}_2\text{O}_3\text{Aq}$ (v *Hyponitrous acid*, p 564)—4 By the action of SnCl_2 in HCl on HNO_3Aq or a nitrate—5 By passing NO through SO_2Aq or an acid sulphate—6 By the action of HNO_3Aq on Cu in presence of much Cu_2NO_3 , if NH_4NO_3 is present, much N_2O and N, with little NO , are produced (Acworth, *C J* 28, 828)

Preparation—1 Pure NH_4NO_3 is slowly heated in a retort to a temperature at which gas begins to be given off The gas flame is then lowered, and the decomposition allowed to proceed $\text{NH}_4\text{NO}_3=\text{N}_2\text{O}+2\text{H}_2\text{O}$ It is best to make the

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NH_4NO_3 by neutralising pure dilute HNO_3Aq with pure NH_3Aq or $(\text{NH}_4)_2\text{CO}_3\text{Aq}$, evaporating till the B P gets to $c\ 120^\circ$ and a drop solidifies on a cold plate, allowing to cool, and breaking up into small pieces If the temperature rises above 250° , decomposition to N, O, and H_2O may occur with explosive violence, in the change $\text{NH}_4\text{NO}_3=\text{N}_2\text{O}+2\text{H}_2\text{O}$ c 31,100 cal are reproduced, whereas the change $\text{NH}_4\text{NO}_3=\text{N}_2+\text{O}+2\text{H}_2\text{O}$ is accompanied by the production of c 48,700 cal (*Th* 2, 205) To obviate explosions, Case neuve (*D P J* 257, 435) recommends to dry NH_4NO_3 carefully over a low flame, to bring the salt while warm into a retort or flask, to heat with a very small flame which is gradually increased till decomposition begins, and then to withdraw the flame The gas is collected over hot water, brine, or Hg If the gas is to be used as an anæsthetic, the greatest care should be taken that the NH_4NO_3 used is pure, and especially that it is quite free from NH_4Cl , else the N_2O may contain Cl, the gas should be passed through KOH Aq and FeSO_4Aq to absorb traces of Cl, NO , and $\text{NO}-2$ A mixture of 5 parts SnCl_2 , 10 parts HCl Aq S G 1 21, and 9 parts HNO_3Aq S G 1 38, is heated to boiling, when pure N_2O is evolved in a regular stream, any alteration in the proportions may cause explosions (Campari, *C C* 1888 1569)

Properties—A colourless gas, with a slightly sweetish smell and taste Supports combustion almost as well as O When breathed, N_2O produces insensibility, which lasts for a short time only It is often used as an anæsthetic in dental operations, for this purpose it is usually much compressed in iron bottles N_2O is decomposed by heat, the decomposition being complete at $c\ 900^\circ$ H and N_2O forms an explosive mixture, most inflammable gases burn in N_2O N_2O is liquefied at 0° by a pressure of c 30° atmos, the liquid occupies c $\frac{1}{30}$ of the volume of the gas (Faraday, *A* 56, 157) Liquid N_2O is colourless and very mobile, a drop burns when let fall on the skin, on evaporation much cold is produced, liquid N_2O boils in liquid CO_2 Metals dropped into liquid N_2O generally hiss as hot iron does in water Glowing C swims on the liquid and burns briskly Hg sinks and freezes, S, I, P do not react Water freezes when added to liquid N_2O , but so sudden an evolution of vapour occurs that an explosion generally takes place (Faraday, *A* 56, 157), nitric and sulphuric acids are frozen by liquid N_2O , alcohol, ether, and CS, mix, without freezing By evaporating liquid N_2O in an air stream, Wills (*C J* [2] 12, 21) obtained solid N_2O in some quantity, the solid is a snow like substance, more compact than solid CO, when slightly warmed it melts, at $c\ -99^\circ$, and then boils, at $c\ -92^\circ$

Reactions—1 Decomposed by heat to N and O, change is complete at $c\ 900^\circ$ (Langer a V Meyer, *Pyrochemische Untersuchungen* [Brunswick, 1885], 65, cf Berthelot, *C R* 77, 1448)—2 Decomposed to N and O, by electric sparks 3 Passed over red-hot iron, Fe_2O_3 , and N are formed—4 Potassium or sodium burns in N_2O setting free N—5 Mixed with hydrogen, or other combustible gas such as CO, PH_3 , H_2S , or a hydrocarbon, and brought to a flame, combustion ensues at the expense of the O of the N_2O (explosions occur with H)—6. *Easily burnt sub*

O O

stances, when inflamed and plunged into N_2O , burn almost as rapidly as in O, e.g. C, P, S, Na, &c.—7 Mixed with oxygen, and subjected to the electric discharge, NO_2 is formed.—8 A solution of N_2O in water is neutral to litmus, N_2O is, however, related to the acid $H_2N_2O_3$, as it is obtained by the decomposition of this acid in aqueous solution (*v. Hyponitrous acid*, p. 569). The hyponitrites may be regarded as compounds of the negative radicle N_2O with more positive oxides, e.g. Ag_2O , N_2O .

References to older memoirs—Gay Lussac, *G A* 58, 29, W. Henry, *Annals of Phil* 24, 299, 344, Pléischl, *S* 38, 461, Davy, *G A* 6, 105, Natterer, *P* 62, 133, Dumas, *C R* 27, 463.

NITRIC OXIDE NO (*Nitrogen dioxide Deutoxide of nitrogen Nitrous gas or air*) Mol w 29.97 VD 15, not changed from -100° to 1200° (Dacomo a V Meyer, *B* 20, 1832, Langer a Meyer, *Pyrometrische Untersuchungen* (1885) 66). SH p (equal wt of $H_2O = 1$) 18° to $172^\circ = 23173$ (Regnault, *Acad* 26, 1), SH v (equal wt of $H_2O = 1$) 1652 (Clausius, *Mechan Wärmetheorie*, 1, 62). S c 05 at ordinary temp S (in alcohol) 309 at 2° , 282 at 11.8° , 266 at 20° (Bunsen, *Gasom Methoden*, [2nd ed.] 227). S (in H_2SO_4 , SG 1.84) at $18^\circ = 3.5$, S (in H_2SO_4 , SG 1.5) at $18^\circ = 1.7$ (Lunge, *B* 18, 1391). HF [N, O] = $-21,575$, [N $_2$ O, O] = $-25,410$.

NO is liquefied at the following temperatures and pressures (Olszewski, *C R* 100, 940) —

Pressure in atmos	71.2	57.8	49.9	41
Temperature	-93.5°	-97.5°	-100.9°	-105°
Pressure in atmos	31.6	20	10.6	5.4
Temperature	-110.9°	-119°	-129°	-138°
Pressure	1 atmo	138 mm	18 mm	
Temperature	-153.6°	-167°	-176.5°	

The critical temp is c -93.5° according to Olszewski, but according to Cailliet (*C R* 85, 1016) NO is liquefied at -11° under a pressure of 104 atmos, but not at 8° even under 270 atmos pressure. NO solidifies at -167° (Olszewski, *l.c.*).

NO was discovered by Priestley and called by him *saltpetre gas*.

Formation—1 NO is probably formed in the first stages of those combustions in air whereby nitrites and nitrates are produced, e.g. in burning air with compressed O, in slowly burning P, ether, &c., in air (*v. Nitrogen, Reactions*, No 1, p. 558). NO is also probably formed when electric sparks are passed through a mixture of N and O.—2 By reducing HNO_3 Aq or HNO_2 Aq, e.g. by reaction with Cu, by passing in SO_2 , by electrolysis, &c. (*v. Nitric acid, Reactions*, Nos 3, 4, 5, p. 520).—3 By reacting on KNO_3 with $FeCl_3$ Aq and HCl Aq, or with $FeSO_4$ Aq and H_2SO_4 Aq.

Preparation—1 A quantity of HCl Aq is divided into 2 equal parts, one portion is converted into $FeCl_3$ Aq by dissolving iron wire in it until it is saturated, the two portions are then mixed and placed in a retort or flask, KNO_3 is added in quantity nearly equal to that of the Fe used, and the whole is warmed ($6FeCl_3$ Aq + $8HCl$ Aq + $2KNO_3$ = $6FeCl_2$ Aq + $2KCl$ Aq + $4H_2O$ + $2NO$). $FeSO_4$ Aq and H_2SO_4 Aq may be substituted for $FeCl_3$ Aq

and HCl Aq.—2 Cu cuttings are added to HNO_3 Aq, SG 1.2, in a flask arranged so that it can be surrounded by cold water when desired, action begins after a little and red fumes of NO_2 are evolved, when the gas in the flask is quite colourless it is collected over cold water, or if required dry it is passed over solid dry KOH and then through conc H_2SO_4 . About 130 c.c. acid are used for 15 grams Cu. The HNO_3 Aq used should not be more conc than SG 1.2, and temperature must be kept as low as possible, else N_2O and N_2O_3 may be formed. If the action proceeds for some time the gas contains considerable quantities of N_2O , as the reaction of Cu with much $Cu(NO_3)_2$ Aq in presence of HNO_3 produces this gas (Acworth, *C J* 28, 828). But with all precautions, NO prepared by this method is always liable to contain N_2O and N (*v. Acworth, l.c.*). Carus (*A* 94, 138) says NO can be obtained pure by passing the products of the reaction of Cu with HNO_3 Aq into $FeSO_4$ Aq, which absorbs NO but not N_2O or N, and then gently warming the solution thus obtained.—3 Kammerer (*B* 18, 3064) recommends to fill a Wolff's bottle with Cu turnings, add enough cold saturated $NaNO_2$ Aq to fill the bottle $\frac{1}{2}$, and then allow conc H_2SO_4 to drop in little by little.—4 Thiele (*A* 253, 246) prepares NO by adding conc $NaNO_2$ Aq (free from carbonate) to $FeCl_2$ or $FeSO_4$ in HCl Aq.—5 SO_2 is prepared by heating Cu with conc H_2SO_4 , and passed into slightly warmed HNO_3 Aq SG 1.15, the issuing gas is passed through wash bottles wherein excess of SO_2 is absorbed ($3SO_2 + 2HNO_3$ Aq + $2H_2O = 3H_2SO_4$ Aq + $2NO$).

Properties—A colourless gas, at great pressure and very low temperature, a colourless liquid, becomes solid at -167° . As NO combines with O immediately on coming into contact with air, it cannot be determined whether pure NO is or is not tasteless and odourless. Sl. sol water, v. sl. sol H_2SO_4 , less sol H_2SO_4 Aq. NO does not change the colour of litmus. Substances which produce much heat when burning, e.g. P and C, continue to burn in NO if plunged into the gas when burning vigorously, burning S or H, or a burning taper, ceases to burn in NO . A mixture of NO and H is not explosive. NO is reduced to N_2O by Zn, Fe, SO_2 , &c. At a very high temperature, c. 1700° , NO is decomposed to N and O.

The SG of NO has been determined from -100° to 1200° (*v. beginning of this article*), for this temperature interval the molecular formula is NO .

NO is a neutral oxide. In the compound $NOCl$, formed by the union of NO and Cl , NO acts as the more positive radicle, it also perhaps takes the part of the positive radicle in nitrosulphuric acid, which may be regarded as $SO_2.OH.ONO$.

Reactions—1 Decomposed into N and O by heat, but only at a very high temperature, c. 1700° , according to Langer and V Meyer (*Pyrochemische Untersuchungen*, 66).—2 *Electric sparks* produce N_2O (which then goes to N and O) and O (Berthelot, *Bl* [2] 26, 101).—3 Reduced to N_2O by many easily oxidised substances, e.g. moist zinc or iron filings (NH_3 also is produced), sulphur dioxide and water (reductions occurs even in presence of O if H_2SO_4 is

also present, *v* Lunge, *C J* 47, 465), hot copper, &c —4 Heated with potassium or sodium, K_2O or Na_2O , and N are formed —5 Passed over red hot carbon N and CO , are obtained —6 Mixed with hydrogen and inflamed, H_2O and N are formed, NH_3 is formed by passing a mixture of NO and H over spongy Pt (L Wright, *C J* 89, 357) —7 The copper-*unc* couple acting on NO in presence of water produces NH_3 (Gladstone a Tribe, *C J* 43, 341) —8 Stannous chloride, in presence of $HClAq$, produces NH_4OH , HCl , N, and $SnCl_4$, the action ceases at 100° (Divers a Haga, *C J* 47, 623) —9 When a flame is brought to a mixture of NO and carbon disulphide, CO_2 , SO_2 , and N are produced very rapidly, and with a brilliant white flash —10 In contact with water in the dark slowly produces HNO_3Aq , N, and a little N_2O (Cooke, *C N* 58, 115, *cf* Russell a Lapraik, *C J* 32, 37) —11 Contact with conc potash solution causes a slow formation of KNO_3Aq , N_2O , and N (Gay-Lussac, *Gm* 2, 378, Russell a Lapraik, *C J* 32, 35) —12 With alkaline solution of pyrogallol N_2O and N are formed (R a L, *l c*) —13 Reduced to NH_3 , with separation of I, by conc hydroiodic acid solution (Chapman, *C J* [2] 5, 166) —14 An alkaline solution of stannous hydroxide ($=K_2SnO_2$) produces $K_2N_2O_2$ and K_2SnO_3 (Divers a Haga, *C J* 47, 361) —15 Ferrous hydroxide, in presence of much conc $KOHAq$, reduces N_2O to NH_3 (D a H, *l c*) —16 Oxidised to HNO_3Aq by an alkaline solution of potassium permanganate, with separation of $MnO_2 \cdot xH_2O$ (Wanklyn a Cooper, *P M* [5] 6, 288) —17 Oxidised to NO , and N_2O , by nitric acid of SG >1.15 —18 With nitric acid in sulphuric acid, SO_2 , OH , ONO is produced —19 With oxygen and water NO is wholly oxidised to HNO_3Aq (Lunge, *C J* 47, 465) —20 With oxygen and conc sulphuric acid forms SO_3 , OH , ONO (L, *l c*)

Combinations —1 With oxygen to form NO_2 . According to Lunge (*C J* 47, 465), NO_2 is the chief or only product when excess of O is used, but with excess of NO both NO_2 and N_2O , are formed —2 With ferrous salts in solution, to form deep brown coloured liquids Gay (A Ch [6], 5, 145) finds that the quantity of NO absorbed is independent of the kind of ferrous salt used and of dilution It is proportional to the quantity of Fe in solution, and varies with temperature and pressure The relation between quantity of NO absorbed and pressure is a special one, it resembles that which holds good in the solution of NH_3 by water The solutions lose all NO *in vacuo*, or by passing a stream of H into them The quantity of NO absorbed at 8° and 760 mm nearly agrees with the formula $2NO \cdot 3FeSO_4$, between 8° and nearly 25° the formula $NO \cdot 2FeSO_4$, approximately expresses the quantity of NO absorbed, and at $c 25^\circ$ the NO agrees with that required by $NO \cdot 5FeSO_4$ NO is also absorbed by chromous, stannous, and mercurous salt solutions —3 With antimony chloride, to form $NO \cdot 2SbCl_5$, also with aluminum, bismuth, and ferric chloride (Beeson, *C R* 108, 1012) —4 NO combines with liquid N_2O , to form N_2O , (*q v*)

References to older memoirs —Gay-Lussac, A Ch [8] 23, 229, Millon, *C R* 14, 908, Carus, A 94, 188

NITROGEN TRIOXIDE N_2O_3 . (Nitrous anhydride Nitrogen sesquioxide) Mol w 75.9 (*infra*)

Formation —1 By passing NO into liquid N_2O_4 at $c 20^\circ$, more or less pure liquid N_2O_3 is formed (Dulong, A Ch 2, 317, Péligot, A 39, 327, Ramsay, *C J* 57, 590) According to Hasenbach (*J pr* [2] 4, 1), N_2O_3 is produced by passing a mixture of NO and NO_2 through a hot tube, and then into a vessel surrounded by a freezing mixture Ramsay a Cundall (*C J* 47, 672) showed that no contraction occurs when NO_2 and NO are mixed at the ordinary temperature —2 By reacting on starch or As_2O_3 with HNO_3Aq SG 1.3 to 1.35 (Lunge, B 11, 1641, Stenhouse a Groves, *C J* 31, 545) —3 Fritzsche (*J pr* 22, 14) gently warmed red fuming HNO_3 , condensed the vapour, again very gently warmed this liquid, and passed the vapour into a strongly cooled receiver, to 92 parts of the liquid thus obtained (chiefly N_2O_3), he added 45 parts of cold water very slowly, the liquid being kept very cold Two layers of liquid were thus obtained, the under was very deep blue, and was regarded by Fritzsche as fairly pure N_2O_3 , the upper layer, which was grass green, was a solution of N_2O_3 and NO_2 in HNO_3Aq By placing both liquids in a retort kept at 0° , the lower layer boiled, and a pure indigo blue liquid condensed in the very strongly cooled receiver, this liquid consists chiefly of N_2O_3 , according to Fritzsche F (*l c*) obtained a deep blue liquid, containing, according to him, at least 93.4 p c N_2O_3 , by distilling a large quantity of nitric acid which had been used in a battery, condensing the distillate in a vessel surrounded by snow and $CaCl_2$, and redistilling several times at the lowest possible temperature —4 By the reaction of O on excess of NO, considerable quantities of N_2O_3 , are produced (Lunge, *C J* 47, 466) —5 By dropping water on to 'chamber crystals' (nitrosyl sulphate, $SO_3 \cdot OH \cdot ONO$) (Streiff, B 5, 285)

Preparation —Pure N_2O_3 , has not been obtained 1 The gas obtained by reacting on starch or powdered As_2O_3 with HNO_3Aq SG 1.35 is nearly pure N_2O_3 , according to Lunge (B 11, 1229, 1641), the starch is made into a paste with water, and the acid is added from a dropping funnel, the flask is gently warmed, and then plunged into cold water Stenhouse a Groves (*C J* 31, 545) also recommend HNO_3Aq SG, the reaction proceeds at 70° with production of nearly pure N_2O_3 —2 Pure NO is passed into cold liquid N_2O . The product is not pure N_2O_3 (Ramsay, *C J* 57, 591) But *v* Lunge, *Z anorg, Chem* 7, 209 [1894]

Properties —The liquid obtained by condensing the gaseous product of the reaction between As_2O_3 and HNO_3Aq , SG 1.3, at 70° , passing the vapour of this liquid over P_2O_5 , and condensing again, is deep blue, it does not solidify at -90° , it is miscible with liquid N_2O , O passed into liquid N_2O_3 , mixed with N_2O_3 , very slowly, if at all, combines with the N_2O_3 , liquid N_2O_3 appears to dissociate slightly to NO and N_2O , at -90° (Ramsay, *C J* 57, 597) Gaines (*C N* 48, 97) says that N_2O_3 liquefies at -14.4° According to Geuther, liquid N_2O_3 boils at 3.5° , and has the following SG 1.464 at -8° , 1.4555 at -4° , 1.451 at -1° , 1.449 at 0° , 1.4485 at 1° , 1.447 at 2° Birhausa

(*C R* 109, 68) says that N_2O_3 solidifies at -82° , Ramsay found the blue liquid to remain liquid at -90° (Birhauss's liquid probably contained N_2O_4 , as it was formed by the action of O on excess of NO_2 .)

Molecular weight of nitrogen trioxide—Ramsay determined the lowering of the freezing-point of N_2O_4 , after passing in NO and calculating the quantity of N_2O_3 produced by the increase in weight, his results, on the whole, are in favour of the formula N_2O_3 for the liquid compound. There has been much discussion as to the existence of this oxide in the state of gas. Some chemists deny the existence of a compound nN_2O_3 , and say that the reactions of this supposed compound are those of a mixture of NO and NO_2 (containing variable quantities of N_2O_4 according to the temperature). As NO rapidly combines with O, a mixture of NO and NO_2 would also surely combine with O, but Ramsay & Cundall have shown (*C J* 47, 187) that O acts very slowly on the blue liquid which, on the hypothesis of the non existence of N_2O_3 , is a mixture of NO and N_2O_4 , this result is confirmed by Ramsay's later observations (*C J* 57, 597), and also by observations made by Richardson (*C J* 51, 397). Lunge has shown (*D P J* 233, 63) that the gas from the blue liquid which has the empirical composition N_2O_3 is not wholly transformed into NO_2 even in presence of 10 times as much O as is required on the assumption that the gas in question is a mixture of NO and N_2O_4 . Moreover, Ramsay & Cundall (*C J* 47, 672) showed that although no change in appearance or volume occurs when NO_2 gas is mixed with NO, yet on lowering the temperature of the mixed gases, by $HClAq$ and ice, a dark blue liquid was formed (liquid NO_2 is colourless, and liquid N_2O_4 is yellow red). Again, Ramsay (*C J* 57, 597) found that the blue liquid does not freeze at -90° , but part of a mixture of NO and N_2O_4 would freeze at -10° , as this is about the freezing point of N_2O_4 . Some of the reactions of N_2O_3 with alkalis, H_2SO_4 , &c., are scarcely those of a mixture of two substances (*v. Reactions*, 1, 2, and 5). According to Gay Lussac (*G A* 58, 29) a mixture of excess of NO with O allowed to stand for a long time over $KOHAq$ forms KNO_2Aq , and the gases disappear in the ratio required to form N_2O_3 .

As regards the existence of N_2O_3 as a gas, Lunge's experiments show that the oxidation of the gas obtained by heating starch with HNO_3Aq S G 1 55 to NO_2 is not completed by a very large excess of O (*D P J* 233, 63, *B* 12, 357). Now, as Lunge has also shown (*C J* 47, 465), that NO_2 is the sole product, or almost the sole product, of the action of excess of O on NO, and therefore as soon as O is added to NO there must be present a mixture of NO, NO_2 , and O, it is difficult to escape the conclusion that the action of a large excess of O on a mixture of NO and NO_2 must result in the formation of much NO_2 , and that, therefore, the gas obtained by HNO_3Aq acting on starch, contains N_2O_3 , which is not oxidised by O to N_2O_4 . Ramsay & Cundall (*C J* 47, 672) found that no contraction occurred on mixing NO_2 with NO, if any N_2O_3 were formed contraction must have occurred. B & C prepared a blue liquid having the composition N_2O_3 (by action of HNO_3Aq on As_2O_3), and determined the

V D of the gas obtained by gently warming the liquid. The V D was 22 35 at 18.2° , V D corresponding to $N_2O_3 = 88$. If the gas was supposed to be a mixture of NO, NO_2 , and N_2O_3 (without any N_2O_4), there must have been present 17 63 p.c. N_2O_3 to raise the S G of $NO + NO_2$ to 22 35, hence the maximum percentage of N_2O_3 gas which could be present was 17 63, but as the gas certainly contained some N_2O_4 , the percentage of N_2O_3 gas must have been less than 17 63. R & C show that a mixture of NO, NO_2 , and N_2O_3 , having the same percentage of N and O as N_2O_3 , would have the V D 23 42 under the conditions of their experiment, assuming a formula given by Willard Gibbs to be correct, hence they conclude that the gas examined by them was more probably a mixture of NO, NO_2 , and N_2O_3 , than a mixture of these gases with N_2O_4 . R & C also draw attention to the remarkable result of Lunge's experiments, which he has himself stated, that the dissociation of N_2O_3 (supposing it to exist as a gas) is almost independent of temperature. According to Luck (*Fr* 8, 402) and Moser (*W* 2, 139), the absorption-spectrum of the gas supposed to be N_2O_3 is identical with that of NO_2 (containing N_2O_4). On the question of the existence or non existence of gaseous N_2O_3 , besides the memoirs referred to, *v. Lunge*, *B* 11, 1232, 1641, 12, 357, 15, 495, Witt, *B* 12, 2188, Geuther, *A* 245, 96.

Reactions—1 Dissolves in water at 0° , forming a blue liquid, on warming, NO is evolved and HNO_3Aq remains, if the solution is neutralised by alkali, a mixture of a nitrite (MNO_2) and nitrate (MNO_3) is obtained. According to Fremy (*C R* 79, 61) addition of a little water causes evolution of pure NO, and HNO_3 is formed, but addition of much cold water produces a fairly stable solution, which may be kept unchanged for some days, and which slowly evolves NO and N_2O_4 on warming. Addition of sand, powdered $CaSO_4$, or other indifferent substance, causes rapid decomposition to HNO_3Aq and NO. The solution is a strong reducer, cold SO_2Aq forms various nitro derivatives of S oxy acids, warm SO_2Aq forms NO, N, NH_3 , and H_2SO_4 (*F*, *1 c*, *cf. Reinsch*, *J pr* 28, 399).—2 Passed into *alkali solution*, nitrite and nitrate are formed (*cf. Nitrous acid and Nitrites*, p 567). Gay Lussac (*G A* 58, 29) says that KNO_2 is formed by allowing O, mixed with a large excess of NO, to stand over $KOHAq$, and that the gases disappear in the ratio required to form N_2O_3 .—3 Decomposes *urea* evolving N.—4 With *benzenoid primary amido compounds*, forms diazo-compounds (*q v. vol 1 p 397*). With *paraffinoid amido compounds*, the general result is to substitute OH for NH_2 , and evolve N.—5 With *conc sulphuric acid*, nitrosyl sulphate (SO_2OHONO) is formed. As NO_2 reacts with *conc* H_2SO_4 to form SO_2OHONO and HNO_3 , and as NO passed into H_2SO_4 containing NO_2 forms SO_2OHONO and H_2O , it is sometimes argued that the reaction of N_2O_3 with H_2SO_4 is easily explained by the view that regards N_2O_3 as a mixture of NO and NO_2 , if this view is held, then the two reactions

- (1) $2NO_2 + H_2SO_4 = SO_2OHONO + HNO_3$;
- (2) $HNO_3 + 2NO + 3H_2SO_4$

= SSO_2 , $\text{OH ONO} + 2\text{H}_2\text{O}$ must proceed simultaneously and at equal rates, and the supposed mixture must never contain more NO than is shown by the ratio NO NO_2 , otherwise some NO would pass on unabsorbed. According to Lunge (*lc*) (confirmed by Groves, *C S Proc* 1, 24), NO only slowly converts H_2SO_4 containing HNO_2 into SO_2 , OH ONO .

Combinations—According to Weber (*P* 118, 471) N_2O_5 combines with SnCl_4 and TiCl_4 , e.g. to form $\text{SnCl}_4\text{N}_2\text{O}_5$.

NITROGEN DIOXIDE NO_2 , and TETROXIDE N_2O_4 (Nitrogen peroxide Nitroso nitric anhydride) The mol w of the compound N_2O_4 varies with temperature, at low temperatures and small pressure (c -12° at 115 mm) the mol w is 91.86 = N_2O_4 , and at moderately high temperatures (c 150°) the mol w is 45.93 = NO_2 (*v infra*). N_2O_4 melts at -10° (Deville & Troost, *C R* 64, 257, Ramsay, *C J* 57, 590). Boils at 21.64° (760 mm, Thorpe, *C J* 87, 224) SG (liquid) $d_4^{20} = 1.4903$ (Thorpe, *lc*) V.D. at -12° and 115.4 mm = 42.54 (Natanson, *W* 27, 606), at 130° a 718 mm = 23.26 (Richardson, *C J* 51, 397), at 183° a 760 mm = 22.7 (Deville & Troost, *C R* 64, 237), at 27° a 16 to 35 mm = 23.1 (Troost, *C R* 86, 1395) SH (liquid) 46 (Ramsay, *C J* 57, 590), for SH at different temperatures *v* Berthelot & Ogier (*A Ch* [5] 30, 382). For heat of fusion *v* Ramsay (*lc*) H.F. $[\text{N}_2\text{O}_4] = -2.005$, $[\text{NO}_2] = 19.570$, $[\text{N}_2\text{O}_4\text{Aq}] = 5.750$ (*Th* 2, 199). For electrical resistance of liquid N_2O_4 *v* Boguski (*Z P C* 5, 69). For absorption-spectrum *v* Kundt (*P* 141, 157), Gernez (*C R* 74, 465), Luck (*Z* [2] 6, 237). For thermal expansion *v* Thorpe (*C J* 87, 224).

Molecular weights of nitrogen peroxide—The V.D. at low temperatures near the liquefying point corresponds with the formula N_2O_4 , and the same formula expresses the molecular composition at c 27° under a pressure of 16 to 35 mm, when heated to c 150° the V.D. shows that the molecular formula is NO_2 . Measurements of V.D. are given by Deville & Troost (*C R* 64, 257), Troost (*C R* 86, 1395), Playfair & Wanklyn (*C J* 15, 156), E & L Natanson (*W* 27, 606), Richardson (*C J* 51, 397). That the change in V.D. measures a process of dissociation from N_2O_4 to NO_2 has been shown by Natanson (*lc*), Naumann (*A Suppl* 6, 205, *B* 11, 2045), Salet (*C R* 67, 488). The amount of dissociation is increased by raising temperature or lowering pressure (*v* especially Natanson, *lc*). Ramsay determined the lowering of the freezing point of acetic acid by dissolving N peroxide in it (*C J* 53, 621), and also the lowering of f.p. of $\text{C}_2\text{H}_5\text{Cl}$ and CHCl_3 by the peroxide (*C J* 57, 590), both results tend to show that the mol w of the liquid compound is N_2O_4 .

Formation—1 By passing electric sparks through a mixture of O and N—2 By bringing NO into contact with excess of air or O—3 By heating $\text{Pb}(\text{NO}_3)_2$ —4 By the action of $\text{AgNO}_3 + \text{NO}_2\text{Cl}$ (Exner, *C C* 1872 278), but the existence of NO_2Cl is denied by Williams (*C J* 49, 226).—5 According to Hasenbach (*J pr* [2] 4, 1), by passing O into the liquid obtained by heating starch with HNO_3Aq and condensing, but the observations of Ramsay & Cundall (*C J* 47, 187) show that O very slowly, if at all, combines

with N_2O_4 —6 By the reaction between NOCl and KNO_3 (Girard & Pabst, *Bl* [2] 30, 351).

Preparation—1 A mixture of dry NO and about an equal volume of dry O is passed over dry KOH, and then into a vessel cooled to -20° , solid N_2O_4 is thus obtained (Péligot, *A* 9, 259, 39, 327, cf Dulong, *G A* 58, 53, a Lunge, *C J* 47, 465)—2 Dry powdered $\text{Pb}(\text{NO}_3)_2$ is heated in a tube of hard glass, and the gas is passed into a U tube cooled to c -8° , a stream of dry O is passed through the liquid in the U tube for some time (Péligot, *lc*, Thorpe, *C J* 87, 224)—3 By very gently heating As_2O_3 with HNO_3Aq SG 15 and a little conc H_2SO_4 , and passing O into the liquid thus obtained (cf Ramsay & Cundall, *C J* 47, 187, with Lunge, *B* 11, 1641)—4 By distilling fuming HNO_3 at a low temperature, and fractionating the distillate in a stream of dry O (Fritzsche, *J pr* 22, 21) Setliff (*C C* 1888 461) has determined the yields of NO_2 obtained by the various processes.

Properties—At temperatures below -10° a white, crystalline (? prismatic), deliquescent solid, between -10° and 21.6° a liquid, which is nearly colourless at -9° , yellow at 0° , and orange at 21° , at temperatures above 21.6° , a gas, the colour of which deepens from orange yellow to nearly black as temperature rises. At c 180° – 200° decomposition to NO + O begins and is complete at c 620° (Richardson, *C J* 51, 397, cf Reactions, No 1). The colourless liquid is N_2O_4 , as this dissociates into NO_2 , the colour becomes deeper and deeper red (*v* Salet, *C R* 67, 488, also *v supra*). The gaseous compound at ordinary temperatures is a mixture of NO_2 and N_2O_4 , this gas is reddish brown, has a pungent odour, is irrespirable, and stains the skin yellow. NO_2 dissolves in cold water, forming HNO_3Aq and HNO_2Aq , it is dissolved, apparently without change, by CHCl_3 , $\text{C}_2\text{H}_5\text{Cl}$, CS_2 (Friedburg, *C N* 47, 52), and $\text{C}_2\text{H}_5\text{NO}_2$. As solution of NO_2 in HNO_3 does not show the absorption-lines of gaseous and liquid N_2O_4 , it is probable that the solution in HNO_3 contains a compound or compounds of the two substances (Gernez, *C R* 74, 465). NO_2 probably combines with NO at low temperatures to form N_2O_3 (cf Combinations, No 1).

NO_2 is an acidic oxide, inasmuch as it reacts with H_2O to form HNO_3Aq and HNO_2Aq , because of this reaction it may be looked on as nitro-nitric anhydride, or perhaps better (as there is no nitroso nitric acid) as a mixed anhydride. There are no salts corresponding with NO_2 . Numerous nitro compounds are known, obtained by substituting the monovalent radicle NO_2 for H.

Reactions—1 Heat dissociates N_2O_4 to 2NO_2 (*v* beginning of this article). At c 180° a 755 mm decomposition to NO + O begins, at c 400° about 33 p.c. of the NO_2 is decomposed, at c 500° 60 p.c. is decomposed, and at c 620° the change is complete, on cooling, the NO + O recombine to form NO (Richardson, *C J* 51, 397)—2 Electric sparks are said to decompose NO_2 to N and O, according to Richardson (*C J* 51, 402) a very small quantity of a white solid is produced—3 NO_2 dissolves in water, added to a little ice-cold H_2O a green solution is obtained, which becomes blue on dilution, and then colour-

less, the solution gives the reactions of HNO_3 and HNO_2 . On warming NO_2Aq , or on adding NO_2 to hot water, NO is evolved and HNO_2Aq remains ($3\text{NO}_2 + \text{H}_2\text{O} + \text{Aq} = 2\text{HNO}_2\text{Aq} + \text{NO}$). The cold colourless solution of NO_2 in water may be raised to the boiling-point without a complete decomposition of the HNO_2Aq to HNO_3Aq and NO , addition of Pt wire or powdered glass, &c., causes decomposition of NO_2Aq with evolution of NO . NO_2Aq decomposes HIAq and iodides in solution with separation of I , it also deoxidises KMnO_4Aq —4 With aqueous solution of an alkali NO_2 produces a nitrite and a nitrate—5 Sulphuric acid in solution absorbs NO_2 , evolving NH_3 , and depositing S —6 NO_2 passed into conc sulphuric acid forms HNO_3 and nitrosyl sulphate (lead chamber crystals, SO_2OHONO) (*v* Lunge, *D P J* 233, 65)—7 NO_2 dissolves in very conc nitric acid, forming a yellow liquid, which is probably not merely a solution of NO_2 in HNO_3 , inasmuch as it does not show the absorption-lines of gaseous or liquid N_2O_4 , whereas a solution in CS_2 does show these lines (*v* Gernez, *C R* 74, 465)—8 NO_2 reacts with many carbon compounds replacing H by NO_2 , with alcoholic iodides it usually produces nitrates of the alcoholic radicle, e.g. $\text{C}_2\text{H}_5\text{I} + \text{N}_2\text{O}_4 = \text{C}_2\text{H}_5\text{NO}_2 + \text{NO} + \text{I}$ —9 With cold boron chloride, crystals BCl_2NOCl are formed (Geuther, *J pr* [2] 8, 854)—10 Carbon monoxide is said to be partly oxidised to CO_2 , while part of the CO combines with NO_2 —11 When NO_2 is mixed with hydrogen, and the mixed gases are passed over spongy Pt, H_2O and NH_3 are formed—12 Ordinary combustibles are extinguished in NO_2 , but strongly burning phosphorus and carbon continue to burn—13 Many metals decompose NO_2 , forming oxides and N , Fe , Cu , &c., at a red heat, K and Na at the ordinary temperature—14 Liquid N_2O_4 reacts with mercury to form HgNO_2 and NO without any nitrite, the equation $2\text{N}_2\text{O}_4 + 2\text{Hg} = 2\text{HgNO}_2 + 2\text{NO}$ is almost realised quantitatively (Divers a Shimidzu, *C J* 47, 630) With silver, liquid N_2O_4 forms AgNO_3 , copper probably forms $\text{Cu}_2(\text{NO}_3)_2$ (*D a S*, 1c)—15 Liquid N_2O_4 reacts with liquid sulphur dioxide to form $(\text{NO}_2)_2\text{S}_2\text{O}_4$ (De la Provostaye, *A Ch* [3] 73, 362)—16 With sulphuric anhydride, NO_2 gas forms $\text{NO}_2\cdot\text{S}_2\text{O}_4$, O NO_2 (Weber, *P* 123, 337)

Combinations—1 With nitric oxide, at low temperatures, to form a blue liquid which is chiefly N_2O_3 (Dulong, *A Ch* 2, 317, Peligot, *A* 39, 327, Ramsay, *C J* 57, 590, Richardson, *C J* 51, 397) Ramsay a Cundall (*C J* 47, 672) found that no contraction occurs when dry NO_2 and NO are mixed at the ordinary temperature. According to Hasenbach (*J pr* [2] 4, 1), N_2O_3 is formed by passing NO and NO_2 through a hot tube, and then into a well-cooled receiver—2 With chlorine, and bromine, to form NO_2Cl and NO_2Br , respectively (Gay-Lussac, *A Ch* [3] 23, 203, Landolt, *A* 116, 177, but denied by Williams, *C J* 49, 226)—3 With antimony chloride, to form $8\text{SbCl}_3\cdot\text{N}_2\text{O}_4$, obtained by heating in a sealed tube at 100° (Besson, *C R* 108, 1012)—4 NO_2 also combines with the chlorides of aluminium, antimony (SbCl_3), bismuth, and iron, the compounds are decomposed by water, and also by a gentle heat (*B*, 1c)—5 With phosphorus pentfluoride, to form crystals of $\text{NO}_2\cdot\text{PF}_5$, very

easily decomposed (Tassel, *C R* 110, 1264)—6 With magnesium hydrogen phosphate, to form $2\text{MgHPO}_4\cdot\text{NO}_2$ (Luck, *Fr* 1874 255)—7 With amylene (and some other hydrocarbons) to form $\text{C}_6\text{H}_{10}(\text{NO}_2)_2$

Constitution of N_2O_4 —From the reactions of liquid N_2O_4 with Hg and Ag , whereby HgNO_3 , or AgNO_3 , and NO are produced, Divers a Shimidzu conclude that N_2O_4 is best represented as NO NO_2 (*C J* 47, 630) This formula is in keeping with the reaction $\text{AgO NO} + \text{NO}_2\text{Cl} = \text{AgCl} + \text{NO ONO}_2$ (Exner, *C C* 1872, 273), this argument, however, assumes AgNO_2 to have the structure AgO NO (*v Nitrites*, p 567), it also assumes the existence of NO_2Cl The formation of nitrosyl sulphate by the reaction of N_2O_4 with H_2SO_4 is also in keeping with the formula NO NO_2 , $\text{NO NO}_2 + \text{H}_2\text{SO}_4 = \text{NO HSO}_4 + \text{HNO}_3$ Also, if nitrous acid is assumed to be HO NO , then the formation of this acid and HNO_2 by the reaction of N_2O_4 with water points to the formula NO ONO_2 ($\text{NO ONO}_2 + \text{H}_2\text{O} = \text{NO OH} + \text{HO NO}_2$) The formation of diazo benzene nitrate by the reaction of N_2O_4 with amido benzene is in keeping with the formula NO ONO_2 , thus $\text{C}_6\text{H}_5\text{NH}_2 + \text{NO ONO}_2 = \text{NO ON N C}_6\text{H}_5 + \text{H}_2\text{O}$ The formation of $\text{C}_6\text{H}_5\text{ONO}_2$ by the reaction of N_2O_4 with $\text{C}_6\text{H}_5\text{I}$ is best expressed by writing N_2O_4 as NO ONO_2 , ($\text{C}_6\text{H}_5\text{I} + \text{NO ONO}_2 = \text{C}_6\text{H}_5\text{ONO}_2 + \text{I} + \text{NO}$) The fact that $\text{C}_6\text{H}_{10}(\text{NO}_2)_2$, when reduced by Sn and HClAq yields NH_4Cl and not an NH_2 compound, induced V Meyer (*A* 171, 5) to assign to N_2O_4 the formula NO_2NO_2 and to write $\text{C}_6\text{H}_5(\text{NO}_2)_2$ as $\text{C}_6\text{H}_5(\text{ONO}_2)_2$, because had the C been in direct union with N , an NH_2 compound would have been formed on reduction N_2O_4 is dissociated by heat to 2NO_2 , this reaction seems to point to the constitution $\text{O}_2\text{N NO}_2$ or perhaps to $\text{NO O}_2\text{NO}$ Whichever formula is adopted, the reactions of N_2O_4 make it evident that some substances separate the group NO_2 , and others the group ONO , and that the way in which the molecule N_2O_4 splits up depends largely on the conditions of the reaction

NITROGEN PENTOXIDE N_2O_5 (*Nitric anhydride*) Mol w uncertain, as compound has not been gasified

Formation—1 By passing Cl over AgNO_3 ($2\text{AgNO}_3 + \text{Cl}_2 = 2\text{AgCl} + \text{N}_2\text{O}_5 + \text{O}$)—2 By dehydrating conc HNO_3 by P_2O_5 ($2\text{HNO}_3 + \text{P}_2\text{O}_5 = 2\text{HPO}_3 + \text{N}_2\text{O}_5$)—3 By the reaction of NO_2Cl on AgNO_3 at 60° – 70° ($\text{AgNO}_3 + \text{NO}_2\text{Cl} = \text{AgCl} + \text{N}_2\text{O}_5$, Odet a Vignon, *C R* 69, 1142, 70, 96, but the existence of NO_2Cl is very doubtful, *v* Williams, *C J* 49, 226)

Preparation—1 Perfectly dry AgNO_3 is placed in a dry U tube, which is connected by a glass tube, without corks or caoutchouc, with another dry U tube having a small reservoir at the bottom, the U tube containing AgNO_3 is immersed in a water bath, and the other U tube is surrounded by ice and salt, a very slow stream of dry CO_2 is passed through the apparatus, the AgNO_3 being kept at c 180° (to expel every trace of moisture), a very slow stream of dry Cl is then passed from a gasholder (the Cl being stored over conc H_2SO_4 and driven out by the inlet of H_2SO_4), the AgNO_3 being heated to 95° , and the temperature then lowered to 58° – 68° . After a time crystals of N_2O_5 appear

in the cooled U tube, while a little liquid (N_2O_4 or N_2O_5) collects in the small reservoir. Not more than 3 to 4 litres Cl should pass in 24 hours. To transfer the N_2O_4 to another vessel, the Cl is replaced by a current of dry CO_2 , the U tube is connected by a glass tube with the vessel, which is immersed in a freezing mixture, the U tube is removed from its freezing mixture, and the N_2O_4 is poured into the vessel (Deville, *A Ch* [3] 28, 241). — 2 Very conc HNO_3 is surrounded by ice and salt, rather more P_2O_5 than the wt of HNO_3 is added little by little, so that the temperature does not rise above 0° , the thick mass is brought into a wide retort and distilled at as low a temperature and as slowly as possible, the retort being cooled if the liquid froths, pure N_2O_4 condenses in the cooled receiver, but towards the end of the reaction liquid $2N_2O_5 \cdot H_2O$ is formed (Berthelot, *Bl* [2] 21, 53, a modification of the process given by Weber, *J pr* [2] 6, 342). About 80 grams N_2O_4 are obtained from 150 gr HNO_3 .

Properties and Reactions—Very lustrous, translucent, rhombic prisms, melts at 30° , with partial decomposition to N_2O_4 and O , and boils between 45° and 50° . N_2O_4 should be kept in stoppered bottles over H_2SO_4 . At $c 15^\circ$ the crystals become yellowish, but are colourless when kept in a freezing mixture (Weber, *lc*). $SG c 1.64$. $H F [N_2O_4] = -31,600$ (Berthelot, *A Ch* [5] 6, 145), $[N_2O_4, Aq] = 29,820$ (*Th* 2, 199), N_2O_4 decomposes slowly when kept, rapidly in sunlight, giving N_2O_4 and O (Berthelot, *Bl* [2] 21, 53). Burning P burns brightly in slightly warmed N_2O_4 , C burns in the vapour of N_2O_4 , but not in the solid or liquid compound, K burns brightly, but most of the metals are without action on N_2O_4 , with S there is formed $(NO)_2S_2O_8$ (Weber, *lc*). N_2O_4 reacts with water to form HNO_3 . With conc HNO_3 , it produces $H_2N_2O_{11}$ ($= 2N_2O_4 \cdot H_2O$, or $N_2O_4 \cdot 2HNO_3$), *v Dinitric acid*, under Nitric acid, p 520. The reactions of N_2O_4 point to the constitution $NO_2 \cdot O \cdot NO_2$.

PERNITRIC OXIDE NO_3 or N_2O_5 . According to Hautefeuille a Chappuis (*C R* 92, 80, 134, 94, 1111, 1306), a very unstable compound of this composition is obtained by the action of the silent electric discharge on a mixture of N and O at a low temperature. When a certain quantity is formed, the substance begins to decompose to NO_2 and O . Berthelot (*Bl* [2] 35, 227) says that the compound is obtained by the action of the induction discharge on a mixture of O and NO_2 . The compound is described as a liquid which does not solidify in a freezing mixture of ice and salt, and is extremely unstable, decomposing to NO_2 and O .

Nitrogen, oxyacids. The only oxyacids of N which have been isolated are nitric (HNO_3) and dinitric ($H_2N_2O_{11}$), nitrous acid (HNO_2) and hyponitrous acid ($H_2N_2O_4$) are known in aqueous solution. Nitric and dinitric acids are described under the heading Nitric acid (p 517). The present article contains descriptions of nitrous and hyponitrous acids and their salts.

Nitrous acid and Nitrites, HNO_2 and MNO_2 . Nitrites are usually, if not always, accompanied by nitrates in nature. Nitrites are present in the atmosphere, Warington (*C J* 39, 229) has shown that when distilled water is exposed to the air it soon gives the reactions of

nitrites. Small quantities of nitrites are present in many river and well waters (v Warington, *C J* 55, 537, Munro, *C J* 49, 632). The juices of many plants contain nitrites (Genadius, *Am Ch* 5, 7). Nitrites are sometimes, but not generally, found in saliva (Wurster, *B* 22, 1901). Nitrites are formed in the soil by oxidation of various nitrogenous compounds (v NITRIFICATION, p 521). $NaNO_2$ is formed by exposing Pt black to air and $NaOHAq$ (Loew, *B* 23, 1443). NH_4NO_2 is produced by passing O and air over Pt black at 180° – 300° (Ilosva, *Bl* [3] 2, 734). NH_4NO_2 is also produced by burning H in air (Struve, *J* 1870 199, 209, Schönbein, *J* 1862 94, Zöller a Grete, *B* 10, 2145), and also during the slow combustion in air of P or ether (Berthelot, *A Ch* [5] 12, 440, *C R* 108, 543, Kolbe, *A* 119, 176, Ilosva, *Bl* [8] 2, 734). It was formerly stated that NH_4NO_2 is formed by the direct union of N and H_2O (by evaporating H_2O in air), but this has been disproved (Carus, *A* 174, 31, Weith a Weber, *B* 7, 1745). Berthelot, however, asserts that NH_4NO_2 is produced by subjecting H_2O and pure N to the action of a very powerful induction coil (*Bl* [2] 27, 338). NH_4NO_2 is also formed by the oxidation of NH_3 by ozone (Carus, *lc*, Goppelsröder, *J pr* [2] 4, 139, 338). Fe nitrite is said to be formed by passing air over reduced Fe at 190° – 250° (Ilosva, *Bl* [3] 2, 734). Alkali nitrites are formed, along with nitrates, by the reaction of N_2O_4 on alkali solutions (v Nitrogen trioxide, *Reactions*, No 2, p 564). Nitrous acid, along with HNO_3 , is produced by the action of cold water on N_2O_5 (v Nitrogen trioxide, *Reactions*, No 1, p 564). Nitrites are also formed by the reduction of nitrates, *eg* by heating KNO_3 , O is evolved and KNO_2 remains, or by the action of amalgamated Zn on KNO_3 , Aq , KNO_2 , Aq is produced.

The normal nitrites have the composition MNO_2 and $M'(NO_2)_2$, several basic nitrites are known. Most of the normal nitrites are soluble in water and alcohol, $AgNO_2$ is one of the least soluble nitrites, from it most of the other nitrites may be obtained. Nitrites are decomposed by heat, evolving NO or NO_2 , and leaving metallic oxide or metal, solutions of nitrites are decomposed by boiling, generally with evolution of NO and formation of nitrates. Nitrites in solution are decomposed by dilute H_2SO_4 , giving HNO_2 , which soon decomposes to HNO , Aq and NO . Acidulated solutions of nitrites reduce $KMnO_4$, Aq , KCr_2O_7 , Aq , H_2SAq , &c., they ppt Au from $AuCl_3$, Aq , and Hg from mercurous solutions. These solutions decompose $KIAq$ with separation of I , and give a brown colour with $FeSO_4$, these reactions serve to distinguish nitrites from nitrates. (For details regarding the detection and estimation of nitrites, a manual of analysis must be consulted.) Nitrites may be regarded either as $MO \cdot NO$ or as MNO_2 , Divers (*C J* 47, 226) endeavours to show that the second view is preferable.

Nitrous acid HNO_2 . This acid is known only in aqueous solution, and it is doubtful whether HNO_2 , Aq has been obtained free from HNO_3 , Aq . Frey (*C R* 79, 61) says that a solution of N_2O_4 in a considerable quantity of cold water may be kept for some days without change, and that on boiling NO and N_2O_4 are

evolved. The solution very probably contains HNO_2 , but whether it is free from HNO_3 or not is undecided. This solution acts as a strong reducer. Thomson gives the thermal data $[\text{N}_2^0, \text{O}^0, \text{Aq}] = -6,820$, $[\text{H}_2\text{N}_2\text{O}^0, \text{Aq}] = 80,770$, $[\text{2NO}, \text{O}^0, \text{Aq}] = 86,880$, $[\text{H}_2\text{NO}, \text{O}^0, \text{Aq}] = 52,845$ (*Th* 2, 199).

Nitrites MNO_2 , and $\text{M}^{\text{IV}}(\text{NO}_2)_2$, also basic salts, $x\text{MO} \cdot y\text{N}_2\text{O}_5$, and $x\text{M}_2\text{O}_3 \cdot y\text{N}_2\text{O}_5$. The nitrites have been examined chiefly by Fischer (*P* 74, 116), Lenz (*P* 118, 282), Hampe (*A* 125, 295), Stromeyer (*A* 96, 230), Lang (*J* pr 86, 299).

Ammonium nitrite NH_4NO_2 . A crystalline mass, decomposed by heat to N and H_2O (*v* Nitrogen, Preparation, No 2, p 557). Obtained by decomposing NH_4ClAq by AgNO_3Aq . Berthelot (*Bl* [2] 21, 55) says this salt is formed, along with N, by the action of dry NH_3 on NO and O. It was formerly stated that NH_4NO_2 is produced by evaporating water in air free from NH_3 , but this was disproved by Carius (*A* 174, 81) and by Weith & Weber (*B* 7, 1745). According to Berthelot (*Bl* [2] 27, 238) NH_4NO_2 is produced by the action of a powerful induction-coil on a mixture of H_2O and pure N. NH_4NO_2 is also formed by oxidising NH_3Aq by ozone (Carius, *lc*, Goppebröder, *J* pr [2] 4, 139, 383), and also, along with ozone and H_2O_2 , by burning H in air (Struve, *J* 1870 199, 209, Zöllner & Grete, *B* 10, 2145).

Barium nitrite $\text{Ba}(\text{NO}_2)_2 \cdot \text{H}_2\text{O}$ (Fischer)

Cadmium nitrite $\text{Cd}(\text{NO}_2)_2 \cdot \text{H}_2\text{O}$ (Hampe, Lang).

Calcium nitrite $\text{Ca}(\text{NO}_2)_2 \cdot \text{H}_2\text{O}$ (Fischer, Hampe, Lang).

Cobalt nitrite. This salt is not known, but several double salts of $\text{Co}(\text{NO}_2)_2$ have been isolated.

Cobalt-potassium nitrite

$2\text{Co}(\text{NO}_2)_2 \cdot 6\text{KNO}_3 \cdot x\text{H}_2\text{O}$ (x varies from 0 to 1, 3, and 4). Prepared by mixing KNO_3Aq and $\text{Co}(\text{NO}_2)_2\text{Aq}$, or CoCl_2Aq , adding excess of acetic acid, washing the pp with K acetate solution, and then with 80 p c alcohol, and drying below 100° . A bright yellow powder, consisting of small 4 sided prisms, very slightly sol water, insol alcohol or ether. At 200° gives NO_2 , Co_2O_3 , and KNO_2 (Fischer, *P* 67, 245, Saint-Evre, *J* pr 54, 85, 58, 185, Braun, *Fr* 6, 42, 7, 313, Stromeyer, *A* 96, 220, Erdmann, *J* pr 97, 385, Sadtler, *Am S* [2] 49, 189). Various other Co-K nitrites are formed under different conditions of concentration and acidification (*v* especially Sadtler, *lc*). If Ca salts are present a triple nitrite of Co, Ca, and K is ppd (Erdmann, *lc*). A triple nitrite of Co, Pb, and K is described by Stromeyer (*lc*). Erdmann (*lc*) also describes a compound of $\text{Co}(\text{NO}_2)_2$, KNO_2 , and NH_4 .

Cobalt-cassium nitrite

$\text{Co}(\text{NO}_2)_2 \cdot 3\text{CsNO}_2 \cdot \text{H}_2\text{O}$ (Rosenblatt, *B* 19, 2531).

Cobalt-sodium nitrites

$2\text{Co}(\text{NO}_2)_2 \cdot 4\text{NaNO}_2 \cdot \text{H}_2\text{O}$ and $2\text{Co}(\text{NO}_2)_2 \cdot 6\text{NaNO}_2 \cdot \text{H}_2\text{O}$ (Sadtler, *lc*).

Copper nitrites, basic salts (Hampe, van der Meulen, *B* 12, 758).

Lead nitrites. These salts were examined by Prout, Chevreul (*G* 4, 176), and Berzelius (*G* 4, 194, 46, 166). Péligot (*A* 39, 838) revised and classified the work, Bromers (*A* 72, 88), von Lorenz (*W* 4, B [2nd part] 84

1133), and Meissner (*J Z* [2] 3, 26), more recently have examined these salts. The lead nitrites are all basic salts, many seem best looked on as compounds of basic nitrites with basic nitrates. They are obtained by digesting $\text{Pb}(\text{NO}_3)_2\text{Aq}$ with Pb, von Lorenz says that 14 different salts are thus obtained.

Magnesium nitrite $\text{Mg}(\text{NO}_2)_2 \cdot 3\text{H}_2\text{O}$ (Lang); $\text{Mg}(\text{NO}_2)_2 \cdot 2\text{H}_2\text{O}$ (Hampe).

Mercuric nitrite $\text{Hg}(\text{NO}_2)_2 \cdot 2\text{H}_2\text{O}$ (Lang).

Nickel nitrite $\text{Ni}(\text{NO}_2)_2$ (Lang). The double salts, $\text{Ni}(\text{NO}_2)_2 \cdot 4\text{KNO}_3$, $\text{Ni}(\text{NO}_2)_2 \cdot 2\text{KNO}_3 \cdot \text{Ba}(\text{NO}_3)_2$, $\text{Ni}(\text{NO}_2)_2 \cdot \text{Ca}(\text{NO}_3)_2 \cdot 2\text{KNO}_3$ have been isolated (Erdmann, *J* pr 97, 385).

Platino-nitrites *v* PLATINUM, vol 1v.

Potassium nitrite KNO_2 . Colourless, microscopic, prismatic crystals (according to Lang, *J* pr 86, 295, the crystals are $2\text{KNO}_2 \cdot \text{H}_2\text{O}$), very sol water, insol absolute alcohol. KNO_3Aq with H_2SO_4 evolves NO , and forms KNO_3Aq . KNO_2 is prepared by passing into cold KOHAq the gas obtained by heating starch with HNO_3Aq S G 135, evaporating till crystallisation begins, pouring off from the crystals of KNO_2 , adding dilute acetic acid and 2 vols alcohol, separating the lower layer which forms and evaporating it (Fischer). Stahlschmidt (*P* 128, 466) reduces KNO_3Aq by Zn in presence of NH_4Aq . Erdmann (*J* 1866 154) reduces KNO_3 by fusion with Fe, and crystallises from water.

Double salts of KNO with the nitrites Ba, Cd, Ca, Co, Cu, Pb, Mg, Hg, Ni, Pd, Ag, Sr, and Zn are described by Hampe, Lang, and Fischer.

Silver nitrite AgNO_2 . Obtained by adding AgNO_3Aq to the solution produced by passing into KOHAq the gas formed by heating starch with HNO_3Aq S G 135, after neutralising this solution by acetic acid. Small white crystals. Dissolves in 300 pts water at ordinary temperatures, easily in boiling water. The decomposition of AgNO_2 by heat has been examined by Divers & Shimidzu (*C J* 47, 630). Treated with H_2S , AgNO_2 yields Ag_2S , S, NO, NH_3 , and NH_4OH (Divers & Haga, *C J* 51, 48).

Sodium nitrite NaNO_2 . A crystalline salt, prepared similarly to KNO_2 . Liard (*Bl* [2] 27, 434) recommends to fuse NaNO_3 with an equivalent quantity of K_2SO_4 and to extract with alcohol.

Strontium nitrite $\text{Sr}(\text{NO}_2)_2$ (Hampe).

Zinc nitrite $\text{Zn}(\text{NO}_2)_2 \cdot 3\text{H}_2\text{O}$ (Lang, Hampe).

HYPONITROUS ACID and **HYPONITRITES**, $\text{H}_2\text{N}_2\text{O}_3\text{Aq}$ and $\text{M}_2\text{N}_2\text{O}_3$. Hyponitrites were first obtained by Divers in 1871, by reducing nitrates by Na-amalgam (*Pr* 19, 425). The salts were examined by Zorn (*B* 10, 1306, 11, 1630, 2217, 12, 1509, 15, 1007, 1258), van der Plaats (*B* 10, 1507), Menke, (*C J* 38, 401), Berthelot & Oger (*C R* 96, 80, 84). In 1884 Divers and Haga (*C J* 45, 78) showed that the silver salt is AgNO . The investigation has been continued by Divers and Haga (*C J* 47, 364, 55, 760), Dunstan and Dymond (*C J* 51, 646), and Berthelot and Maquenne (*C R* 108, 1286, 1303).

Zorn prepared ethyl hyponitrite and determined its molecular formula (from VD) to be $(\text{C}_2\text{H}_5)_2\text{N}_2\text{O}_3$, hence the formula for the salts is better written $\text{M}_2\text{N}_2\text{O}_3$ than MNO .

Hyponitrites are produced (1) by reducing

nitrate or nitrite in solution by Na amalgam (Divers), (2) by electrolyzing nitrite solution, using Hg as negative electrode (Zorn, Divers a Haga), (3) by the reaction of $\text{Fe}(\text{OH})_2$ on alkaline nitrite solution, or on NO in presence of alkali (Zorn, Dunstan a Dymond), (4) by decomposing oxysulphates [salts of $\text{NH}(\text{SO}_3\text{H})\text{OH}$] by alkali (Divers)

Silver hyponitrite $\text{Ag}_2\text{N}_2\text{O}_2$ (*Nitrosyl silver*)

Preparation—1 Na amalgam is added to KNO_3Aq in the ratio $\text{KNO}_3, 4\text{Na}$, the liquid being kept cold. When evolution of gas ceases the solution is neutralised by acetic acid, and AgNO_3 is added, the pp is washed in the dark with cold water, dissolved in cold dilute HNO_3Aq , and reprecipitated by $\text{Na}_2\text{CO}_3\text{Aq}$, the pp is thoroughly washed in the dark and dried *in vacuo* over H_2SO_4 (Divers, *Pr* 19, 425, D a Haga, *C J* 45, 78)—2 Zorn (*B* 12, 1509) reduces $\text{Ba}(\text{NO}_3)_2$ by Na amalgam, and pptd by AgNO_3Aq , the pp of $\text{Ag}_2\text{N}_2\text{O}_2$ thus obtained is pure. The $\text{Ba}(\text{NO}_3)_2$ is obtained by boiling $\text{Ba}(\text{NO}_3)_2\text{Aq}$ with Pb, prepared by ppg dilute Pb acetate solution by Zn, the boiling is continued till all the Pb is changed to PbO , the PbO in solution is removed by CO_2 , and the last traces by H_2S , the filtrate is evaporated to a thick oil, which cools to a mass of $\text{Ba}(\text{NO}_3)_2$, the salt is obtained pure by crystallising from 80 p.c. alcohol. For preparing large quantities of $\text{Ag}_2\text{N}_2\text{O}_2$, Zorn (*l.c.*) places the Na amalgam, in large pieces, in the middle bulb of a Kipp's apparatus, the $\text{Ba}(\text{NO}_3)_2\text{Aq}$ being in the lower bulb, and the upper half fills with distilled water. By regulating the stopcocks so that a slow stream of gas escapes, the reduction proceeds satisfactorily—3 Pure FeSO_4Aq is mixed with enough milk of lime to ppt the Fe as $\text{Fe}(\text{OH})_2$, NaNO_3Aq is added (1 part NaNO_3 to 10 parts FeSO_4), and the apparatus is kept cold. When the reaction is finished the solution is pptd by AgNO_3Aq (Zorn, *B* 15, 1258, cf Dunstan a Dymond, *C J* 51, 646)

Properties and Reactions— $\text{Ag}_2\text{N}_2\text{O}_2$ is a yellowish amorphous salt. By standing in NH_3Aq over H_2SO_4 , it is obtained in small crystals (Zorn). The salt is not hygroscopic, insoluble, may be kept in boiling water without decomposition, is decomposed at c. 100° , probably giving AgNO_3 , when quickly heated to c. 150° , $\text{Ag}_2\text{N}_2\text{O}_2$ explodes, evolving brown vapours. $\text{Ag}_2\text{N}_2\text{O}_2$ dissolves in HNO_3Aq and $\text{H}_2\text{SO}_4\text{Aq}$, it is pptd by NH_3Aq or $\text{Na}_2\text{CO}_3\text{Aq}$. Unacted on by CO_2 . Decomposed by NaOHAq at c. 70° . Decomposed by $\text{H}_2\text{PO}_4\text{Aq}$, H_2S , or boiling $\text{H}_2\text{C}_2\text{H}_3\text{O}_2\text{Aq}$, with production of $\text{H}_2\text{N}_2\text{O}_2\text{Aq}$. $\text{Ag}_2\text{N}_2\text{O}_2$ reacts with $\text{C}_2\text{H}_5\text{I}$ to form $(\text{C}_2\text{H}_5)_2\text{N}_2\text{O}_2$ (Zorn, *B* 11, 1630), the formula of which compound is molecular, as its V.D. has been determined.

Barium hyponitrite BaN_2O_4 is described by Zorn (*B* 15, 1007)

Calcium and Strontium hyponitrites

$\text{CaN}_2\text{O}_4 \cdot 4\text{H}_2\text{O}$ and $\text{SrN}_2\text{O}_4 \cdot 5\text{H}_2\text{O}$ are described by Maquenne (*C R* 108, 1808)

Hyponitrous acid $\text{H}_2\text{N}_2\text{O}_2\text{Aq}$ Known only in solution. Moist $\text{Ag}_2\text{N}_2\text{O}_2$ is suspended in water, such a quantity of dilute HClAq is added that a little $\text{Ag}_2\text{N}_2\text{O}_2$ remains unchanged, and the liquid is filtered from AgCl (van der Plaats, *B* 10, 1507). $\text{H}_2\text{N}_2\text{O}_2\text{Aq}$ is an acid liquid, fairly stable, may be boiled with HNO_3Aq or

$\text{H}_2\text{C}_2\text{H}_3\text{O}_2\text{Aq}$ without decomposition, reduces KMnO_4Aq , and separates I from KIAq . The solution slowly decomposes, and after a few weeks not a trace of $\text{H}_2\text{N}_2\text{O}_2$ remains. The liquid thus obtained is neutral, probably N_2O and H_2O are formed. It was proved by van der Plaats (*l.c.*) that conc. H_2SO_4 evolves N_2O from $\text{H}_2\text{N}_2\text{O}_2\text{Aq}$.

Constitution of hyponitrites—The compound $(\text{C}_2\text{H}_5)_2\text{N}_2\text{O}_2$ reacts not as an ethereal salt but as a diazo compound, with reducers it evolves N and forms $\text{C}_2\text{H}_5\text{OH}$, water produces $\text{C}_2\text{H}_5\text{OH}$, CH_3CHO , and N. The constitution of this compound is $\text{C}_2\text{H}_5\text{O} \cdot \text{N} \cdot \text{N} \cdot \text{O} \cdot \text{C}_2\text{H}_5$, and hence the constitution of the acid is probably $\text{HO} \cdot \text{N} \cdot \text{N} \cdot \text{OH}$ (Zorn, *B* 11, 1630). This formula is confirmed by the production of hyponitrites, along with sulphates, by the action of KOHAq on $\text{NH}(\text{SO}_3\text{K})\text{OH}$. This reaction shows that the O in hyponitrites is in direct union both with N and H (Divers a Haga, *C J* 55, 760)

Nitrogen, oxybromide of, NOBr (*Nitrosyl bromide*) Mol. w. not determined directly, probably = 109.72, from analogy with NOCl . A very dark brown liquid. Obtained by passing NO into Br at -7° to -15° (Landolt, *A* 116, 177), also by distilling nitrosyl sulphate (NOHSO_4) with KBr (Girard a Pabst, *Bl* [2] 30, 531). Landolt gives b.p. as -2° , Girard a Pabst as 19° . Easily decomposed to NO and Br by heat. With cold water forms HBrAq and HNO_3Aq , with KOHAq forms KBrAq and KNO_3Aq .

Nitryl tribromide Landolt (*A* 116, 117) supposed that a compound NOBr_3 is obtained (along with NOBr) by passing Br into well-cooled Br. That the amount of NO absorbed by Br, at c. 15° , is nearly independent of pressure, and corresponds approximately with the quantity required to form NOBr_3 , was shown by Pattison Muir (*C J* 28, 844). The experiments of Frohlich (*A* 224, 270), and of Roozeboom (*R T C* 4, 381) have shown that the so called nitryl tribromide is probably only a mixture of NOBr and Br.

Nitrogen, oxychlorides of The compound NOCl has been isolated, the existence of NO_2Cl is very doubtful, according to Hautefeuille a Ohappuis (*A Ch* [6] 2, 282), a compound $\text{N}_2\text{Cl}_2\text{O}_2$ is formed by passing N with O and Cl through an induction apparatus. The non existence of NOCl_2 , said by Gay Lussac to be produced by the reaction of conc. HClAq with HNO_3 (*A Ch* [3] 23, 203), has been proved by Tilden (*C J* 27, 639), and this has been confirmed by Goldschmidt (*A* 205, 372).

NITROSYL CHLORIDE NOCl (*Chloronitrous gas*) Mol. w. 65.34 V.D. 33 at c. 15° to 700° (Sudborough a Millar, *C J* 59, 78) S.G. at -18° 1.433, at -12° 1.4165 (Geuther, *A* 245, 97, Tilden, *C J* 27, 630). Boils at c. -8° (Tilden). An orange red gas, liquefied by passing through a U tube cooled by ice and salt, the liquid is reddish yellow. Dissociation into NO and Cl begins at c. 780° , at 985° about 42 p.c. is dissociated (S a M, *l.c.*) For absorption spectrum v. Magnanini (*Z P C* 4, 427). Formed by combination of NO with Cl (Gay-Lussac, *A Ch* [3] 23, 203), by the reaction of PCl_3 , PCl_5 , or AsCl_3 , with NO, or N_2O , and by the reaction of POCl_3 with KNO . (Naquet, *J* 1860 102), by passing

HCl into N_2O_4 at a low temperature by heating conc HClAq and HNO_3 , and by heating NO H SO₄ with NaCl

$NOCl$ is more readily prepared by warming HNO_3 Aq SG 1.42 with c 4 vols HClAq SG 1.16, passing the gases over $CaCl_2$ and then into conc H_2SO_4 until the acid is saturated, then adding perfectly dry NaCl, and heating gently (Tilden, *C J* 27, 630, cf Girard a Pabst, *Bl* [2] 80, 631)

$NOCl$ reacts with H_2O to form HClAq and HNO_3 Aq, with KOHAq it produces KClAq and KNO_3 Aq, Hg decomposes it, forming NO and $HgCl_2$, it dissolves Au and Pt more slowly (Tilden, *lc*) With conc H_2SO_4 , nitrosyl sulphate ($NOHSO_4$) is formed (Tilden) $NOCl$ reacts energetically with SO_2 to produce NO_2 , SO_2 , Cl (Weber, *P* 123, 333) Forms double compounds with many metallic chlorides, e.g. $AlCl_3$, $SbCl_3$ (v Sudborough, *C J* 59, 655)

NITRYL CHLORIDE NO_2Cl A compound with this composition was supposed to be formed by the reaction of $POCl_3$ with $AgNO_3$ or $Pb(NO_3)_2$, also by passing a very slow stream of Cl over $AgNO_3$ at c 95° (Odet a Vignon, *C R* 69, 1142, 70, 96), Hasenbach obtained a liquid which he took to be NO_2Cl by passing Cl and NO_2 through a hot tube and cooling the issuing gases (*J pr.* [2] 4, 1), Williamson (*Pr* 7, 16) poured SO_2 , OH Cl on to KNO_3 , and obtained a gas which formed HNO_3 Aq and HClAq with cold water, and was supposed by W to be NO_2Cl , Muller (*A* 122, 1) supposed that NO_2Cl was formed by the reaction of NO_2 with PCl_5 , and Schiff thought he had obtained the compound as a product of the reaction of HNO_3 with PCl_5 , Meissner (*J Z* 10, 27) failed to obtain any NO_2Cl by the processes used by Odet a Vignon, Schiff, Muller, and Williamson Geuther (*A* 245, 96) could not form NO_2Cl by the reaction of PCl_5 with HNO_3 , NO_2 , or a nitrate Williams (*C J* 49, 222) showed that NO_2Cl is not produced by the reaction (1) of $POCl_3$ with $Pb(NO_3)_2$, (2) of $POCl_3$ with HNO_3 , (3) of SO_2 , OH Cl with KNO_3 , (4) of SO_2 , Cl_2 with KNO_3 , he also established a very large probability in favour of the conclusion that the products of the reaction of NO_2 with Cl , at different temperatures, are $NOCl$, and N_2O , holding more or less Cl in solution

Nitrogen, phosphide of, v PHOSPHORUS NITRIDE

Nitrogen, phosphochloride of, $N_3P_2Cl_4$ (*Phosphorus chloronitride Nitrogen chlorophosphide Phosphorus nitrogen chloride*) Mol. w 847.13 V D 176.7 (Gladstone a Holmes, *C J* 17, 225, Wichelhaus, *B* 3, 163) SG 1.98 (G a H) Melts at c 110° (G a H), at 114° (W); boils at 240° (G a H), at 250° – 260° (W) Crystallises in trimetric forms, a b c = 4417.1 1.8165 (Groth, *B* 3, 166), c/a = 109.9. (G a H)

Formation—By heating PCl_5 with NH_4Cl , NH_3 , or NH_2HgCl

Preparation—1 PCl_5 is saturated with dry NH_3 , the product is distilled with water, and the crystals which form on the sides of the receiver are washed, dried, and recrystallised from hot ether (Wöhler a Liebig, *A* 11, 146)—2 A mixture of 1 pt. PCl_5 and 2 pts. NH_4Cl is heated in

a flask connected with a reversed condenser; the mass gets red, then brown, and the $N_3P_2Cl_4$ sublimes, the compound is dissolved out in ether and crystallised, or is blown over in steam (Gladstone, *C J* 3, 185, modified by Coultbridge, *C J* 53, 398) The yield is not more than 10 p c of the PCl_5 used—3 An intimate mixture of white pp (NH_2HgCl) and PCl_5 is gently heated in a flask, the product is treated with water, which dissolves out $HgCl_2$ and NH_4Cl , the residue is dried and the $N_3P_2Cl_4$ is dissolved out (from PCl_5 , N_2H_4) by ether (Gladstone a Holmes, *C J* 17, 225)

Properties—Hard, lustrous, trimetric, crystals, sol alcohol, ether, or $CHCl_3$, insoluble in water, but slowly decomposed (G a H, W) Gives off white fumes when heated in air, may be sublimed in H or H_2S Solution in alcohol slowly decomposes, also that in ether if a trace of water is present (even in absence of water, according to Wichelhaus) Is not acted on by I (G), nor by hot H_2SO_4 Aq, HClAq, or HNO_3 Aq (Wöhler a Liebig), decomposed by hot fuming HNO_3 (G)

Reactions—1 Water forms pyrophosphoric acid ($P_2N_2H_6O_8$) and HCl, according to G a H, but W says the products are many and complicated—2 Ammonia or potash decomposes $N_3P_2Cl_4$ in alcoholic solution, the reaction being similar to that of water (G a H)—3 Ammonia gas passed over melted $N_3P_2Cl_4$ forms phospham $nPN(NH_2)$, and HCl (Coultbridge, *C J* 53, 398)—4 Heated with *aniline*, $P_2N_2(NHCH_2)_4$ is formed (Hofmann, *B* 17, 1909, Coultbridge, *lc*), *ortho toluidine*, and *phenylhydrazine* produce similar reactions (G, *lc*)—5 Nascent hydrogen forms PH_3 (W, *lc*)—6 Heated with copper oxide, NO_2 and N are evolved—7 Passed over red hot iron, N is evolved and $FeCl_3$ and Fe phosphide remain (W a L)

Nitrogen, selenide of ($?NSe$) Espenschied (*A* 113, 101) passed NH_3 , diluted with H, over well cooled $SeCl_4$, shook the product with water, and washed the ppd mixture of Se and N selenide with CS_2 (to extract Se), he thus obtained an orange-yellow powder, which was extremely explosive and very dangerous to handle Analyses pointed to the formula NSe , but E supposed that the substance probably contained H besides N and Se For details of preparation and properties v Espenschied (*lc*)

Nitrogen, silicide of, v SILICON NITRIDE, in vol iv

Nitrogen, sulphide of, nNS Mol w unknown SG 2.1166 at 15° (Michaëlis, *Z* [2] 6, 460), 2.22 at 15° (Berthelot, *A Ch* [5] 27, 202) Sublimes at 135° , melts at 158° , and decomposes with slight explosions at 160° (M, *lc*, at 207° according to B, *lc*) Berthelot (*lc*) gives $[N,S] = -31,900$ N sulphide was prepared, but not pure, by Soubeiran (*A Ch* [2] 67, 71), Fordos and Gélis obtained it approximately pure (*A Ch* [3] 32, 385), it has been examined further by Michaëlis (*Z* [2] 6, 460), and Demarcay (*C R* 91, 854, 1066, 92, 726) N sulphide easily explodes when rubbed or struck with a hammer

Preparation—1 SO_2 or S_2Cl_2 is dissolved in 8–10 vols CS_2 , and dry NH_3 is passed in, the liquid darkens in colour, and a cochineal red pp is produced, which dissolves after a time and a

brown powder is *ppd.*, passage of NH_3 is continued until the brown *pp* dissolves, and the liquid becomes orange-yellow with a few nearly colourless flocks of NH_4Cl floating in it (addition of more NH_3 decomposes the *NS* in solution), the liquid is filtered and allowed to evaporate, when orange red *NS* separates, the *NS* is collected and washed with CS_2 , to remove traces of *S*. Omitting intermediate products, the reaction may be represented approximately as $8\text{NH}_3 + 3\text{SCl}_2 = 2\text{NS} + \text{S} + 6\text{NH}_4\text{Cl}$ (F & G) — 2 SOCl_2 is surrounded by cold water, and NH_3 is passed into it, after a time the cold water is removed, the solid mass is mixed by a glass rod, and NH_3 is passed over it as long as absorption continues, the nearly white mass thus obtained is treated with CS_2 , and the solution on evaporation deposits *NS*, the *insol* in CS_2 consists of NH_4Cl , $(\text{NH}_4)_2\text{SO}_4$, and polythionates of NH_4 (Michaelis).

Properties — Orange-red, rhombic (or ? triclinic) crystals, with a faint odour, which becomes more marked at $c\ 120^\circ$, the vapour rapidly attacks the mucous membrane of the nose and eyes, detonates at $c\ 160^\circ$, explodes violently when rubbed or struck by a hammer, adheres strongly to glass or paper when rubbed thereon, is electric *insol*, but slowly decomposed by water, slightly *sol* alcohol, ether, turpentine, CS_2 is the best solvent, this solution slowly decomposes.

Reactions — 1 *Water* slowly decomposes *NS*, forming NH_4Aq , $(\text{NH}_4)_2\text{S}_2\text{O}_8\text{Aq}$, and $(\text{NH}_4)_2\text{S}_2\text{O}_4\text{Aq}$ — 2 *Potash* forms NH_3 , $\text{K}_2\text{S}_2\text{O}_8\text{Aq}$, and $\text{K}_2\text{SO}_4\text{Aq}$ — 3 *Hydrogen chloride* produces NH_4Cl , chlorides of *S*, and probably a compound of *NS* with SCl_2 — 4 *Chlorine* reacts on *NS* in CHCl_3 with formation of a compound of *S* chloride and *NS*, Demarcay (C R 31, 854) gives the formula SNCl to the product, and says that addition of *NS* to this produces $(\text{SN})_2\text{Cl}$ — 5 A large excess of *sulphur chloride* reacting on *NS* in CHCl_3 produces a compound of *S*, *N*, and *Cl*, to which Demarcay (C R 91, 1066) assigns the formula $\text{S}_2\text{N}_2\text{Cl}$ — 6 *Nitric acid* forms $\text{S}_2\text{N}_2\text{NO}_3$, and *sulphuric acid* produces HCl and $\text{S}_2\text{N}_2\text{HSO}_4$ (D, l c, also C R 92, 726).

Combinations — With *sulphur dichloride* to form several compounds, especially 2NS SCl_2 , 4NS SCl_2 , and 6NS SCl_2 (Michaelis).

Nitrogen, sulphochloride of NS_2Cl ($=\text{N}_2\text{S}_2\text{SCL}_2$) According to Soubeiran (A Ch 87, 87, 101), this substance is formed by heating, in a sealed tube at 100° , the compound $2\text{NH}_4\text{SCl}_2$, which is obtained by passing a slow stream of NH_3 into SCl_2 . NS_2Cl is a citron-yellow solid, decomposed by heat to *N*, *S*, and S_2Cl_2 , decomposed by water to NH_4Cl and $\text{H}_2\text{S}_2\text{O}_8$. NS_2Cl is also said to be formed by passing CO_2 into a hot solution of *NS* in S_2Cl_2 (S., l c). M M P M

NITROGEN GROUP OF ELEMENTS *Nitrogen, phosphorus, vanadium, arsenic, molybdenum, antimony, didymium, erbium, tantalum, bismuth*. Of these ten elements, *N*, *As*, *Sb*, and *Bi* are found uncombined, the others occur only in combination. *N* is found in vast quantities in the air, compounds of *N* and also of *P* occur in large quantities in rocks and in animal and vegetable matter. Compounds of *As*, *Sb*, and *Bi* are widely distributed in minerals, but no very large quantities are found. Compounds of *V* are found widely spread about, but they occur only in comparatively small quantities. The compounds of *Nb*, *Di*, *Er*, and *Ta* have been found in but a few minerals, these four bodies are classed among the rare elements. *Sb* has been known as a metal from about the end of the fifteenth century, and *Bi* from the sixteenth century, *P* was prepared in 1669, *As* in 1694, and *N* in 1772, *V* was obtained by Roscoe in 1867, investigations on this element having been carried on from the beginning of this century, *Di* was isolated in 1842, *Nb* and *Ta* were obtained nearly pure in 1864, after a long series of investigations conducted by different chemists from 1801 onwards, researches on *Er* have been carried on since 1788 to the present day, but the element has not yet been isolated. Within recent years great doubt has been cast on the elementary character of *Di*, and although the investigation of *Er* is yet far from complete it is very probable that the substance known by this name is not a simple body. Of the names given to the ten elements, *As* is derived from the name by which the chief ore of this metal was known in ancient times, *N*, *P*, and *Di* express characteristic properties of the elements, *Er* is derived from the locality where the minerals were found from which the compounds of this element have been prepared, *V*, *Nb*, and *Ta* are taken from mythological personages, and the origins of the names *Bi* and *Sb* are not known with certainty.

N is obtained from air by removing the *O* by hot *Cu*, or other deoxidiser, *P* is prepared by heating *Ca* phosphate with SiO_2 to combine with the *Ca*, and *C* to remove the *O*, *As*, *Sb*, and *Bi* are prepared by removing *O* from the oxides by *C*, *V* and *Nb* by reducing the chlorides at a high temperature by *H*, and *Di* by reducing the chloride by *Na* or *K*, to obtain *Ta*, the compound Na_2TaF_6 is reduced by *Na*, *Er* has not yet been isolated. *N* is a gas at ordinary temperatures, but has been liquefied at a very low temperature under great pressure, *P* is a soft solid with a low melting point and not high *S G*, the other elements are hard, lustrous, and metal like, none has a very high *S G*, *Ta* = 11 (?) is the heaviest, and all melt at or below a full red heat (*M P* of *V*, *Nb*, and *Ta*, not determined, *Er* not isolated). The following table presents some of the chief properties of the ten elements —

	NITROGEN	VANADIUM	NIOBIUM	DIDYMIUM	TANTALUM
Atomic weights	14.01	51.2	94	144	182
One or more compound of each element, except <i>Di</i> , has been gasified, specific heat of <i>Di</i> only has been determined directly. Mol. w of $\text{N}=\text{At. w} \times 2$; mol. w of none of the others known.					
Melting points	—	—	—	600°–700°	—

	NITROGEN	VANADIUM	NIOBIUM	DIDYMIUM	TANTALUM
<i>Specific gravities (approx)</i>	(liquid) 885	5.5	7	6.5	11 (?)
<i>Specific heats</i>	—	—	—	0.4563	—
<i>Atomic weight</i>	15.8	91	134	221	166(?)
<i>Spec grav (approx)</i>					
<i>Occurrence and preparation</i>	In large quantity in air. Many compounds, especially nitrates and NH_3 compounds, are widely distributed in large quantities. Prepared by removing O from air.	Vanadates of Pb, Ca, Bi, &c occur as comparatively rare minerals. Small quantities of V compounds are very widely distributed. Prepared by long continued heating VCl_2 in H.	Niobates occur in a few rare minerals. Prepared by repeatedly heating NbCl_5 in H.	Silicate of D ₁ is found in several rare minerals. Prepared by heating D_1Cl_5 with K, or by electrolyzing molten D_1Cl_5 .	Tantalates occur with niobates in a few rare minerals. Prepared by heating K_2TaF_7 with K, or Na_2TaF_6 with Na.
<i>Physical properties</i>	Colourless, tasteless, odourless gas, liquefied at very low temp and great pressure, liquid boils at -195°	Grey, lustrous crystalline powder	Steel grey, lustrous solid	White solid, with slightly yellow tinge, hard, ductile	Grey, lustrous, solid, not obtained free from impurities
<i>Chemical properties</i>	Combines slowly with O under influence of electric discharge, also to a small extent with H. At high temp combines with B, Cr, Mg, Si, and V, and perhaps also with Al, Fe, and Zn. Compounds exhibit most diverse chem properties, oxides are generally acidic, one hydride is strongly alkaline, another is acidic. Never replaces H of acids to form salts. Forms the oxychloride NOCl . An allotropic form perhaps exists, proof is not satisfactory. Atom is trivalent in NH_3 .	Burns in air to V_2O_5 , then to V_2O_3 , and finally to V_2O_4 . Burns in Cl to VCl_5 . Combines with N to form VN. Dissolved by conc H_2SO_4 and HNO_3 , forms vanadates with molten alkalis. Some oxides are basic, e.g. V_2O_3 , others are basic and acidic, e.g. V_2O_5 , which forms salts with acids and also with alkalis. Reacts both as a metal and a non-metal. Forms oxychlorides VOCl_2 , VOCl_3 , and VOCl_4 . Does not form a hydride. Atom is tetravalent in VCl_4 , perhaps trivalent in VOCl_2 .	Burns in air to Nb_2O_5 . Combines with Cl to NbCl_5 . Sol in conc H_2SO_4 , not in HNO_3 or HClAq . Nb_2O_5 forms niobates with basic oxides, also dissolves in H_2SO_4 , but no definite salts isolated. No hydride known with certainty, but there are indications of existence of hydride. Forms a nitride. Oxychloride is NbOCl_2 . Compounds formed by replacing H of acids by Nb not yet isolated, but probably exist NbF_5 and NbOF_3 form many double salts. Atom pentavalent in NbCl_5 .	Burns in air to D_1O_5 , D_2O_5 exists and acts as a peroxide. D_1O_5 is basic, easily soluble in dilute acids. Decomposes cold water slowly, and hot water rapidly. D ₁ is almost certainly a mixture or compound of two or more elements. D_1OCl is isolated. No hydride. No compounds gasified.	Burns in air to Ta_2O_5 . Combines with Cl to form TaCl_5 . Ta_2O_5 forms tantalates with alkalis, seems to have no basic properties. Sol only in HFAq , and H_2SO_4 + HFAq . Forms a nitride. No hydride known. No oxychlorides isolated. Atom pentavalent in TaCl_5 .

	PHOSPHORUS	ARSENIC	ANTIMONY	ERBIUM	BISMUTH
<i>Atomic weights</i>	30.96	74.9	120	166	208
<i>Molecular weights</i>	61.92 and 123.84	149.6 and 299.6	(?) 120 (?) 240	—	(?) 208 or (?) 416
	Compounds of all these elements, except Er, have been gasified, and S H of each, except Er, has been determined directly				
<i>Melting-points (approx)</i>	45°	500° (under pressure)	425°	—	270°
<i>Specific gravities (approx)</i>	1.9	5.7	6.7	—	9.9
<i>Specific heats</i>	202	0.83	0.53	—	0.308
<i>Atomic weight</i>	16.3	13.2	18	—	20.5
<i>Spec grav</i>					
<i>Occurrence and preparation</i>	Many phosphates occur in large quantities and widely distributed in rocks and waters, also in bones and plants. Compounds of P with C, N, and O, are found in nerve and brain matter. Prepared by heating $\text{Ca}_3(\text{PO}_4)_2$ with SiO_2 and C.	Is found native in small quantities, compounds with S and other elements are widely distributed, but are not found in very large quantities. Prepared by reducing As_2O_3 by C.	Sb found native in small quantity, Sb_2S_3 occurs in comparatively small quantities, fairly widely distributed, other compounds also occurs sparingly. Prepared by reducing Sb_2O_3 by C.	As silicate in a few rare minerals. Er has not yet been isolated.	Bi found native, chief compounds are Bi_2S_3 and Bi_2O_3 , not found in any large quantities, but fairly widely distributed. Prepared by reducing Bi_2O_3 by C.
<i>Physical properties</i>	Soft, wax like, crystalline solid, also a red amorphous solid. Non conductor of electricity.	Grey, hard, brittle, crystalline, solid, also as a black amorphous powder. Fair conductor of electricity.	White, lustrous, brittle, very crystalline, solid. Fair conductor of electricity.	Not isolated.	White, with slightly reddish tinge, brittle, crystalline, not good conductor of electricity.
<i>Chemical properties</i>	Burns in air to P_2O_3 and P_2O_5 . Combines directly with Cl, Br, and I, to form PX_3 and PX_5 . Oxidised by HNO_3 to H_3PO_4 . Oxides are anhydrides, several oxy-acids known. Does not form salts by replacing H of acids. Hydride PH_3 resembles NH_3 , but is less alkaline. Exists in two allotropic forms. Atom trivalent and pentavalent in gaseous molecules.	Burns in air to As_2O_3 , As_2O_5 , also known. Combines directly with Cl, Br, and I, to form AsX_3 . Oxidised by HNO_3 to H_3AsO_4 . Oxides are acidic, and As_2O_3 is also feebly basic. Does not form salts by replacing H of acids. Hydride AsH_3 is scarcely if at all alkaline. Exists in two allotropic forms. Atom trivalent in gaseous molecules.	Burns in air to Sb_2O_3 , Sb_2O_5 , also known. Combines directly with Cl, Br, and I, to form SbX_3 and SbX_5 . Oxidised by HNO_3 to Sb_2O_3 and $\text{Sb}_2\text{O}_5 \cdot x\text{H}_2\text{O}$. Oxides are acidic and basic. Forms a few salts by replacing H of acids. Decomposes steam, evolving H. Hydride SbH_3 is not alkaline. Atom trivalent in gaseous molecules.	Reactions of Er not known, as the metal has not yet been isolated. Very probably Er is a mixture or compound of two or more elements. Er_2O_3 is basic, no acidic oxide known. No hydride known. No compound yet gasified.	Burns in air to form Bi_2O_3 , Bi_2O_5 , also known. Combines directly with Cl, Br, and I, to form BiX_3 . Oxidised by HNO_3 to Bi_3NO_5 . Oxides are basic, Bi_2O_3 acts as a basic peroxide with perhaps very feeble acidic properties. Many salts formed by replacing H of acids by Bi. Decomposes steam, evolving H. No hydride isolated. Atom trivalent in gaseous molecules.

General formulae and characters of chief compounds MH_3 , $M = N, P, As, Sb, Bi, M_2O_3$, $M =$ any element of the group except Nb and Ta M_2O_5 , $M = N, P, V, Nb, Sb, Ta, Bi, M_2O_3$, $M =$ any of the elements except Er M_2S_3 , $M = P, V, As, Sb, Bi, Bi, M_2S_3$, $M = P, V, As, Sb$, some other sulphides known, *e.g.* NS, VS, TaS_3 , no sulphide of Nb or Er isolated MCl_3 , $M =$ any of the elements except Er and Ta MCl_3 , $M = P, Nb, Sb, Ta$, some other haloid compounds exist, *e.g.* P_2I_4, VCl_4 , no haloid compounds of Er isolated with certainty HMO, H_2MO, H_3MO, H_4MO , most of these acids except for $M = N, P, V, As, Sb$, acids containing Nb and Ta also known, no acids of Di, Er , or Bi isolated M_2SO_3 , &c, $M = Di, Er, Bi, VO(SO_3), (VO_2)_2(SO_3)_2$, &c $As_2O_3, Sb_2O_3, Sb_2O_5, As_2O_5$, &c

The *hydrides* NH_3, PH_3, AsH_3 , and SbH_3 show a gradation of properties, from the strongly alkaline NH_3 to the neutral AsH_3 and SbH_3 , NH_3 combines readily with acids, PH_3 only with such strong acids as HI or HCl , AsH_3 and SbH_3 do not combine with acids AsH_3 and SbH_3 are easily decomposed by heat, while NH_3 and PH_3 are stable in this respect The hydrides are all oxidised by mixing with O and heating, NH_3 being the most difficult to change in this way A hydride of Nb (? NbH) probably exists N forms also the hydrides N_2H and N_2H_4 , the former is a fairly strong acid, the latter is basic Besides PH_3 , two hydrides of P, *viz.* P_2H_2 and P_2H_4 , exist

Regarded broadly, the *oxides* may be divided into three classes (1) *acidic oxides*, those of N and P, (2) *basic*, those of Di, Er , and Bi , (3) *acidic and basic*, those of V, As, and Sb—acidic and ?basic, oxides of Nb and Ta The distinctly acidic oxides of N are N_2O, NO_2 , and N_2O_5 , with water N_2O forms HNO_2Aq , and N_2O_5 forms HNO_3Aq , but NO_2 produces both HNO_2 and HNO_3 , N_2O may be called the anhydride of $H_2N_2O_5Aq$, inasmuch as it is formed by heating $H_2N_2O_5Aq$, but the acid has not been obtained from N_2O NO can hardly be classed as acidic or basic, there are compounds in which NO may be regarded as playing the part of the more positive radicle, *e.g.* $NOCl$ and $NOHSO_4$, and there are others in which NO seems to form the negative radicle, *e.g.* $(NO)_2H_2$, and perhaps $NOOH, P_2O_5$, and P_2O_5 react with water to form H_2PO_4Aq and H_3PO_4Aq respectively, P_2O_5 forms both of these acids The oxides M_2O_3 , where $M = Di, Er$, or Bi , react with acids to form salts M_2X ($X = SO_3, 2NO_2$, &c), Bi_2O_3 forms salts Bi_2X and evolves O_2 , this oxide probably forms bismuthates— Bi_2O_3, xM_2O —when fused with large excess of alkalis, but these bismuthates cannot be isolated V_2O_5 probably forms salts with acids, the compound $V_2O_5(SO_3)_2, H_2O$ has been isolated V_2O_5 with strong acids forms salts xV_2O_5, yA ($A =$ acidic oxide, SO_3 , &c), and with alkalis it produces salts of the type xV_2O_5, yM_2O, V_2O_5 reacts with alkalis to produce vanadates xV_2O_5, yM_2O , it also combines with several anhydrides to form salts xV_2O_5, yA ($A =$ acidic oxide, P_2O_5, SO_3 , &c), the acids HVO_3 and $H_2V_2O_7$ have been isolated As_2O_3 does not form an acid with water, but with $KOHAq$ it produces $KAsO_3, As_2O_3$ with water

forms H_2AsO_4, As_2O_3 combines with a few anhydrides of strong acids to form such compounds as xAs_2O_3, ySO_3 Neither Sb_2O_3 nor Sb_2O_5 forms an acid with water, a few unstable salts Sb_2O_3, M_2O have been isolated, three weakly acidic hydrides of Sb_2O_3 are known, from each of which salts are derived With acidic oxides Sb_2O_3 combines to form xSb_2O_3, yA ($A =$ acidic oxide, SO_3 , &c), some of these compounds are fairly well-defined salts, *e.g.* $Sb_2O_3, 3SO_3$ The oxides Nb_2O_5 and Ta_2O_5 form salts when fused with alkalis, xM_2O, yM_2O , these oxides dissolve in some strong acids, probably with formation of salts, although none has yet been isolated

The *oxyacids* of the nitrogen elements are numerous, oxyacids of all except Di, Er , and Bi are known The table on p 575 presents the composition of the most marked of the acids, and the relations between them, their corresponding oxides, and their salts, the symbol Aq added to the formula of an acid means that that acid is known only in aqueous solution, M here stands for a monovalent metal, $RO =$ basic oxide generally, including M_2O, MO, M_2O_3

The acids of N and P possess the characteristics of acids much more fully than any of the other oxyacids of the N elements $HNO_2, HNO_3, HPO_3, H_3PO_4, H_2P_2O_7, H_4P_2O_7, H_4AsO_4$, these acids are produced by the reaction of their corresponding oxides with water—the other acids of the N elements are not obtained directly from their oxides, although in many cases the oxides are formed by heating the acids HNO_2Aq is a very strong acid, about equal to $HClAq$, putting the strength or affinity of HNO_2Aq as 100, that of H_3PO_4Aq is approximately about 6, and that of H_2AsO_4Aq about 4 One can scarcely give the name *acid* to the hydrated oxides of Nb and Ta, and it is very doubtful whether $Sb_2O_3, 3H_2O$ ($= H_2SbO_3$) can be called an acid It should be remembered that the oxides of Nb and Ta have not been very fully examined

The chief *haloid compounds* of the nitrogen elements are MX_3 and MX_5 , no haloid compound of Er has been isolated with certainty, TaX_5 is not known, and pentahaloid compounds of N, V, As, Di, and Bi have not yet been prepared V forms VCl_4 , which has been gasified unchanged Most of the haloid compounds are formed by the direct union of their elements, they are decomposed by water, forming $HXAq$, and oxyacids or hydrated oxides in the cases of N, P, V, As, Nb, and Ta, and oxyhaloid compounds in the cases of Di (? Er) and $Bi, SbCl_5$ gives $SbOCl$ if little water is used, but Sb_2O_3Aq if much warm water is employed

The *sulphides*, M_2S_3 , of P, V, As, and Sb react with alkali sulphides to form alkali thio-salts, Di_2S_3 and Bi_2S_3 show no acidic properties, Er and Nb sulphides have not been isolated, and the sulphides of Ta have been studied but slightly, NS is an explosive compound, which forms NH_3 salts of thionic acids when treated with water or potash

The nitrogen elements form Group V of the elements in the classification based on the periodic law This group is composed as follows —

Even series	N = 14	V = 51	Nb = 94	Di = 144	Ta = 182
Odd series	P = 81	As = 75	Sb = 120	Er = 166	Bi = 208

Acid	Salts	Corresponding oxide	Remarks
<i>Nitrogen</i>			
$\text{H}_2\text{N}_2\text{O}_2\text{Aq}$	$\text{M}_2\text{N}_2\text{O}_2$	N_2O	Acid not formed from N_2O , but N_2O obtained by heating $\text{H}_2\text{N}_2\text{O}_2\text{Aq}$, and by action of H_2SO_4 on $\text{M}_2\text{N}_2\text{O}_2\text{Aq}$
HNO_2Aq	MNO_2	N_2O_3	Acid obtained by dissolving N_2O_3 in fair quantity of cold water, solution slowly decomposes to HNO_2Aq and NO . Doubtful whether HNO_2Aq has been obtained quite free from HNO_3 .
HNO_3	MNO_3 , also M_2NO_4 , and many basic nitrates $x\text{N}_2\text{O}_5, y\text{RO}$, also probably a few acid salts $\text{M}'\text{NO}_3, x\text{N}_2\text{O}_5$	N_2O_5	N_2O_5 reacts with water to form HNO_3Aq , N_2O_5 obtained by removing H_2O from HNO_3 by P_2O_5 .
$\text{H}_2\text{N}_2\text{O}_{11}$	None	N_2O_5	Acid formed by adding very conc HNO_3 to melted N_2O_5 , and cooling
<i>Phosphorus</i>			
$\text{H}_2\text{H}_2\text{PO}_3$	$\text{M H}_2\text{PO}_3$	None	Acid obtained by action of equivalent quantity of $\text{H}_2\text{SO}_4\text{Aq}$ on $\text{Ba}(\text{H}_2\text{PO}_3)_2$, which is produced by P reacting with BaO Aq . H_2PO_3 decomposes by heat to H_3PO_3 and PH_3 .
H_3PO_3	M HPO_3 and M_2PO_3	None	H_3PO_3 obtained by H_2S on PbPO_3 in water and evaporation <i>in vacuo</i> . $\text{H}_2\text{PO}_3\text{Aq}$ fairly stable, boiled with $\text{H}_2\text{SO}_4\text{Aq}$ forms $\text{H}_2\text{PO}_4\text{Aq}$ and $\text{H}_3\text{PO}_4\text{Aq}$.
$[\text{H}_2\text{H}_2\text{P}_2\text{O}_5]$	$\text{M}_2\text{H}_2\text{P}_2\text{O}_5$	$\text{P}_2\text{O}_4(?)$	Acid unknown. Na salt got by heating $\text{NaH}_2\text{P}_2\text{O}_5, 5\text{H}_2\text{O}$ to 160° . Pb salt got by heating $\text{PbH}_2(\text{HPO}_3)_2$ to 140° <i>in vacuo</i> .
H_4HPO_3	M HPO_3 and $\text{M H}_2\text{PO}_3$	P_2O_5	Acid obtained by slow action of cold water on P_2O_5 , decomposed by heat to H_3PO_3 and PH_3 .
$\text{HPO}_3, \text{H}_2\text{PO}_4, \text{H}_3\text{P}_2\text{O}_7$	$\text{M PO}_3, \text{M}_2\text{PO}_4, \text{M HPO}_3, \text{M H}_2\text{PO}_4, \text{M}_2\text{P}_2\text{O}_7, \text{M}_2\text{H}_2\text{P}_2\text{O}_7$	P_2O_5	Acids obtained by action of water on P_2O_5 . A little cold water gives HPO_3 , cold water and P_2O_5 in ratio $\text{P}_2\text{O}_5, 2\text{H}_2\text{O}$ give H_2PO_4 , much water gives $\text{H}_3\text{P}_2\text{O}_7$, H_3PO_4 heated gives $\text{H}_2\text{P}_2\text{O}_7$, and this at higher temperature gives HPO_3 . P_2O_5 is not obtained by heating the acids.
<i>Vanadium</i>			
$\text{HVO}_3, \text{H}_4\text{V}_2\text{O}_7$	$\text{MVO}_3, \text{M}_2\text{V}_2\text{O}_7$, also salts of types $\text{M}_2\text{VO}_4, \text{M}_2\text{V}_2\text{O}_7, \text{M}_2\text{V}_2\text{O}_{11}$, &c	V_2O_5	Acids not obtained directly from the oxide, but V_2O_5 obtained by heating the acids.
<i>Arsenic</i>			
—	$\text{MAsO}_2, \text{M}_2\text{AsO}_4$	As_2O_3	As_2O_3 in water does not form an acid, but reacts with alkalis &c to form salts.
$\text{HAsO}_3, \text{H}_2\text{AsO}_4, \text{H}_4\text{As}_2\text{O}_7$	$\text{MAsO}_3, \text{M}_2\text{HAsO}_4, \text{M}_2\text{AsO}_4, \text{M}_2\text{As}_2\text{O}_7$	As_2O_5	H_2AsO_3 obtained by action of H_2O on As_2O_5 , H_2AsO_4 gives $\text{H}_4\text{As}_2\text{O}_7$, and at higher temperature HAsO_3 is formed; on heating $\text{HAsO}_3, \text{As}_2\text{O}_5$, and H_2O are produced.

Acid	Salts	Corresponding oxide	Remarks
Niobium $x\text{Nb}_2\text{O}_5, y\text{H}_2\text{O}$ (? acids)	$x\text{Nb}_2\text{O}_5, y\text{RO}$, some be long to types MnNbO_5 and $\text{M}_2\text{Nb}_2\text{O}_7$, others are more complex	Nb_2O_5	Hydrates of Nb_2O_5 are known, but salts are not obtained by reaction of these with alkalis, but by fusing Nb_2O_5 with alkalis and alkaline carbonates
Antimony H_3SbO_4 (? acid)	MSbO_3	Sb_2O_3	H_3SbO_4 is not obtained directly from Sb_2O_3 , but Sb_2O_3 is formed by heating H_3SbO_4 . H_3SbO_4 scarcely exhibits acidic properties. MSbO_3 are few, produced by dissolving Sb_2O_3 in alkalis
—	$\text{M}_2\text{Sb}_2\text{O}_7$ (? $\text{MSbO}_3, \text{MSbO}_3$)	? Sb_2O_4	Said to be formed by fusing Sb_2O_3 with alkalis and alkaline carbonates
$\text{HSbO}_3, \text{H}_3\text{SbO}_4, \text{H}_4\text{Sb}_2\text{O}_7$	$\text{MSbO}_3, \text{M}_2\text{H}_2\text{Sb}_2\text{O}_7, \text{M}_4\text{Sb}_2\text{O}_7$	Sb_2O_5	Acids are not obtained directly from Sb_2O_5 , but oxides are formed by heating the acids. H_3SbO_4 gives HSbO_3 at 175° . $\text{H}_4\text{Sb}_2\text{O}_7$ is obtained from its salts, it is easily decomposed to HSbO_3 . HSbO_3 dissolves in KOH aq, but does not form salts thus. MSbO_3 obtained by fusing Sb_2O_3 or HSbO_3 with alkalis or alkaline carbonates. $\text{M}_2\text{Sb}_2\text{O}_7$ are formed by action of RO on MSbO_3 .
Tantalum $\text{Ta}_2\text{O}_5, 2\text{H}_2\text{O}$ (? acid)	MTaO_3 , also complex salts of general form $x\text{Ta}_2\text{O}_5, y\text{RO}$	Ta_2O_5	Acid not formed directly from oxide, but oxide obtained by heating the acid. Salts formed by fusing Ta_2O_5 or $\text{H}_2\text{Ta}_2\text{O}_7$ with basic oxides

The division into two families is not marked, the properties of the members of the group vary gradually from N, which is a typical non metal, to Bi, which is distinctly metallic. The properties of these elements and their compounds show that on the whole V, Nb, and Ta form a family, As, Sb, and Bi another family, and N and P a third family. Di and Er have not been much examined, and it is very probable that neither is an elementary substance. For detailed descriptions of the elements and their chief compounds, v. NITROGEN, PHOSPHORUS, &c

M P M M

NITROGLYCERIN v GLYCERIN

NITRO-HARMALINE v HARMALINE

NITRO-HEMIPIC ACID v HEMIPIC ACID

NITRO-HEPTANE (?) $\text{C}_7\text{H}_{15}\text{NO}_2$ (193°–197°) S G 12 937 Occurs among the products of the action of HNO_3 (S G 1 38) on light petroleum oils (95°–100°) (Beilstein a Kurbatoff, B 18, 2029) Oil, sol warm KOH aq

NITRO-HEPTYL-BENZENE $\text{C}_7\text{H}_{15}, \text{C}_7\text{H}_9\text{NO}_2$ (178° at 10 mm) Formed by nitration (Auger, Bk [2] 47, 50) Heavy straw-coloured oil

DI-NITRO-HEPTYLENE $\text{C}_7\text{H}_9(\text{NO}_2)_2$ [182°] From heptene and HNO_3 (S G 1 3) (Morris, C. J. 41, 176). Tables (from alcohol) Sol alcohol, ether, and benzene, volatile with steam

NITRO HEXADECYL-BENZENE

$\text{C}_{16}\text{H}_{33}(\text{C}_{16}\text{H}_{33})\text{NO}_2$ Nitro cetyl benzene [c 36°] Formed by nitration of cetyl benzene (Krafft, B 19, 2984)

NITRO-p-HEXADECYL TOLUENE

$\text{C}_6\text{H}_4(\text{NO}_2)\text{Me C}_{16}\text{H}_{33}$ [40°] Obtained by nitration of hexadecyl toluene (Krafft a Götting, B 21, 3182)

DI NITRO HEXANE $\text{C}_6\text{H}_{14}, \text{CH}(\text{NO}_2)_2$ Obtained from hexyl methyl ketone and HNO_3 (Chancel, C R 94, 399) Heavy oil Yields π hexoic acid on reduction $-\text{KC}_6\text{H}_{11}\text{N}_2\text{O}_4$ yellow plates (from hot water) $-\text{AgA}'$

Tetra nitro hexane $\text{C}_6\text{H}_6(\text{NO}_2)_4$ Crystals, obtained by passing NO_2 into an ethereal solution of diallyl at 0° (Henry, B 2, 279)

NITRO-HEXOIC ACID $\text{C}_6\text{H}_9\text{NO}_4$ [112°] Formed by reducing di nitro hexoic acid in alcoholic solution by sodium amalgam (Kullhem, A 167, 45, Kachler, A 191, 159) Four sided prisms, v sol water and alcohol With tin and HCl aq it yields methyl isopropyl ketone, hydroxylamine, and CO_2 On adding H_2SO_4 to a solution of the K salt mixed with KNO_3 , there is produced a blue colour, which can be taken up by ether $-\text{NaA}'$ 8aq $-\text{BaA}'$, 3aq $-\text{BaC}_6\text{H}_9\text{NO}_4$, $-\text{AgA}'$

Di-nitro-hexoic acid $\text{C}_6\text{H}_{10}\text{N}_2\text{O}_6$ [215°] Formed by boiling camphor with HNO_3 (Kull-

hem, *A* 163, 231, Kachler, *A* 191, 144) Plates (from water) — $\text{NH}_4\text{A}'$ — $\text{NaA}'4\text{aq}$ — CaA' , 3aq slender needles — BaA' , 5aq — BaA' , 3aq — AgA' Formerly supposed to be $\text{C}_6\text{H}_4\text{N}_2\text{O}_6$, v *Di nitro heptonic acid*, under CAMPHOR, vol 1 p 672

NITROHEXYLENE $\text{C}_6\text{H}_4\text{N}_2$ (?) (210°–215°) A product of the action of HNO_3 (S G 134) on the fraction 95°–100° of the petroleum of Baku (Belstein a Kurbatoff, *B* 13, 1920)

m **NITRO HIPPURIC ACID** $\text{C}_8\text{H}_6\text{N}_2\text{O}_5$, 16 $\text{C}_6\text{H}_4(\text{NO}_2)\text{CO NH CH}_2\text{CO}_2\text{H}$ [162°] S 36 at 23° Formed by the action of HNO_3 and H_2SO_4 on hippuric acid, and occurs in dogs' urine after administration of *m* nitro benzoic acid (Berta gmini, *A* 78, 100, Schwanert, *A* 112, 69, Conrad, *J pr* [2] 15, 254) Needles Split up by HClAq into glycoicoll and *m* nitro benzoic acid CaA' , 3aq — BaA' , 2 — CuA' , 5aq — ZnA' , 6aq — PbA' , 5aq — AgA'

p Nitro hippuric acid $\text{C}_8\text{H}_6\text{N}_2\text{O}_5$ [129°] Occurs in urine after a dose of *p* nitro toluene (Jaffé, *B* 7, 1673) Orange prisms (from alcohol) — BaA' , 4aq — AgA' long needles — Urea salt $\text{HA}'\text{CON}_2\text{H}$, [180°] Occurs in dogs' urine after a dose of *p* nitro benzoic aldehyde (Sieber a Smirnoff, *M* 8, 90) Pearly plates

NITROHYDANTOIN $\text{C}_8\text{H}_7\text{N}_3\text{O}_4$, 16 $\text{NH} \begin{array}{c} \text{CO NH} \\ \text{CO CH(NO)} \end{array}$ [170°] Formed from hydantoin and HNO_3 (Franchumont a Klobbie, *R T C* 7, 12)

DI *p* NITRO HYDROBENZOIN

Acetyl derivative

$\text{C}_6\text{H}_4(\text{NO})\text{CH}(\text{OAc})\text{CH}(\text{OAc})\text{C}_6\text{H}_4\text{NO}$, [340°] From the dibromide of di *p* nitro di phenyl ethylene and alcoholic KOAc (Elbs a Bauer, *J pr* [2] 34, 346) Small yellow crystals, m sol alcohol, ether, and glacial HOAc

NITROHYDROCHLORIC ACID v CHLORHYDRIC ACID, *Reactions*, No 17, vol 11 p 3

NITRO-HYDROCINNAMIC ACID v NITRO β PHENYL PROPIONIC ACID

DI NITRO-HYDRO-*p*-COUMARIC ACID v DI NITRO *p* OXY β PHENYL PROPIONIC ACID

NITRO HYDRO ψ -CUMOQUINONE $\text{C}_6\text{H}_4\text{NO}$, 16 $\text{C}_6\text{Me}_3(\text{NO})(\text{OH})_2$ [106°] Formed from nitro ψ cumoquinone and SO_2 (Nef, *A* 237, 18) Yellow needles (from ether)

NITRO-HYDRO-(β)-NAPHTHOQUINONE

$\text{C}_{10}\text{H}_6\text{N}_2\text{O}_2$, 16 $\text{C}_{10}\text{H}_6(\text{NO})(\text{OH})_2$ [159.5°] Formed from nitro (β) naphthoquinone and SO_2 (Ziertling, *B* 23, 177, v Groves, *C J* 45, 299) Red needles, sol boiling water

NITRO HYDROQUINONE *Mono methyl ether* $\text{C}_6\text{H}_4(\text{NO}_2)(\text{OMe})(\text{OH})$ [83°] Formed from HNO_3 and $\text{C}_6\text{H}_4(\text{OMe})(\text{OH})$ in ether (Weselsky a Benedikt, *M* 2, 369) Orange needles

Di methyl ether $\text{C}_6\text{H}_4(\text{NO}_2)(\text{OMe})_2$ [71°] Formed from $\text{C}_6\text{H}_4(\text{OMe})_2$ and cold dilute (1 10) HNO_3 (Habermann, *B* 11, 1034, Muhlhauser, *A* 207, 253) Felted needles

Mono-ethyl ether $\text{C}_6\text{H}_4(\text{NO}_2)(\text{OEt})(\text{OH})$ [83°] Yellow needles (W a B)

Di-ethyl ether $\text{C}_6\text{H}_4(\text{NO}_2)(\text{OEt})_2$ [49°] Formed by nitration (Nietzki, *A* 215, 148)

Mono-benzyl derivative

$\text{C}_6\text{H}_4(\text{NO}_2)(\text{OC}_6\text{H}_5)(\text{OH})$ [158°] Formed by boiling the benzyl derivative of nitro arbutin with dilute H_2SO_4 (Schiiff a Pellizzari, *A* 221, 871, G 14, 501) Yellow needles (from water).

Vor. III

Di-benzyl derivative

$\text{C}_6\text{H}_4(\text{NO}_2)(\text{OC}_6\text{H}_5)_2$ [83°] Obtained by nitration of $\text{C}_6\text{H}_4(\text{OC}_6\text{H}_5)_2$ Yellow needles (from alcohol)

Di propionyl derivative

$\text{C}_6\text{H}_4(\text{NO}_2)(\text{O COEt})_2$ [86°] Obtained by nitration (Hesse, *A* 200, 247)

Di nitro hydroquinone $\text{C}_6\text{H}_4\text{N}_2\text{O}_6$, 16

$\text{C}_6\text{H}_4(\text{NO}_2)_2(\text{OH})$ [5 2 4 1] [136°] Obtained by saponification of its di acetyl derivative (Nietzki, *B* 11, 469, *A* 215, 145) and also by boiling di-nitro arbutin with dilute H_2SO_4 (Streeker, *A* 118, 293) Flat golden needles (from water) Its alkaline solutions are violet — BaA' violet black needles with bronze lustre

Di acetyl derivative [96°] Obtained by nitrating the di acetyl derivative of hydroquinone Yellow needles

Mono methyl ether [102°] Needles

Di methyl ether The two crystalline isomerides $\text{C}_6\text{H}_4(\text{NO}_2)_2(\text{OMe})_2$ [3 2 1 4] [177°] and $\text{C}_6\text{H}_4(\text{NO}_2)_2(\text{OMe})_2$ [5 2 1 4] [202°] are formed by nitrating $\text{C}_6\text{H}_4(\text{OMe})_2$ (Nietzki a Reehberg, *B* 23, 1216)

Mono ethyl ether [71°] Yellow needles

Di ethyl ether The two crystalline isomerides $\text{C}_6\text{H}_4(\text{NO}_2)_2(\text{OEt})_2$ [3 2 1 4] [130°] and $\text{C}_6\text{H}_4(\text{NO}_2)_2(\text{OEt})_2$ [5 2 1 4] [176°] are formed by nitration of the di ethyl derivative of hydroquinone (Nietzki, *A* 215, 150, N a R)

Methylethyl ether

$\text{C}_6\text{H}_4(\text{NO}_2)_2(\text{OMe})(\text{OEt})$ [144°] Formed from $\text{C}_6\text{H}_4(\text{NO}_2)_2(\text{OMe})(\text{OEt})$ (Fiala, *M* 6, 914)

Benzyl ether $\text{C}_6\text{H}_4(\text{NO}_2)(\text{OC}_6\text{H}_5)(\text{OH})$

[137°] Formed by nitration (S a P) Golden needles — KA' aq explosive scarlet needles — $\text{C}_6\text{H}_4\text{N}_2\text{O}_6 \cdot 2\text{NH}_3$ Loses NH_3 in air, forming $\text{NH}_4\text{A}'$, which is stable

Tri nitro hydroquinone *Di methyl ether* $\text{C}_6\text{H}_4(\text{NO}_2)_3(\text{OMe})$ [101°] Formed by adding a solution of $\text{C}_6\text{H}_4(\text{OMe})_2$ in HOAc to a cold mixture of H_2SO_4 and fuming HNO_3 (H) Yellow needles (from alcohol)

Di ethyl ether $\text{C}_6\text{H}_4(\text{NO}_2)_3(\text{OEt})_2$ [130°] Formed from either $\text{C}_6\text{H}_4(\text{NO}_2)_2(\text{OEt})_2$ by further nitration (Nietzki) Yellow needles Aniline forms red crystals of $\text{C}_6\text{H}_4(\text{NO}_2)_3(\text{NHPh})(\text{OEt})_2$ [133°] whence alcoholic potash yields yellow crystals of $\text{C}_6\text{H}_4(\text{NO}_2)_3(\text{OH})(\text{OEt})_2$ [152°]

Tetra nitro hydroquinone *Di isobutyl ether* $\text{C}_6(\text{NO}_2)_4(\text{OC}_4\text{H}_9)_2$ Formed by nitrating $\text{C}_6\text{H}_4(\text{OC}_4\text{H}_9)_2$ (Schubert, *M* 3, 686) Needles (from alcohol), v sol hot HOAc

DI-NITRO HYDROTOLUQUINONE

$\text{C}_6\text{HMe}(\text{NO}_2)_2(\text{OH})_2$ [149°–153°] Obtained by saponifying its mono acetyl derivative, which is got by nitrating the di acetyl derivative of hydro toluquinone (Kehrmann a Brasch, *J pr* [2] 39, 377) Yellowish red crystals (containing aq)

Acetyl derivative

$\text{C}_6\text{HMe}(\text{NO}_2)_2(\text{OAc})(\text{OH})$ [146°] Yellow crystals

Di acetyl derivative

$\text{C}_6\text{HMe}(\text{NO}_2)_2(\text{OAc})_2$ [154°–157°] Colourless silky needles

NITRO-DI IMIDO-HYDROQUINONE

$\text{C}_6\text{H}(\text{NO})_2(\text{NH})_2(\text{OH})_2$ [3 5 2 4 1] Formed from di amido hydroquinone sulphate, HOAc, and HNO_3 (S G 14) (Nietzki a Schmidt, *B* 22, 1658) Needles or prisms. Reduced by SnCl_2 to tri amido hydroquinone

NITRO IMIDO-DI PHENYL-SULPHOXIDE

$\text{NH} \begin{smallmatrix} \text{C}_6\text{H}_5 \\ \text{C}_6\text{H}_4(\text{NO}_2) \end{smallmatrix} \text{SO}$ Formed by nitration of imido di phenyl sulphoxide (Bernthsen, *B* 17, 2858)

(a)-Di nitro imido-di phenyl sulphoxide

$\text{NH} \begin{smallmatrix} \text{C}_6\text{H}_5 \\ \text{C}_6\text{H}_3(\text{NO}_2)_2 \end{smallmatrix} \text{SO}$ Formed, together with the (β)-isomeride by nitration of imido di phenyl sulphide (Bernthsen, *A* 230, 115) Orange needles Reduced by SnCl_2 and HCl to (α) di-amido imido di phenyl sulphide

Acetyl derivative $\text{C}_{12}\text{H}_9\text{Ac}_2\text{N}_2\text{SO}_2$

(β) Di nitro imido-di phenyl sulphoxide

Lemon yellow powder Reduced by SnCl_2 and HCl to (β) di amido imido di phenyl sulphide

NITRO-DI IMIDO-RESORCIN

$\text{C}_6\text{H}(\text{NH})_2(\text{NO}_2)(\text{OH})_2$ [6 4 2 3 1] Formed from di amido resorcin sulphate, HOAc , and HNO_3 (Nietzki & Schmidt, *B* 22, 1659) Needles — $\text{K}_2\text{A}''$ orange needles

NITRO-INDAZINE $\text{C}_8\text{H}_5(\text{NO}_2)_2\text{N}_2$ [181°]

Formed, together with nitro cresol, by heating diazotised (4, 1, 2) nitro *o* toluidine sulphate with water (Witt, Nollung, & Grandmougin, *B* 23, 3636) Needles Yields a methyl derivative $\text{C}_8\text{H}_5\text{Me}(\text{NO}_2)_2\text{N}_2$ [159°], an acetyl derivative [140°], and a bromo derivative $\text{C}_8\text{H}_4\text{Br}(\text{NO}_2)_2\text{N}_2$ [229°]

DI NITRO-INDIN *v* INDINDI NITRO-INDIGO *v* INDIGONITRO INOSITE *v* INOSITENITRO-IODO- compounds *v* IODO-NITRO-compoundsNITRO-ISATIN *v* ISATINNITRO-ISATOIC ACID *v* ISATOIC ACIDNITRO-LACTIC ACID *v* NITRO OXY PROPIONIC

ACID

TRI-NITRO-LAURENE $\text{C}_{11}\text{H}_5(\text{NO}_2)_3$ (?) [84°] From laurene, HNO_3 , and H_2SO_4 (Fittig, *A* 145, 150)

NITROLIC ACIDS Compounds of the form $\text{RCH}(\text{NO})(\text{NO}_2)$ or $\text{RC}(\text{NO}_2)\text{NOH}$ They are formed by the action of nitrous acid (*ie* KNO_2 and H_2SO_4) on the sodium derivatives of primary nitro paraffins, and by the action of hydroxyl amine on the compounds $\text{RC}(\text{NO}_2)\text{Br}$. Their alkaline solutions are red (V Meyer, *B* 7, 1510, cf vol 1 p 101) The compounds of the form $\text{RR}'\text{C}(\text{NO})(\text{NO}_2)$ formed by the action of nitrous acid on secondary nitro paraffins are called pseudonitrols The pseudonitrols are also formed by the action of NO_2 upon ketones, thus acetone yields $(\text{CH}_3)_2\text{C}(\text{NO})(\text{NO}_2)$ (Scholl, *B* 21, 506) The pseudonitrols may perhaps be formulated $\text{RR}'\text{C}(\text{N}=\text{O})\text{NO}_2$ (V Meyer, *B* 21, 1291) The pseudonitrols do not form salts, their solutions are blue

NITRO-MALONIC ETHER $\text{CH}(\text{NO}_2)(\text{CO}_2\text{Et})_2$ Formed from malonic ether (1 pt) and HNO_3 (5 pts of *SG* 15), the product being extracted with ether (Franchimont & Klobbie, *R T C* 8, 283) Heavy oil, decomposing carbonates and forming a white crystalline compound with NH_3 , decomposing at 150°

Methyl nitro-malonnate forms the analogous $\text{CH}(\text{NO}_2)(\text{CO}_2\text{Me})_2\text{NH}_3$ [c 166°]

NITRO-MANDELIC ACID $\text{C}_8\text{H}_7\text{NO}_4$ *ie* $\text{C}_6\text{H}_5(\text{NO}_2)\text{CH}(\text{OH})\text{CO}_2\text{H}$ [140°] Formed from di *o*-bromo-*o*-nitro acetophenone and very dilute KOH (Engler & Wöhrl, *B* 20, 2201) Formed

also from *o* nitro benzoic aldehyde, KC_6H_5 , MeOH , and HCl (Engler & Ziehl, *B* 22, 207) Small crystals, *v* sol water

Methyl ether MeA' [74.5°] Sol alcohol

m Nitro mandelic acid [120°] Formed from *m* nitro *o* amido phenyl acetic acid and HNO_3 (Plochl & Loe, *B* 18, 1181) and also from di *o* bromo *m* nitro acetophenone and dilute KOH (E & W) Yellowish rhombohedra with bitter taste — $\text{NH}_4\text{A}'$ — AgA'

Ethyl ether EtA' [63°].

m Nitro mandelic imido ether

$\text{C}_8\text{H}_7(\text{NO}_2)\text{CH}(\text{OH})\text{C}(\text{NH})\text{OEt}$ [84°] Formed from *m* nitro-benzoic aldehyde, KC_6H_5 , alcohol, and HCl (Beyer, *J pr* [2] 31, 393) Dendritic needles (from ligron) — BHCl [129°] Needles Cold dilute HClAq converts it quickly into *m* nitro mandelic ether [68°]

p Nitro mandelic acid [126°] Formed in like manner (E & Z) Yields *p* oxy phenyl glyoxylic acid on boiling with aqueous Na_2CO_3

Methyl ether MeA' [87°] Prisms

Ethyl ether EtA' [76°] Needles.

NITRO-MESIDINE *v* NITRO AMIDO MESITYLENE

NITRO-MESITOL $\text{C}_6\text{HMe}_3(\text{NO}_2)(\text{OH})$ [64°] From nitro amido mesitylene and HNO_3 (Knecht, *B* 15, 1376, *A* 215, 98) Yellow plates (from water), volatile with steam

NITRO-MESITYLENE $\text{C}_6\text{HMe}_3(\text{NO}_2)$ [42°] (255°) Formed, together with much di-nitro mesitylene, from mesitylene and HNO_3 (*SG* 1 38) (Fittig, *A* 141, 132, 147, 2) Formed also from nitro amido mesitylene and HNO_3 (Ladenburg, *A* 179, 170, Klobbie, *R T C* 6, 31) Triclinic prisms (from alcohol) CrO_3 in HOAc oxidises it to $\text{C}_6\text{HMe}_3(\text{NO}_2)_2\text{CO}_2\text{H}$ [6 2 1 4] [200°-225°] (Emerson, *Am* 8, 268)

Di-nitro mesitylene $\text{C}_6\text{HMe}_3(\text{NO}_2)_2$ [86°] Obtained by dropping mesitylene into cooled fuming HNO_3 . Trimetric prisms, *m* sol hot alcohol

Tri-nitro-mesitylene $\text{C}_6\text{Me}_3(\text{NO}_2)_3$ [232°] Formed from mesitylene, HNO_3 , and H_2SO_4 . Colourless needles (from hot alcohol) or triclinic prisms (from acetone) Yields NH_3 , and di amido mesitylene on reduction by tin and HClAq

NITRO MESITYLENE SULPHONIC ACID $\text{C}_6\text{H}_3\text{NSO}_3$ *ie* $\text{C}_6\text{HMe}_3(\text{NO}_2)(\text{SO}_3\text{H})$ [131°] *S* 100 in the cold Formed from mesitylene sulphonic acid and HNO_3 (Roze, *Z* [2] 6, 74, *A* 164, 65) Prisms (containing $1\frac{1}{2}$ aq) — $\text{KA}'\text{aq}$ — BaA'_2 , 4aq — CaA'_2 , 3aq — PbA'_2 , aq

(a) **NITRO-MESITYLENIC ACID** $\text{C}_6\text{H}_3\text{NO}_4$ *ie* $\text{C}_6\text{H}_2\text{Me}_2(\text{NO}_2)(\text{CO}_2\text{H})$ [5 3 2 1] [212°]

Formed by nitrating mesitylene acid (Schmitz, *A* 193, 162) Colourless crystals (from alcohol) — BaA'_2 , 4aq needles, *v* *e* sol water

Ethyl ether EtA' [64°] Tables

(β) Nitro-mesitylenic acid

$\text{C}_6\text{H}_2\text{Me}_2(\text{NO}_2)(\text{CO}_2\text{H})$ [5 3 4 1] [223°] Found, in small quantity, in preparing the preceding acid (Fittig, *A* 141, 149, 147, 48, Schmitz) Formed also by oxidising nitro mesitylene (*q v*) monoclinic crystals (from alcohol) Melts at 179° when crystallised from water — BaA'_2 , 4aq — BaA'_2 , 2aq — BaA'_2 , 6aq — CaA'_2 , 6aq — MgA'_2 , 11aq — AgA'

Ethyl ether EtA' [72°]. Needles.

NITRO-MESITYL-PHTHALIMIDE

$C_6H_5 \cdot \langle \begin{smallmatrix} CO \\ CO \end{smallmatrix} \rangle N C_6H_4Me_3(NO_2)$ [210°] Formed by nitration of mesityl phthalimide (Eisenberg, *B* 15, 1018) Prisms, sol alcohol

Di-nitro-mesityl-phthalimide

$C_6H_5 \cdot \langle \begin{smallmatrix} CO \\ CO \end{smallmatrix} \rangle N C_6H_3Me_3(NO_2)_2$ [242°] Formed from mesityl phthalimide, HNO_3 , and $H_2SO_4(E)$ Needles, sol alcohol

NITRO METHANE CH_3NO_2 Mol w 61 (101°) *S* G $\frac{15}{15}$ 1.1441, $\frac{25}{25}$ 1.1330 *M* M 1858 (Perkin, *C* J 55, 687) *S* V 59 5 (Schniff, Lossen, *A* 254, 73) *H* F p 18,600 *H* F v 17,440 (Thomson, *Th*) Formed, unaccompanied by methyl nitrite, from MeI and $AgNO_3$ (V Meyer, *A* 171, 32) Formed also by heating potassium chloroacetate with potassium nitrite (Kolbe, *J pr* [2] 5, 427, Preibisch, *J pr* [2] 7, 480, 8, 316) Heavy oil With alcoholic potash it gives a pp of $CH_3KNO_3(EtOH)$, the aqueous solution of which is ppd by $HgCl_2$, the pp being explosive Iron and acetic acid reduce it to methylamine Fuming H_2SO_4 forms hydroxylamine and CO $HClAq$ (*S* G 1 14) at 150° forms formic acid and hydroxylamine Benzoic aldehyde yields $C_6H_5 \cdot CH \cdot CH \cdot NO_2$ (characteristic) (Priebs, *A* 225, 319) Nitro methane converts dichlorhydrin $CH_2Cl \cdot CH(OH) \cdot CH_2Cl$ into di-chloroformin $CH_2Cl \cdot CH(CHO) \cdot CH_2Cl$ (Pfungst, *J pr* [2] 32, 237) $ZnEt_2$, followed by water, yields methyl-diethyl hydroxylamine (Bevad, *J R* 20, 125)

Di nitro-methane Potassium salt $CHK(NO_2)_2$ Formed by passing H_2S into a solution of $GBrK(NO_3)_2$ mixed with ammonia (Vilhers, *Bl* [2] 41, 282) Yellow explosive crystals

Tri nitro methane $CH(NO_2)_3$ Nitroform [15°] Formed by boiling with water tri nitro acetonitrile, a product of the action of HNO_3 and H_2SO_4 on fulminuric acid (Schischkoff, *A* 103, 364) White crystals, m sol water Explodes when quickly heated Reduced by tin and HCl to hydroxylamine, NH_3 , and H_2C (V Meyer & Locher, *A* 180, 172)

Tetra nitro-methane $C(NO_2)_4$ Mol w 196 [13°] (126°) Formed from tri nitro methane, conc H_2SO_4 , and fuming HNO_3 (Schischkoff, *A* 119, 248) White crystals, which will not burn

NITRO METHANE TRICARBOXYLIC

ETHER $C(NO_2)(CO_2Et)_2$ Formed from $CH(CO_2Et)_2$ and HNO_3 (*S* G 1 52) (Franchumont & Klobbie, *R T C* 9, 220) Oil

NITRO-METHANE DISULPHONIC ACID $CH(NO_2)(SO_3H)_2$ The salt K_2A' , formed by the action of conc H_2SO_4 on chloropierin crystallises in minute plates, v sl sol cold water (Rathke, *A* 161, 153, 167, 220)

NITRO METHOXY- compounds v Methyl derivatives of Nitro oxy compounds

NITRO-DI-METHYL-AMIDO-BENZENE SULPHONIC ACID $C_6H_4(NO_2)_2SO_3H$ Formed together with $C_6H_4(NO_2)_2NMe_2$ from $C_6H_4(NO_2)_2SO_3H$ and aqueous $NaNO_2$ (Michler & Walder, *B* 14, 2176) Yellow crystals (from water) — BaA' , — CaA' .

NITRO-TETRA-METHYL-DI-AMIDO-BENZOPHENONE $C_6H_4(NO_2)_4$ $NMe_2 \cdot C_6H_4(NO_2)_2CO \cdot C_6H_4(NO_2)_2NMe_2$ [144°] Formed from $CO(C_6H_4NMe_2)_2$ by nitration (Nathansohn

& Muller, *B* 22, 1883) Needles (from warm alcohol), sl sol ether

Di-nitro di-methyl-amido-benzophenone $C_6H_4(NO_2)_2NMe_2 \cdot CO$ [142°] Obtained from $C_6H_4NO_2 \cdot CO \cdot C_6H_4NMe_2$ and fuming HNO_3 (Fischer *A* 206, 88) Nodules (from dilute $HOAc$)

Tetra-nitro di-methyl-di-amido-benzophenone $CO(C_6H_4(NO_2)_2NMe_2)_2$ [225°] Formed by boiling its di nitro-derivative with phenol (Van Romburgh, *R T C* 6, 252, 365) Plates, v sl sol alcohol

Di nitro-derivative

$CO(C_6H_4(NO_2)_2NMe_2)_2$ Formed by the action of conc HNO_3 on $CO(C_6H_4NMe_2)_2$ or $CS(C_6H_4NMe_2)_2$ (*R*, Baither, *B* 20, 3296) Yellow crystals, decomposing at about 210°

NITRO-DI-METHYL-AMIDO-PHENOL Anhydride of the methylo-hydroxide

$C_6H_4N_2O_2 \cdot C_6H_4(NO_2)_2 \cdot \langle NMe_2 \rangle$ Formed from nitro amido phenol, MeI and KOH (Griess, *B* 13, 647) Yellow crystals — $B'HClAq$ — $B'HI$ 2aq — $B'H_2PtCl_6$ 6aq yellow crystals

Di-nitro-di-methyl-amido phenol $C_6H_3N_2O_2 \cdot C_6H_4(NO_2)_2NMe_2(OH)$ [195°] Formed from KCy and alcoholic $C_6H_4(NO_2)_2NMe_2$ at 50° (Lippmann & Fleissner, *M* 6, 808) Triclinic yellow crystals Converted by boiling $KOHAq$ into dimethylamine and di nitro resorcin [145°] — NH_4A' [195°] — KA' — BaA' , 1 1/2 aq — AgA' red crystalline pp

Tri nitro methyl amido phenol Nitro-derivative $C_6H(NO_2)_3(OH)NMe_2(NO_2)$ [188°] Formed by boiling $C_6H(NO_2)_2NMe_2(NO_2)$ with water (Van Romburgh, *R T C* 8, 275) Yellow crystals (from water) Its methyl ether $C_6H(NO_2)_3(OMe)NMe_2(NO_2)$ [99°] is formed by dissolving $C_6H(NO_2)_2NMe_2(NO_2)$ in $MeOH$ The ethyl ether $C_6H(NO_2)_3(OEt)NMe_2(NO_2)$ [98°] is formed in like manner

DI NITRO TETRA METHYL DI p AMIDO DIPHENYL $C_6H_4(NO_2)_2(NMe_2) \cdot C_6H_4(NO_2)_2(NMe_2)$ [188°] Obtained by nitration (Michler, *B* 14, 2164, 17, 118) Red needles

DI NITRO-DI-METHYL-p-AMIDO-DI-PHENYLAMINE [4 2 1]

$C_6H_4(NO_2)_2 \cdot NH \cdot C_6H_4NMe_2$ [168°] Formed from $C_6H_4Cl(NO_2)_2$ and $C_6H_4(NH_2)(NMe_2)$ (Leilmann & Mack, *B* 23, 2739) — Orange plates

p NITRO DI METHYL AMIDO DI PHENYL CARBINOL $C_6H_4(NO_2) \cdot CH(OH) \cdot C_6H_4NMe_2$ [96°] Formed by boiling p-nitro benzoic aldehyde with di methyl aniline and $HClAq$ (Albrecht, *B* 21, 3294) Thin yellow needles

Reactions — 1 Dimethylaniline and $ZnCl_2$ yield $C_6H_4(NO_2) \cdot CH(C_6H_4NMe_2)_2$ [177°] — 2 Boiling alcoholic potash and zinc-dust form an azo compound [199°] — 3 Zinc-dust and HCl reduce it to di methyl-di amido di phenyl carbinol [165°] and di methyl di amido di-phenyl-methane [93°]

Methylo iodide $B'MeI$ [c 175°]

o-Nitro tetra-methyl-di-p-amido tri-phenyl carbinol $C_6H_4(NO_2)_2 \cdot C_6H_4(NO_2)_2 \cdot C_6H_4(NO_2)_2 \cdot CO(OH)(C_6H_4NMe_2)_2$ *o-Nitro malachite green* [163°] Formed by heating di-methyl aniline (3 1/2 pts) with o-nitro-benzoic aldehyde (1 pt) and $ZnCl_2$ (1 pt) on the water bath, and oxidising the resulting leuco base with PbO_2 and dilute H_2SO_4 (O Fischer & Schmidt, *B*, 17, 1890) Small yellow crystals, sol alcohol

***m*-Nitro-tetra-methyl-di-*p* amido-tri-phenyl-carbinol** [31] $C_6H_4(NO_2)C(OH)(C_6H_5NMe_2)_3$ Formed by oxidation of *m* nitro tetra methyl di amido tri phenyl methane (E & O Fischer, *B* 12, 802) — *Picrate* small green needles

***p*-Nitro - tetra - methyl - di - amido - tri - phenyl - carbinol** Formed like the two preceding isomerides, and also by heating dimethylaniline with $BzCl$ and $ZnCl$ (E & O Fischer, *B* 12, 800, 14, 2528) Small golden prisms Dyes a splendid green — *Picrate* minute needles

NITRO - DI - METHYL - AMIDO - PHENYL

HEXYL KETONE $C_6H_4(NO_2)C(=O)C_6H_5$ [65°] Obtained by nitrating $C_6H_4(NMe_2)C(=O)C_6H_5$ (Auger, *Bl* [2] 47, 42) Yellow needles (from alcohol)

o-NITRO - TETRA - METHYL - DI - *p* - AMIDO - TRI-PHENYL-METHANE

$C_6H_4(NO_2)CH(C_6H_5NMe_2)_2$ *o Nitro leuco mala chite green* [160°] Prepared by heating *o* nitro benzoic aldehyde with dimethylaniline and $ZnCl$. (Fischer, *B* 15, 682, 17, 1889) Yellow monoclinic prisms, sl sol alcohol

***m*-Nitro - tetra - methyl - di - amido - tri - phenyl - methane** $C_6H_4(NO_2)CH(C_6H_5NMe_2)_2$ [152°] Obtained in the same way from *m* nitro benzoic aldehyde (E & O Fischer, *B* 12, 802) Yellow crystals, sl sol alcohol

***p* Nitro - tetra - methyl di amido tri phenyl methane** [177°] Obtained from *p* nitro benzoic aldehyde, dimethylaniline, and $ZnCl_2$ at 100° (Fischer, *B* 14, 2526) Golden plates *Methylo-vodide* $B'Me_2I_2$ aq [220°]

Tetra nitro - di - methyl di amido di phenyl methane $CH_2(C_6H_4(NO_2)_2NMeH)_2$ [250°] Formed by boiling its di nitro derivative with phenol (Van Romburgh, *R T C* 7, 233) Orange crystals, sl sol hot alcohol

Di nitro derivative

$CH_2(C_6H_4(NO_2)_2NMeNO_2)_2$ Formed from $CH_2(C_6H_4NMe_2)_2$, acetic acid, and HNO_3 Yellow crystalline substance, decomposing at 217°-220°

Hexa - nitro tetra methyl di amido tri - phenyl methane [200°] Formed by nitrating $CH_2(C_6H_4NMe_2)_2$ (Fischer, *A* 206, 122) Golden needles, sl sol alcohol

NITRO - DI - METHYL - *p* AMIDO-PHENYL

OXAMIC ETHER $C_{12}H_8N_2O_3$ *ie* [134] $C_6H_4(NMe_2)(NO_2)NHCOCO_2Et$ [152°] Formed from di methyl-amido phenyl oxamic ether and nitrous acid (Wurster & Sendtner, *B* 12, 1804) Red needles Gives $C_6H_4(NMe_2)(NH_2)_2$ on reduction by tin and $HClAq$

NITRO - TETRA - METHYL - DI - AMIDO - PHENYL-DI-TOLYL-METHANE

$C_6H_4(NO_2)CH(C_6H_4MeNMe_2)_2$ [224°] Formed from di methyl *m* toluidine and *p* nitro benzoic aldehyde (Kook, *B* 20, 1562) Yields a picrate [199°]

NITRO METHYL-AMINE *v* METHYL NITRO AMINE, *p* 279

Nitro-di-methyl-amine *v*. Di methyl nitro-amine, *p* 280

***o*-NITRO-METHYL-ANILINE** $C_6H_4N_2O_2$ *ie* [12] $C_6H_4(NO_2)NMe_2$ [28°] Formed by heating *o* nitro phenol with alcoholic methylamine at 180° (Hempel, *J pr* [2] 41, 164)

Nitrosamine $C_6H_4(NO_2)N(NO)Me$ [36°]

Nitro-methyl-aniline

[13] $C_6H_4(NO_2)NMe_2$ [66°]. Formed by methyl-

ation of *m* nitro aniline (Nöbling & Stricker, *B* 19, 548) Reddish yellow needles, sol Pot Aq

Acetyl derivative $C_6H_4(NO_2)NAcMe$ [95°] Needles, sol water (Meldola, *C J* 53, 777).

Benzoyl derivative $C_6H_4BzN_2O_2$ [105°]

Nitrosamine $C_6H_4(NO_2)N(NO)Me$ [70°]

p Nitro methyl aniline

[14] $C_6H_4(NO_2)NMe_2$ [152°] Formed by heating [14] $C_6H_4(NO_2)N_2NMeC_6H_4(NO_2)$ [41] with $HClAq$ (Meldola & Salmon, *C J* 53, 774) Short thick yellowish brown prisms or tablets (from alcohol), sl sol hot water, v sol alcohol

Acetyl derivative $C_6H_4(NO_2)NAcMe$ [153°]

Benzoyl derivative [112°] Prisms

Nitrosamine $C_6H_4(NO_2)N(NO)Me$ [100°]

m Nitro di methyl-aniline

$C_6H_4(NO_2)NMe_2$ [13] [61°] (280°-285°)

Large red prisms Obtained by heating *m* nitro aniline hydrobromide (1 mol) with $MeOH$ (2 mols) at 100° (S & B) Prepared by nitration of dimethylaniline in presence of a large excess of H_2SO_4 , *p* nitro dimethylaniline being formed in smaller quantity A mixture of 193 g of HNO_3 (S G 1.35) and 600 g of ordinary H_2SO_4 is allowed to slowly drop into a solution of 200 g of dimethylaniline in 4,000 g of ordinary H_2SO_4 , keeping the mixture cooled below 5° After standing for 4 or 5 hours it is poured into about 10 litres of iced water and filtered from the ppd *p* nitro derivative, more of which separates on partial neutralisation with Na_2CO_3 (50 or 60 g on the whole) The filtrate neutralised with Na_2CO_3 deposits the *m* nitro di methyl aniline, the yield is 160 to 170 g (Groll, *B* 19, 198)

Methylo bromide $C_6H_4(NO_2)NMe_2Br$ Crystalline (Stadel & Bauer, *B* 19, 1941) Yields crystalline $B'MeCl$, $B'MePtCl_6$, and unstable $B'MeOH$ which is converted by *m* nitro phenol into $C_6H_4(NO_2)NMe_2$ $O C_6H_4(NO_2)$ [62°]

***p* Nitro di methyl aniline** $C_6H_4(NO_2)NMe_2$ [163°] Formed by adding HNO_3 to a solution of dimethylaniline in $HOAc$ (Webster, *B* 10, 760) Formed also by oxidising *p* nitroso di methyl-aniline with $KMnO_4$ (Wurster, *B* 12, 528, cf Schraube, *B* 8, 616), by heating *p* chloro nitro benzene with NMe_2 at 180° (Leymann, *B* 15, 1234), and by the action of nitrous acid on $C_6H_4(NMe_2)SO_3H$ (Michler & Walder, *B* 14, 2176), on $C_6H_4(NMe_2)N_2C_6H_4NMe_2$ (Lippmann & Fleissner, *B* 16, 1421), and on the base $C_6H_4(NO_2)(NH_2)NMe_2$ (Hewitt, *B* 21, 2309) Yellow needles Does not form salts

Di nitro-methyl aniline $C_6H_4N_2O_4$ *ie* $C_6H_4(NO_2)_2NMe_2$ [177°]

Formation — 1 From chloro di nitro benzene and methylamine (Leymann, *B* 15, 1234) — 2 By heating $C_6H_4(NO_2)_2NMe_2$ (78°) in $HOAc$ with bromine (L), or CrO_3 (Van Romburgh, *R T C* 8, 250) — 3 By nitrating the formyl, acetyl, or oxalyl derivative of methyl aniline (Norton & Allen, *B* 18, 1995, Norton & Livermore, *B* 20, 2273)

Properties — Yellow crystals, v sol alcohol, sl sol hot water

Di nitro di-methyl-aniline

$C_6H_4(NO_2)_2NMe_2$ [421] [87°] Formed from $C_6H_4Cl(NO_2)_2$ and NMe_2 (Leymann, *B* 15, 1283) Prepared by slowly adding 500 g of dimethyl aniline to a mixture of 6 litres of ordinary conc.

HNO₃ and 6 litres of water, cooled to 0°, the yield being 116° (Mertens, *B* 19, 2123, cf Weber, *B* 10, 763). Large yellow pointed crystals or prisms. V sol alcohol, chloroform, benzene, acetic acid, and HClAq. Heated with 10 p.c. KOH aq it readily decomposes into di nitro phenol [114°] and dimethylamine. Boiled with fuming HNO₃ it is converted into tri nitro phenyl methyl nitramide C₆H₃(NO₂)₃NMe(NO₂). Salts — B'HCl unstable — B'₂H₂Cl PtCl₄, granular pp., or large brownish red pyramids.

Di nitro di methyl aniline

C₆H₄(NO)₂NMe [5317]. Formed by nitration of dimethylaniline with dilute HNO₃ (equal vols of HNO₃ (S G 1.4) and water) without cooling, the yield being 15 p.c. (Mertens, *B* 19, 2125). Golden brown plates. Decomposes at 250°–272°. Soluble in phenol. It is attacked only by very strong KOH. Boiled with fuming HNO₃ it is converted into di nitro phenyl methyl nitramine C₆H₄(NO)₂NMe(NO₂), a crystalline body which explodes above 220°.

Tri-nitro methyl-aniline

C₆H₃N₃O₆ 2e C₆H(NO₂)₃NHMe [111°]. Formed from C₆H₅Cl(NO₂)₃ (picryl chloride) and methylamine in hot alcoholic solution (Van Romburgh, *R T C* 2, 31, 105, 305). Yellow needles (from alcohol). Gives off NMeH₂ when heated with KOH aq.

Nitro derivative C₆H(NO₂)₃NMe(NO₂) [127°]. Obtained from dimethylaniline (10 pts), H₂SO₄ (100 pts), and HNO₃ (200 pts of S G 1.48). Obtained also from methyl aniline and HNO₃ (Romburgh), and by nitration of di methyl amido phenyl naphthyl sulphone (Michler a Schacht, *B* 12, 1790). Pale yellow crystals (from alcohol). Yields picric acid on boiling with aqueous Na₂CO₃. Reduced by tin and HClAq to tri amido phenol.

Tri nitro-di-methyl aniline C₆H(NO₂)₃NMe₂ [138°]. Formed from picryl chloride and alcoholic dimethylamine (Van Romburgh, *R T C* 2, 105, 8, 250). Yellow crystals (from benzene). Forms with picric acid the crystalline compound C₆H₂N₃O₆C₆H(NO₂)₃OH [114°].

Tetra-nitro-methyl-aniline Nitro derivative C₆H(NO₂)₄NMe(NO) [146°]. Formed from the preceding tri nitro di methyl aniline and HNO₃ (S G 1.5) (Van Romburgh, *R T C* 8, 274). Yellowish white needles. On dissolving in MeOH it yields C₆H(NO₂)₃(OMe)NMe(NO₂) [99°], alcohol forms C₆H(NO₂)₃(OH)NMe(NO₂) [98°] in like manner. Aqueous methylamine produces C₆H(NO₂)₃(NMeH)NMe(NO₂) [192°] and C₆H(NO₂)₃(NMeH)₂ [235°]. If a few drops of aniline be poured upon the nitro derivative it catches fire.

NITRO-METHYL-ANTHRAQUINONE

C₁₀H₆O₂(CH₃)₂(NO₂) [270°]. Prepared by nitration of methyl anthraquinone by adding KNO₃ to its solution in H₂SO₄ (Romer a Lunk, *B* 16, 696). Sublimes in small white needles. Sol alcohol and ether, v sol nitrobenzene.

TRI-NITRO-TRI-METHYL-BENZENE

C₆Me₃(NO₂)₃ [128456] [209°]. Got by nitration (Jacobsen, *B* 19, 2517).

NITRO-METHYL-BENZOIC ACID v Nitro TOLUIC ACID

Nitro di-methyl benzoic acid C₆H₂NO₂ 2e C₆H₂Me₂(NO₂)CO₂H [195°]. Formed by ox-

dising nitro ψ cumene (Schaper, *Z* 1867, 13). Slender needles — CaA', 6aq — BaA', 9aq — EtA' p-NITRO-METHYL COUMARILIC ACID



C₆H₂(NO₂)₂ < $\begin{smallmatrix} \diagup \\ \diagdown \end{smallmatrix}$ [178°] Obtained by

saponification of its ethyl ether, which is formed by heating dry p-nitro phenol sodium with chloroacetoacetic ether and dissolving the product (probably p nitro phenoxy acetic ether) in conc H₂SO₄ (Nuth, *B* 20, 1332). Short yellow needles. M sol hot water, alcohol, and ether, nearly insol cold water. Most of the salts are sparingly soluble — AgA' 3aq small needles.

Ethyl ether EtA' [74°] White needles

NITRO-METHYL ETHYL-BENZENE v

NITRO ETHYL TOLUENE

NITRO (B 2 Py 2) DIMETHYL (Py 3)

ETHYL QUINOLINE C₁₀H₇N(NO₂) [109°]

Formed by nitration of di methyl ethyl quinoline (Harz, *B* 18, 3391). Triclinic crystals. Easily soluble in chloroform, m sol ether.

Salts — B'HCl 2e easily soluble yellow crystals — B'₂H₂ PtCl₄ 2aq short thick orange crystals, sol hot water.

Nitro (B 2, 4, Py 2) tri methyl-(Py 3) ethyl quinoline C₁₀H₇(NO₂)₃N [90°]. Formed by nitration of the base (cf p 326). Needles (from alcohol). It is accompanied by the di-nitro derivative C₁₀H₇(NO₂)₂N [1525°] Waldbott, *B* 23, 2272).

DI NITRO METHYLIC ACID (so called)

CH₂N₂O₂ 2e CH₂NHNO₂? By passing NO into ZnMe there is formed CH₃(ZnMe)N₂O₂, which is decomposed by water yielding CH₃(ZnOH)N₂O₂, whence CO₂ yields crystalline (CH₃N₂O₂)₂Znaq, which is converted, by adding Na₂CO₃ to its aqueous solution, into very soluble CH₃NaN₂O₂ aq (Frankland, *C J* 11, 88).

DI NITRO METHYL MESIDINE Nitramine

C₆Me₂(NO₂)₂NMe(NO₂) [138°]. Formed from C₆Me₂H₂NMe, H₂SO₄, and HNO₃ (Klobbie, *R T C* 6, 31). Crystals, sl sol ether.

NITRO METHYL NAPHTHALENE

C₁₀H₇NO₂ [81°]. Formed, together with the di nitro derivative [206°], by nitration of (β) methyl naphthalene (Schulze, *B* 17, 844). Thin yellowish needles.

DI NITRO (Py 3) METHYL-(β) NAPHTHO

QUINOLINE C₁₀H₇(NO₂)₂N. Four bodies of this constitution, melting at 227°, 230°, 205°–212°, and 250°, may be obtained, as well as a tetra nitro derivative [277°], by nitrating the (Py 3) methyl (β) naphthoquinoline which is formed by condensation of (β) naphthylamine with paraldehyde (Seitz, *B* 22, 256).

NITRO DI METHYL PHENYL ACETIC

ACID [3521] C₆H₂(CH₃)₂(NO₂)CH₂CO₂H **Nitro mesitylic acetic acid** [139°]. Formed by heating (531) di methyl phenyl acetic acid with dilute nitric acid (Wisepek, *B* 16, 1679). Long fine needles. Sol alcohol, ether, and hot water, insol cold water. On reduction it gives carbo mesyl (C₆H₂(CH₃)₂< $\begin{smallmatrix} \text{CH} \\ \text{NH} \end{smallmatrix}$ >CO).

Salts — A'₂Ca 4aq thick needles — A'₂Ba 4aq fine needles — A'Ag

Di-nitro-methyl phenyl acetic acid v Di-NITRO TOLYL ACETIC ACID,

(B 4) **NITRO-(Py 3) METHYL QUINOLINE**
 $C_9H_8N_2O_2$, $\begin{smallmatrix} CH\ CH\ CH \\ CH\ CNO_2\ C\ N\ CMe \end{smallmatrix}$ *o*-Nitro
quinaldine [187°] Formed, together with the
 isomeride [82°] by nitration of (Py 3) methyl
 quinoline (Doebner & Miller, *B* 17, 1699, Ger-
 deissen, *B* 22, 245) Formed also by the action
 of aldehyde on *m* nitro aniline Long needles,
 sl sol cold water—B'HCl prisms—B₂H₂PtCl₆,
 small needles

(B 1 or 3) Nitro (Py 3)-methyl-quinoline
 $C_9H_8Me(NO_2)N$ *m* Nitro *quinaldine* Formed
 as above, and also by the action of aldehyde on
m nitro aniline (D & M) Slender needles,
 volatile with steam—B'HCl prisms—
 B₂H₂PtCl₆ needles

(B 1 or 3) Nitro-(B 2) methyl-quinoline
 [117°] Formed by nitrating (B 2) methyl
 quinoline (Noelting & Trautmann, *B* 23, 3655)
 Formed also from (2, 1, 4) nitro toluidine, gly-
 cerin, picric acid, and H₂SO₄ Needles Yields
 a methylo iodide [190°]

(B 4) Nitro-(B 2) methyl-quinoline
 $CMe\ CH—C\ CH\ CH$
 $CH\ C(NO_2)\ CN\ CH$ [122°] Formed from
 nitro *p* toluidine (N & T) Yellow needles

(B 1) Nitro-(B 4) methyl-quinoline
 $CH\ C(NO_2)\ C\ CH\ CH$
 $CH\ CMe\ CN\ CH$ [93°] Formed from
 (4, 2, 1)-nitro-*o* toluidine and also by nitrating
 (B 4) methyl quinoline

Nitro-methyl-quinoline [127°] Got by
 nitrating lepidine (Busch & Königs, *B* 23, 2687)

Nitro-(Py 3) methyl quinoline tetrahydride
Nitroso derivative $C_{10}H_{11}(NO_2)N\ NO$ [152°]
 A product of the action of nitrous acid on (Py 3)
 methyl quinoline tetrahydride (Möller, *A* 242,
 814) Golden plates (from alcohol benzene)

Nitro (Py 4) methyl-quinoline tetrahydride
 $C_9H_8(NO_2) \begin{smallmatrix} CH_2\ CH_2 \\ NMe\ CH_2 \end{smallmatrix}$ *Nitro karoiline* [94°]

Formed by careful nitration of karoiline (Feer &
 Königs, *B* 18, 2390) Long red needles

(B 1) Nitro (B 2, 4) di methyl quinoline
 $CMe\ C(NO_2)\ C\ CH\ CH$
 $CH\ CMe\ CN\ CH$ [108°] Formed by
 nitrating di methyl quinoline, and also from
 nitro *m*-xyldine [125°], glycerin, H₂SO₄, and
 picric acid (N & T) Yellow needles

Nitro-(B 2, 4, Py 3)-tri methyl quinoline
 $CMe\ CH\ C\ CH\ CH$
 $CH\ CMe\ CN\ CMe$ [92°] Formed by nita-
 ting tri methyl quinoline (Panajotoff, *B* 20, 35)
 Needles—B₂H₂PtCl₆, jag crystalline

Di-nitro-(Py 4)-methyl-quinoline tetra-
 hydride $C_9H_8(NO_2)_2 \begin{smallmatrix} CH_2\ CH_2 \\ NMe\ CH_2 \end{smallmatrix}$ *Di nitro*
karoiline [148°] Formed from karoiline, HOAc,
 and HNO₃ (Feer & Königs, *B* 18, 2390) Golden
 yellow plates, sol alcohol

DI-NITRO-DI-METHYL-SULPHAMIDE
 $C_9H_8N_2SO_2$, $\begin{smallmatrix} SO_2\ (NMe\ NO_2)_2 \end{smallmatrix}$ [90°] Formed
 from di methyl sulphamide and HNO₃ (Franchi
 mont, *R T C* 8, 419) Crystals (from benzene)

NITRO-METHYL-THIENYL METHYL
KETONE $C_9H_8NSO_2$, $\begin{smallmatrix} C_6Me_4H\ CO\ CH_3 \end{smallmatrix}$
 [121°] Obtained by nitrating $C_6Me_4H\ S\ CO\ CH_3$
 (Demuth, *B* 19, 1861) Needles (from ether)

NITRO DI-METHYL-THIOPHENE
 $C_9H_8NSO_2$, $\begin{smallmatrix} SC_6Me_4H\ (NO_2)_2 \end{smallmatrix}$ *Nitro thioxene*

Oil, formed from di methyl thiophene vapour,
 HOAc, and HNO₃ (Messinger, *B* 18, 1638)

NITRO METHYL-*o*-TOLUIDINE $C_8H_8N_2O_2$,
 $\begin{smallmatrix} CH_3\ Me\ (NO_2)\ NHMe \end{smallmatrix}$ [2 4 1] [134°] Obtained
 by oxidation from nitroso methyl *o* toluidine
 $C_8H_8Me(NO)\ NHMe$ (Kock, *A* 243, 309) Green
 ish yellow needles, sol benzene

Nitro-methyl *p* toluidine
 $C_8H_8Me(NO_2)\ NHMe$ [4 2 1] [85°] Formed from
 nitro-*p* toluidine and MeI (Gattermann, *B* 18,
 1487, Niementowski, *B* 20, 1874) Red needles
 (from alcohol) or tables (from benzene)

Acetyl derivative [64°] (250-255° at
 270 mm) Plates—B'C₆H₄(NO₂)₂OH [212°]

Di-nitro-methyl-*o*-toluidine *Nitro deriva-*
tive [2 4 6 1] $C_8H_8Me(NO_2)_2\ NMe\ NO$, [120°]
 Formed from di methyl *o* toluidine and HNO₃
 (Van Romburgh, *R T C* 3, 395) Pale yellow
 crystals Potash converts it into methylamine
 and di nitro *o* cresol

Di-nitro-methyl-*p* toluidine
 $C_8H_8Me(NO_2)_2\ NHMe$ [4 6 2 1] [129°] Obtained
 from $C_8H_8Me(NaCMe)$ and HNO₃ (A L Thomsen,
B 10, 1582, Gattermann, *B* 18, 1487) Red
 needles (from dilute alcohol)

Nitrosamine $C_8H_8Me(NO_2)_2\ NMe\ NO$
 [125°] Yellow needles

Nitro derivative $C_8H_8Me(NO_2)_2\ NMe\ NO$,
 [139°] Formed from di methyl-*p* toluidine (1 pt),
 H₂SO₄ (2 pts), and HNO₃ (10 pts) (Van Rom-
 burgh, *R T C* 3, 404) The same compound
 [130°] appears to be formed by the action of
 dilute HNO₃ on [1 4] $C_8H_8Me\ NaCMe$ (Norton &
 Livermore, *B* 20, 2268) Nearly colourless
 crystals (from boiling alcohol) Yields di nitro
p cresol on boiling with potash

Tri nitro-methyl *m* toluidine *Nitro-de-*
rivative [3 6 4 2 1] $C_8H_8Me(NO_2)_3\ NMe\ NO$
 [102°] Formed from di methyl *m* toluidine and
 HNO₃ (S G 1.5) (Van Romburgh, *R T C* 3,
 413) Pale yellow crystals

Tri nitro methyl *p* toluidine
 $C_8H_8Me(NO_2)_3\ NHMe$ [4 6 3 2 1] [138°] Formed
 by further nitration of nitro methyl *p* toluidine
 (G) Yellowish crystals

Nitro di-methyl *m* toluidine
 $C_8H_8Me(NO_2)_2\ NMe_2$ [84°] Obtained by ox-
 idising nitroso di methyl *m* toluidine with KMnO₄
 (Wurster & Riedel, *B* 12, 1800) Yellow needles

Di nitro di-methyl-*m* toluidine
 $C_8H_8Me(NO_2)_2\ NMe_2$ [107°] Prepared by nita-
 ration of di methyl *m* toluidine dissolved in
 HOAc (W & R) Yellow needles

Di nitro di-methyl *m* toluidine
 $C_8H_8Me(NO_2)_2\ NMe_2$ [168°] Formed from di
 methyl-*m* toluidine, HNO₃, and H₂SO₄ (W & R)
 Less sol alcohol than the isomeride [107°]

(a) **NITRO NAPHTHALENE** $C_{10}H_8NO_2$ Mol
 w 173 [56°] (Mills, *P M* [5] 14, 27), [61°]
 (D'Aguiar, *B* 5, 370, Schiff) (304°) (De Coninck,
B 5, 12) S V S 141 (Schiff, *A* 223, 265)
 S G 1 1331 (Schröder, *B* 12, 1613) S (88 p c
 alcohol) 28 Formed by nitration of naph-
 thalene (Laurent, *A Ch* [2] 59, 378, Beilstein
 & Kuhlberg, *A* 169, 88, Guareschi, *B* 10, 294)
 Formed also by elimination of NH₃ from (a) nitro
 (a)-naphthylamine (Liebermann, *A* 183, 235)

Preparation—By shaking naphthalene with
 nitric acid in the cold, and recrystallising from
 alcohol or ligroin (B & K, cf Piria, *A* 78, 32)
 Prepared also by stirring naphthalene (250 pts.)

with a mixture of HNO_3 (200 pts) and H_2SO_4 (800 pts) at $40^\circ\text{--}50^\circ$ (Witt, *Chem Ind* 10, 215).

Properties—Pale yellow prisms. A few milligrammes projected into a red hot test tube containing nitrogen detonates slightly with a white flame, a larger quantity gives a red flame, with a still larger quantity no flame is seen (Berthelot, *A Ch* [6] 16, 24).

Reactions—1 Readily reduced to (a) naphthylamine. Zinc dust and HClAq yields chlorinated naphthylamines—2 PCl_5 forms (a) chloro naphthalene—3 *Chlorine* yields mono-, di, tri-, and tetra chloro naphthalenes (Atterberg, *B* 9, 316, 926)—4 *Bromine* forms bromo nitro naphthalene [122 5°], di bromo-nitro naphthalene [98°], and two tetrabromides—5 HBrAq at 195° yields $\text{C}_{10}\text{H}_7\text{Br}$ and $\text{C}_{10}\text{H}_6\text{Br}_2$ (Baumhauer, *B* 4, 926)—6 CrO_3 in HOAc forms *c*-nitro phthalic acid (Beilstein a Kurbatoff, *C* 1881, 359)—7 Distillation over *vacuo* distill produces ($\alpha\beta$) naphthazine (G Schultz, *B* 17, 478).

Tetrabromide $\text{C}_{10}\text{H}_6(\text{NO}_2)\text{Br}_4$ [181°] *S* (93.5 p.c. alcohol) 26 at 15.2° . Formed from nitro naphthalene and bromine (Guareschi, *A* 222, 286). White needles. When kept at 137° in a current of air for some time it changes to an isomeride [145°] and $\text{C}_{10}\text{H}_6\text{Br}(\text{NO}_2)$ [122 5°]. When boiled with alcohol for a long time it changes to a second isomeride [173°], *S* (93.5 p.c. alcohol) 13 at 15.2° .

(β) Nitro naphthalene $\text{C}_{10}\text{H}_7\text{NO}_2$ [79°] Formed by eliminating NH_3 from (2, 1) nitro (a) naphthylamine (Lellmann, *B* 19, 236, 20, 891). Formed also by the action of precipitated Cu_2O (2 mols) upon a solution of (β) diazo naphthalene nitrite (1 mol) obtained by adding a solution of 12 g NaNO_2 in 40 g water to a mixture of 7 g of (β) naphthylamine, 15 g HNO_3 , and 250 cc of water. The yield is small (Sandmeyer, *B* 20, 1496). Yellow needles, *v* sol alcohol. Volatile with steam. Reduced by zinc and HOAc to (β) naphthylamine.

' α ' Di-nitro-naphthalene $\text{C}_{10}\text{H}_6(\text{NO}_2)_2$ [1 4°] Mol w 218 [212°] Formed, together with the (1,1') isomeride, by boiling naphthalene with fuming HNO_3 or by heating it with HNO_3 and H_2SO_4 at 100° (Darmstadter a Wichelhaus, *B* 5, 253, D'Aguiar, *B* 5, 370, Beilstein a Kurbatoff, *A* 202, 219). Formed also by heating nitro (a) naphthoic acid [239°] with HNO_3 (*S* G 13) (Ekstrand, *B* 18, 2881).

Properties—Six sided needles (from HOAc), *v* sol HOAc . A few milligrammes thrown into a red hot tube full of nitrogen detonate with reddish white flame (Berthelot, *A Ch* [6] 16, 25).

Reactions—1 *Oxidised* by HNO_3 (*S* G 115) at 150° to *c* nitrophthalic, *s* di nitro benzoic, and picric acids (Beilstein a Kurbatoff, *Bl* [2] 34, 327)—2 PCl_5 yields (γ)-di-chloro naphthalene [107°].

' β ' Di-nitro-naphthalene $\text{C}_{10}\text{H}_6(\text{NO}_2)_2$ [1 1°] [172°] *S* (88 p.c. alcohol) 187 at 19° *S* (benzene) 72 at 19° *S* (chloroform) 11 at 19° . Formed, at the same time as the preceding, by nitrating naphthalene (Darmstadter a Wichelhaus, *A* 152, 301, Beilstein a Kuhlberg, *A* 169, 86). Formed also by heating nitro (a) naphthoic acid [216°] with HNO_3 (*S* G 13) (Ekstrand, *B* 18, 2881), and by eliminating NH_3 from di-nitro (β)-naphthylamine (Gaess, *J. pr*

[2] 43, 37). Tables, more soluble in HOAc , chloroform, benzene, and acetone than the (1,4')-isomeride.

Reactions—1 Dilute HNO_3 at 150° yields di nitro phthalic acid [226°], *s* di nitro benzoic acid, and picric acid—2 PCl_5 forms (γ) di-chloro-naphthalene and (δ) tri-chloro naphthalene.

(γ) Di-nitro-naphthalene $\text{C}_{10}\text{H}_6(\text{NO}_2)_2$ [1 3°] [144°] Obtained from di nitro (a)-naphthylamine [235°] by elimination of NH_3 (Liebermann a Hammerschlag, *A* 183, 272). Yellow needles.

(δ) Di-nitro-naphthalene $\text{C}_{10}\text{H}_6(\text{NO}_2)_2$ [1 3°] [162°] Formed by eliminating NH_3 from di-nitro-(β) naphthylamine [238°] (Graebe a Drews, *B* 17, 1172). Yellow needles, *v* sol alcohol.

' α '-Tri-nitro-naphthalene $\text{C}_{10}\text{H}_5(\text{NO}_2)_3$ Mol w 263 [122°] Formed by further nitration of ' α '-di nitro naphthalene (D'Aguiar, *B* 5, 372, 897). Monoclinic crystals, *v* sol HOAc .

' β '-Tri-nitro-naphthalene $\text{C}_{10}\text{H}_5(\text{NO}_2)_3$ [213°] *S* (88 p.c. alcohol) 06 at 23° . Obtained by further nitration of ' β ' di nitro naphthalene (Laurent, *A* 41, 98, D'Aguiar, Beilstein a Kuhlberg, *A* 169, 96). Formed also by eliminating NH_3 from tri nitro (a) or (β)-naphthylamine (Staedel, *B* 14, 901, *A* 217, 174), and by the action of fuming HNO_3 and conc H_2SO_4 on nitro (a) naphthoic acid (Ekstrand, *B* 19 1181). Prisms (from HOAc), *v* sol HOAc and ether.

(γ) Tri-nitro-naphthalene $\text{C}_{10}\text{H}_5(\text{NO}_2)_3$ [147°] (*B* a *K*), [154°] (*A*). *S* (90 p.c. alcohol) 11 at 18.5° *S* (ether) 38 *S* (benzene) 105 at 18.5° . Obtained by boiling ' α ' di nitro naphthalene for a few minutes with H_2SO_4 and fuming HNO_3 (Beilstein a Kuhlberg, *B* 6, 647). Yellow plates.

' α '-Tetra-nitro-naphthalene $\text{C}_{10}\text{H}_4(\text{NO}_2)_4$ [259°] Formed by boiling ' α ' di nitro naphthalene with HNO_3 and H_2SO_4 for some hours (*A*, *B* a *K*). Crystals (from chloroform), almost insol alcohol.

' β ' Tetra-nitro-naphthalene $\text{C}_{10}\text{H}_4(\text{NO}_2)_4$ [200°] Formed by heating ' β ' di nitro naphthalene with fuming HNO_3 for 4 days at 100° (Lautemann a D'Aguiar, *Bl* 3, 261). Asbestos-like needles (from alcohol). Explodes when heated.

References—Bromo-, Chloro-, and Iodo-

NITRO NAPHTHALENE

NITRO-NAPHTHALENE DICARBOXYLIC ACID $\text{C}_{10}\text{H}_6(\text{NO}_2)(\text{CO}_2\text{H})_2$. Nitro naphthoic acid. Got by oxidising nitro acenaphthene (Quincke, *B* 21, 1454). Yellow needles. On heating it yields an anhydride [220°]— CaA ' aq— $(\text{NH}_4)_2\text{A}$ ' aq.

' α '-NITRO-NAPHTHALENE SULPHONIC ACID $\text{C}_{10}\text{H}_5(\text{NO}_2)(\text{SO}_3\text{H})$ [1 4°] Formed from (a) nitro naphthalene and fuming H_2SO_4 or ClSO_3H (Laurent, *C* 81, 537, Armstrong a. Williamson, *C J Proc* 2, 233, Erdmann, *A* 247, 311). Formed also by nitrating naphthalene (a)-sulphonic acid (Cleve, *Bl* [2] 24, 506). Straw yellow prisms (containing 4 aq), *v* sol. water, sl sol dilute H_2SO_4 . Tastes bitter. Its *K* salt distilled with $\text{K}_2\text{Cr}_2\text{O}_7$ and HCl gives chloro nitro-naphthalene [111°] and di-chloro-nitro naphthalene [85°]. Sodium amalgam reduces it to (a) naphthylamine and H_2SO_4 (Claus a Graeff, *B* 10, 1308).

Salts— KA ' 3 aq *S* 21 at 15° — NH_4A ' 1 3 aq— NaA ' 3 aq— MgA ' 8 aq— CaA ' 2 aq— BaA ' 8 aq.

— $\text{ZnA}'_2, 6\text{aq}$ — $\text{PbA}'_2, 3\text{aq}$ — $\text{MnA}'_2, 2\text{aq}$ —
 $\text{CuA}'_2, 4\text{aq}$ — AgA' monoclinic prisms

Ethyl ether EtA' [101°]

Chloride $\text{C}_{10}\text{H}_7(\text{NO}_2)(\text{SO}_2\text{Cl})$ [113°].

Amide $\text{C}_{10}\text{H}_7(\text{NO}_2)(\text{SO}_2\text{NH}_2)$ [225°]

Nitro-naphthalene- β ' sulphonic acid

$\text{C}_{10}\text{H}_6(\text{NO}_2)(\text{SO}_3\text{H})$ [1 3'] Formed, together with two or more isomerides, by nitration of naphthalene (β)-sulphonic acid (Cleve, *Bl* [2] 26, 444), and, together with a larger quantity of the ' α ' acid and some of the θ acid by heating nitro naphthalene with H_2SO_4 containing excess of SO_2 at 100° (Palmaer, *B* 21, 3260) Yields $\text{C}_{10}\text{H}_6\text{Cl}_2$ [48°]

Salts — KA' — $\text{NH}_4\text{A}'$ — $\text{NaA}'_2, 3\text{aq}$ — $\text{BaA}'_2, 2\text{aq}$ S 115 at 22° — $\text{MgA}'_2, 7\text{aq}$ — $\text{CaA}'_2, 3\text{aq}$ — $\text{ZnA}'_2, 6\text{aq}$ — $\text{PbA}'_2, 3\text{aq}$ — $\text{MnA}'_2, 6\text{aq}$ — $\text{CuA}'_2, 6\text{aq}$ — AgA'

Ethyl ether EtA' [115°]

Chloride $\text{C}_{10}\text{H}_6(\text{NO}_2)(\text{SO}_2\text{Cl})$ [126°] Monoclinic crystals, $a, b, c = 996, 1, 831, \beta = 81^\circ 28'$

Amide [184°] Yellowish needles

(γ)-Nitro-naphthalene sulphonic acid

$\text{C}_{10}\text{H}_6(\text{NO}_2)(\text{SO}_3\text{H})$ [1 3'] Formed, together with the ' β ' and (θ) isomerides, by nitration of naphthalene (β) sulphonic acid (Cleve, *B* 19, 2170) Its chloride on heating with PCl_5 yields $\text{C}_{10}\text{H}_6\text{Cl}_2$ [61°] — KA' — NaA' — $\text{BaA}'_2, 3\text{aq}$ — $\text{PbA}'_2, 3\text{aq}$

Ethyl ether EtA' [115°]

Chloride $\text{C}_{10}\text{H}_6(\text{NO}_2)(\text{SO}_2\text{Cl})$ [140°]

Amide [225°] Long needles

(θ) Nitro-naphthalene sulphonic acid

$\text{C}_{10}\text{H}_6(\text{NO}_2)(\text{SO}_3\text{H})$ [1 2'] The Ba salt is the most soluble of the Ba salts of the acids obtained by nitration of naphthalene (θ) sulphonic acid (Cleve, *Bl* [2] 29, 415, *B* 21, 3264) Occurs also among the products of sulphonation of (α) nitro naphthalene (Palmaer) Its chloride, heated with PCl_5 , yields $\text{C}_{10}\text{H}_6\text{Cl}_2$ [62°] — $\text{KA}'_2, 3\text{aq}$ — $\text{NaA}'_2, 3\text{aq}$ — $\text{NH}_4\text{A}'$ — $\text{BaA}'_2, 3\text{aq}$ S (of BaA'_2) 11 at 100°, 2 at 17° — $\text{CaA}'_2, 2\text{aq}$ — $\text{MgA}'_2, 9\text{aq}$ — $\text{PbA}'_2, 3\text{aq}$ — $\text{MnA}'_2, 10\text{aq}$ — $\text{CuA}'_2, 8\text{aq}$ — $\text{ZnA}'_2, 10\text{aq}$ — AgA'

Ethyl ether EtA' [107°] Needles

Chloride [167°] Yellow needles

Amide [223°] Needles

Nitro naphthalene sulphonic acid

$\text{C}_{10}\text{H}_6(\text{NO}_2)(\text{SO}_3\text{H})$ [1 4'] Formed, together with the (1'') and (1'') isomerides by nitration of naphthalene (α) sulphonic acid, the acids being separated by crystallisation of their chlorides from benzene (Cleve, *B* 23, 958) Yellowish crystalline mass — KA' — $\text{NaA}'_2, 3\text{aq}$ — $\text{CaA}'_2, 2\text{aq}$ S 27 at 17°, 7 at 100° — $\text{BaA}'_2, 3\text{aq}$ S 15 in the cold, 3 at 100° — $\text{PbA}'_2, 6\text{aq}$ — AgA' needles

Methyl ether MeA' [117°]

Ethyl ether EtA' [93°]

Chloride $\text{C}_{10}\text{H}_6(\text{NO}_2)(\text{SO}_2\text{Cl})$ [99°]

Amide [188°] Octahedra

Nitro naphthalene sulphonic acid

$\text{C}_{10}\text{H}_6(\text{NO}_2)(\text{SO}_3\text{H})$ [1 1'] A product of the nitration of naphthalene (α) sulphonic acid (Cleve, *B* 28, 962)

Chloride [101°]

' α ' Nitro naphthalene disulphonic acid

$\text{C}_{10}\text{H}_4(\text{NO}_2)_2(\text{SO}_3\text{H})_2$ [4 2 2'] Obtained from its chloride, which is formed by nitrating naphthalene ' α ' disulphonic chloride $\text{C}_{10}\text{H}_4(\text{NO}_2)_2(\text{SO}_2\text{Cl})_2$ (Alén, *Bl* [2] 89, 63, *Bn* 2, 156) Needles, v e. sol water — $\text{Na}_2\text{A}'_2, 6\text{aq}$ — $\text{K}_2\text{A}'_2, 3\text{aq}$ — $\text{CaA}'_2, 5\text{aq}$ — $\text{BaA}'_2, 5\text{aq}$ — $\text{PbA}'_2, 4\text{aq}$ — $\text{Ag}_2\text{A}'_2, 3\text{aq}$

Chloride $\text{C}_{10}\text{H}_4(\text{NO}_2)_2(\text{SO}_2\text{Cl})_2$ [141°] With PCl_5 it yields θ -tri chloro naphthalene

Amide [287°]

' β ' Nitro naphthalene disulphonic acid

$\text{C}_{10}\text{H}_4(\text{NO}_2)_2(\text{SO}_3\text{H})_2$ [1 3 2'] Formed by nitrating naphthalene ' β ' disulphonic acid (Alén) Scales — $\text{Na}_2\text{A}'_2, 2\text{aq}$ — $\text{K}_2\text{A}'_2, 2\text{aq}$ — $\text{CaA}'_2, 2\text{aq}$ — $\text{BaA}'_2, 2\text{aq}$ — $\text{PbA}'_2, 2\text{aq}$ — $\text{Ag}_2\text{A}'_2, 2\text{aq}$

Chloride $\text{C}_{10}\text{H}_4(\text{NO}_2)_2(\text{SO}_2\text{Cl})_2$ [186°]

With PCl_5 it gives (γ) tri chloro naphthalene

Amide [above 300°] Needles

Di nitro naphthalene ' α '-disulphonic acid

$\text{C}_{10}\text{H}_4(\text{NO}_2)_2(\text{SO}_3\text{H})_2$ Obtained from its chloride which is formed by nitrating naphthalene ' α ' disulphonic acid (Alén) — $\text{K}_2\text{A}'_2, 4\text{aq}$ — $\text{BaA}'_2, 6\text{aq}$ — $\text{Ag}_2\text{A}'_2, 4\text{aq}$

Chloride [219°] Needles (from benzene)

Amide $\text{C}_{10}\text{H}_4(\text{NO}_2)_2(\text{SO}_2\text{NH}_2)_2$ [c 306°]

Peri NITRO (α) NAPHTHOIC ACID

$\text{C}_{10}\text{H}_6(\text{NO}_2)(\text{CO}_2\text{H})$ [1 1'] [215°] S 04 in the cold S (alcohol) 5 Formed, together with the (1,4')-isomeride by nitrating (α) naphthoic acid (Kuchenmeister, *B* 3, 739, Ekstrand, *B* 12, 1393, 18, 73, 2881, 19, 1138, *J pr* [2] 38, 156, 276) Prisms, v sol alcohol Yields on nitration di nitro naphthalene [170°]

Salts — NaA' — $\text{CaA}'_2, 3\text{aq}$ S 2 at 15° —

$\text{BaA}'_2, 6\text{aq}$ — $\text{PbA}'_2, 4\text{aq}$ yellow prisms

Ethyl ether EtA' [69°] Octahedra

Amide $\text{C}_{10}\text{H}_6(\text{NO}_2)(\text{CONH}_2)$ [280°] Needles, converted by conc HClAq at 170 into

$\text{C}_{10}\text{H}_6\text{Cl}_2 < \begin{smallmatrix} \text{NH} \\ \text{CO} \end{smallmatrix} >$ [261°]

Nitro (α) naphthoic acid

$\text{C}_{10}\text{H}_6(\text{NO}_2)(\text{CO}_2\text{H})$ [1 4'] [239°] S 02 at 15° S (alcohol) 5 at 15° Formed as above, and also by saponifying its nitrile which is obtained by nitrating the nitrile of (α) naphthoic acid (Graeff, *B* 14, 1063, 16, 2250, Ekstrand, *J pr* [2] 38, 241) Needles HNO_3 (S G 1.3) converts it into di nitro naphthalene [212°] — $\text{NaA}'_2, 5\text{aq}$ — $\text{CaA}'_2, 2\text{aq}$ S 63 at 15° — $\text{BaA}'_2, 3\text{aq}$ yellow needles — $\text{PbA}'_2, 5\text{aq}$

Methyl ether MeA' [110°]

Ethyl ether EtA' [93°]

Isopropyl ether PrA' [101°]

Nitrile $\text{C}_{10}\text{H}_5(\text{NO}_2)\text{CN}$ [205°]

(γ) Nitro (α -naphthoic acid [255°] Formed by heating its nitrile with HClAq at 160° (Graeff, *B* 16, 2252) Needles (by illumination)

Nitrile [153°] Formed, together with the isomeride [205°] by nitrating (α) naphthonitrile

' α '-Nitro (β) naphthoic acid [220°] Formed, as well as the four following acids, by nitration of (β) naphthoic acid (Ekstrand, *B* 12, 1325) Needles — CaA'_2 S 26 at 15°

Ethyl ether EtA' [82°] Needles

' β ' Nitro (β) naphthoic acid

$\text{C}_{10}\text{H}_6(\text{NO}_2)(\text{CO}_2\text{H})$ [1 or 4 2] [293°] Formed as above (Ekstrand, *B* 18, 1207, *J pr* [2] 42, 375) Obtained also by saponifying its nitrile (Graeff, *B* 16, 2252) Needles, m sol hot alc hol — $\text{KA}'_2, 3\text{aq}$ — $\text{NaA}'_2, 2\text{aq}$ — $\text{CaA}'_2, 3\text{aq}$ S 12 at 15° — $\text{BaA}'_2, 4\text{aq}$

Methyl ether MeA' [112°] Needles

Ethyl ether EtA' [109°] Needles

Isopropyl ether PrA' [76°] Needles

Nitrile $\text{C}_{10}\text{H}_5(\text{NO}_2)\text{CN}$ [173°] Formed by nitration of (β) naphthoic nitrile Needles
 (γ) Nitro- (β) naphthoic acid [269°] Formed as above Needles, v sol alcohol

Ethyl ether EtA' [93°]**(β) Nitro-(β)-naphthoic acid**

$C_{10}H_7(NO_2)(CO_2H)$ [4' or 1' 2] [288°] S (alcohol) 25 in the cold. Formed as above (Ekstrand, *J pr* [2] 42, 292). Needles. Yields di-nitro naphthoic acid [226°] with fuming HNO_3 , $-NH_4A'$ — NaA' 2aq — BaA' , 8aq — CaA' , 4½aq S 15 in the cold.

Ethyl ether EtA' [121°] Plates

(ε)-Nitro (β) naphthoic acid [285°] One of the products of the action of HNO_3 (SG 1 42) on (β) naphthoic acid (Ekstrand, *J pr* [2] 42, 304). Stellate needles (from alcohol).

Ethyl ether EtA' [75°] Needles**Di-nitro-(α) naphthoic acid**

$C_{10}H_6(NO_2)_2CO_2H$ [4' 1' 1] [265°] Obtained by nitration of (α) naphthoic acid and of (1,4')-nitro naphthoic acid (Ekstrand, *B* 17, 1600, 19, 1984, 20, 219, 1353, *J pr* [2], 38, 259). Prisms or needles (from alcohol). Reduced by tin and $HClAq$ to naphthylene diamine [65°] H S in alkaline solution forms $C_{10}H_7N_2CO_2H$ (?), a bluish violet pp — NaA' 6aq — BaA' , 2½aq — CaA' , 3aq needles S 7 in the cold.

Ethyl ether EtA' [143°] Needles**Di-nitro (α) naphthoic acid**

$C_{10}H_6(NO_2)_2CO_2H$ [215°] Formed, together with the preceding acid, by nitrating (α) naphthoic acid (Ekstrand, *B* 19, 1984, *J pr* [2] 38, 270). Needles or plates (from alcohol). Yields nitro amido naphthoic acid [c 110°] on reduction by H S and NH_3 — CaA' .

Ethyl ether EtA' [137°] Needles**Di-nitro-(α)-naphthoic acid**

$C_{10}H_6(NO_2)_2CO_2H$ [4' 1' 1] [218°] Formed by nitrating nitro (α) naphthoic acid [239°], and occurs among the products of the action of fuming HNO_3 on (α) naphthoic acid (Ekstrand, *B* 20, 220, *J pr* [2] 38, 267). Trimetric crystals (from alcohol), *abc* — 973 1 1 442. Yields a lactam of di amido naphthoic acid on reduction — CaA' , 7aq yellow needles, v e sol water.

Ethyl ether EtA' [129°] Needles**Di nitro-(β) naphthoic acid**

$C_{10}H_6(NO_2)_2CO_2H$ [226°] S (alcohol) 17 in the cold. Formed, together with the isomeride [148°] by dissolving (β) naphthoic acid in fuming HNO_3 . Formed also, in like manner, from nitro (β) naphthoic acid [288°] (Ekstrand, *B* 17, 1602, *J pr* [2] 42, 300). Needles (from alcohol). Yields by reduction nitro amido naphthoic acid [235°] and di amido naphthoic acid [c 230°] — NH_4A' aq S 35 in the cold — BaA' , 6aq — CaA' , 4aq S 06 in the cold.

Ethyl ether EtA' [141°] Needles**Di-nitro-(β)-naphthoic acid**

$C_{10}H_6(NO_2)_2CO_2H$ [1' 1' 2] or [4' 4' 2] [248°] S (alcohol) 16 in the cold. Formed as above, and also by nitrating nitro (β) naphthoic acid [293°] (Ekstrand, *B* 17, 1602, *J pr* [2] 42, 286). Rectangular prisms. Reduced by $FeSO_4$ and NH_3 to di amido naphthoic acid [202°] — NH_4A' aq — NaA' 4aq — BaA' , 8aq — CaA' , 5aq

Ethyl ether EtA' [165°]**Tri nitro-(α) naphthoic acid**

$C_{10}H_4(NO_2)_3CO_2H$ [288°] Formed in small quantity by the action of H_2SO_4 and HNO_3 on nitro (α) naphthoic acid [215°] (Ekstrand, *B* 19, 1181, *J pr* [2] 38, 372). Wedge shaped needles (from alcohol). Tastes very bitter — CaA' , 5aq

Ethyl ether EtA' [131°]. Prisms.**Tri nitro (α) naphthoic acid**

$C_{10}H_4(NO_2)_3CO_2H$ [236°] Formed from di nitro (α) naphthoic acid [265°], H_2SO_4 , and fuming HNO_3 (Ekstrand, *B* 19, 1987). Needles

Ethyl ether EtA' [191°] Needles**Tri-nitro (α) naphthoic acid**

$C_{10}H_4(NO_2)_3CO_2H$ [293°] A product of nitration of di nitro (α) naphthoic acid [265°] (E) Cubes, sl sol alcohol.

Ethyl ether EtA' [150°] Needles**NITRO (α) NAPHTHOL**

$C_{10}H_7(NO_2)(OH)$ [2 1] [128°] Formed, together with the isomeride [164°], from acetyl- or benzoyl naphthylamine by nitrating and boiling the product with $NaOHAq$ (Andreoni a Biedermann, *B* 6, 342, Liebermann a Dittler, *A* 183, 245, Worms, *B* 15, 1815, Lellmann, *B* 19, 802). Obtained also by oxidising (β) naphthoquinone (β) oxim (Fuchs, *B* 8, 629), and by the action of nitrous acid on (α) naphthylamine at 100° (Deninger, *J pr* [2] 40, 300). Yellow needles, sl sol water, m sol alcohol, volatile with steam — KA' aq — BaA' , 3aq red needles.

Nitro-(α) naphthol $C_{10}H_7(NO_2)(OH)$ [4 1]

[164°] Formed as above, and also by the action of (nascent) nitrous acid on (α) naphthylamine in the cold (Deninger, *J pr* [2] 40, 300). Golden needles (from water), not volatile with steam, v e sol alcohol. Yields di nitro naphthol [138°] on nitration. By successive reduction and oxidation it may be converted into (α) naphthoquinone — KA' — NaA' , 2aq crimson needles, bluish red when anhydrous — BaA' , aq — BaA' , 3aq (Hubner, *A* 208, 325) — CaA' , 3aq — PbA' , scarlet powder — AgA' .

Nitro-(β) naphthol **Ethyl ether**

$C_{10}H_7(NO_2)(OEt)$ [1 2] [73°] Formed, together with the ethyl ethers of the two following isomerides, by nitrating $C_{10}H_7(OEt)$ dissolved in $HOAc$ (Gaess, *J pr* [2] 43, 25). Needles.

Nitro-(β) naphthol **Ethyl ether**

$C_{10}H_7(NO_2)(OEt)$ [2 2] [114°] Needles. Yields on oxidation nitrophthalic acid [114°].

Nitro-(β)-naphthol $C_{10}H_7(NO_2)(OH)$ [1 2]

[103°] Formed by oxidising the (α) oxim of (β) naphthoquinone (Stenhouse a Groves, *C* 32, 51), or by boiling the acetyl derivative of nitro (β) naphthylamine with $NaOHAq$ (Liebermann a Jacobson, *A* 211, 46). Formed also by the action of (nascent) nitrous acid on (β) naphthylamine (D). Needles. By reduction followed by oxidation it may be converted into (β) naphthoquinone.

Acetyl derivative $C_{10}H_6(NO_2)(OAc)$ [61°]

Yields $C_{10}H_6(NHAc)(OH)$ on reduction by zinc dust and $HOAc$ (Botcher, *B* 16, 1933).

Benzoyl derivative $C_{10}H_6(NO_2)(OBz)$

[142°] Yields $C_{10}H_6(NHBz)(OH)$ on reduction.

Ethyl ether EtA' [104°] Yellow needles

(Witkamp, *B* 17, 393, Gaess, *J pr* [2] 43, 22).

Di nitro (α) naphthol

$C_{10}H_5(NO_2)_2(OH)$ [4 2 1] **Martiusyellow** [138°]

Formed by the action of HNO_3 on (α) naphthol, (α) naphthol sulphonic acid, and (2,1)- or (4,1)-nitro naphthol (Martius, *Z* [2] 4, 80, Darmstadter a Wichelhaus, *A* 152, 299, Liebermann, *A* 183, 249). Formed also by boiling diazonaphthalene chloride or sulphamate with dilute nitric acid (Martius, Neville a Winther, *C* 37, 632). Sulphur yellow crystals, nearly insol boiling water, sl sol alcohol. Not volatile with steam.

Yields phthalic acid on oxidation Its salts dye wool yellow Poisonous 4 g killed a dog (Weyl, *B* 21, 2191) $\text{—NH}_4\text{A}^+ \text{aq} \text{—Na}_2\text{A}^{2-} \text{aq} \text{—CaA}^+ 6\text{aq} \text{—SrA}^+ 3\text{aq} \text{—BaA}^+ 3\text{aq} \text{—AgA}^+$

Ethyl ether EtA' [88°] Needles

Di nitro-(\beta)-naphthol

$\text{C}_{10}\text{H}_7(\text{NO}_2)_2(\text{OH})$ [1' 2'] [194°] Formed by heating (\beta) naphthol with alcohol and HNO_3 (Wallach a Wichelhaus, *B* 3, 846) Prepared by boiling a solution of (\beta) diazonaphthalene chloride with HNO_3 (Graebe a Drews, *B* 17, 1170) Yellow needles Dyes deep yellow On oxidation it yields nitro phthalic acid [160°] $\text{—KA}^+ 2\text{aq}$ yellow needles $\text{—Ba}^+ \text{A}_2\text{aq}$ (Löwe, *B* 23, 2542) —AgA^+ scarlet pp

Ethyl ether EtA' [138°] (Graebe), [144°] (Gaess) Formed by nitrating the compounds $\text{C}_{10}\text{H}_7(\text{NO}_2)(\text{OEt})$ [114°] and [104°] Yellow needles Converted by NH_3 into di nitro naphthylamine [238°]

Di nitro-(\beta)-naphthol

$\text{C}_{10}\text{H}_7(\text{NO}_2)_2(\text{OH})$ [1' 2'] [198°] Formed from its ether by saponifying with alcoholic potash (Gaess) Plates Yields nitrophthalic acid [212° or 218°] on oxidation

Ethyl ether EtA' [215°] Obtained by nitrating $\text{C}_{10}\text{H}_7(\text{NO}_2)(\text{OEt})$ [1 2] (Gaess), and also from the [1 2'] isomeride Needles Possibly identical with the following isomeride

Di nitro-(\beta)-naphthol

$\text{C}_{10}\text{H}_7(\text{NO}_2)_2(\text{OH})$ [4' 1 2']

Ethyl ether EtA' [215°] Formed by nitrating $\text{C}_{10}\text{H}_7(\text{OEt})_2$ (Onufrovitch, *B* 23, 3356) Orange needles Yields (6, 3, 2, 1) di nitro phthalic acid on oxidation

Tri nitro-(\alpha)-naphthol $\text{C}_{10}\text{H}_5(\text{NO}_2)_3(\text{OH})$

[176°] S 28 at 15° Formed from di nitro (\alpha) naphthol, H_2SO_4 , and HNO_3 (Ekstrand, *B* 11, 161, Diehl a Merz, *B* 11, 1662) Small crystals Yields di nitro phthalic acid [213°] on oxidation $\text{—KA}^+ \text{aq}$ S 25 in the cold $\text{—NaA}^+ \text{aq}$ S 3 in the cold $\text{—NH}_4\text{A}^+$ S 15 in the cold $\text{—BaA}^+ 2\frac{1}{2}\text{aq}$ S 09 $\text{—CaA}^+ 3\frac{1}{2}\text{aq}$ S 4 —AgA^+

Methyl ether MeA' [128°] Got by nitrating $\text{C}_{10}\text{H}_7\text{OMe}$ (Staedel, *B* 14, 899, *A* 217, 172) Yellow plates

Ethyl ether EtA' [148°]

Tri nitro-(\beta)-naphthol $\text{C}_{10}\text{H}_5(\text{NO}_2)_3\text{OH}$

Methyl ether MeA' [213°] Got by nitrating $\text{C}_{10}\text{H}_7\text{OMe}$ (Staedel)

Ethyl ether EtA' [186°]

Tetra-nitro-(\alpha)-naphthol $\text{C}_{10}\text{H}_3(\text{NO}_2)_4\text{OH}$

[180°] S (benzene) 45 at 18° Formed by boiling $\text{C}_{10}\text{H}_7\text{Br}(\text{NO}_2)$, [170 5°] with $\text{Na}_2\text{CO}_3\text{Aq}$ (Merz a Werth, *B* 15, 2714) Yields di nitro phthalic acid [227°] on oxidation $\text{—NaA}^+ 2\text{aq}$ S 1 at 19° $\text{—CaA}^+ 2\text{aq}$ $\text{—BaA}^+ 3\text{aq}$ $\text{—AgA}^+ 3\text{aq}$ red needles

DI-NITRO-(\alpha)-NAPHTHOL SULPHONIC ACID $\text{C}_{10}\text{H}_5\text{N}_2\text{SO}_5$

$\text{C}_{10}\text{H}_4(\text{SO}_3\text{H})\text{C}(\text{OH})(\text{NO}_2)\text{C}(\text{NO}_2)\text{OH}$ *Naphthol yellow*

S Formed by warming (\alpha)-naphthol trisulphonic acid with dilute nitric acid at 50° (Lauterbach, *B* 14, 2028) Obtained also from mono- or di sulphonic acids of (\alpha)-naphthol in which one SO_3H is in a different ring to the hydroxyl. Long yellow needles. Very powerful dye, producing a very fast greenish-yellow shade—

KA^+ v sl sol cold water Not poisonous (Weyl, *B* 21, 2191)

Di nitro-(\alpha)-naphthol sulphonic acid

$\text{C}_{10}\text{H}_4(\text{NO}_2)_2(\text{OH})(\text{SO}_3\text{H})$ [4' x 2 4] *Crocein yellow* Formed by warming (\beta) naphthol (\alpha) sulphonic acid with dilute HNO_3 (Nietzki a Zübelen, *B* 22, 454) —KA^+ golden scales

NITRO-(\beta)-NAPHTHOQUINONE

$\text{C}_{10}\text{H}_6(\text{NO}_2)_2\text{O}$ [158°] Formed by nitrating (\beta) naphthoquinone (Stenhouse a Groves, *A* 194, 203, *C J* 33, 416, 45, 299) Crimson plates (from HOAc), sl sol ether and water With alcoholic aniline it forms the compound

$\text{C}_{10}\text{H}_6(\text{NO}_2)(\text{OH})\text{C}_6\text{H}_5\text{N}$ [253°] (Korn, *B* 17, 908)

p Bromo aniline forms the corresponding body

$\text{C}_{10}\text{H}_6(\text{NO}_2)(\text{OH})\text{C}_6\text{H}_4\text{Br}$ [245°] (Brauns, *B*

17, 1133) The corresponding *o* toluidine melts at 240°, the *p* toluidine at 241° Excess of aniline in benzene forms yellow needles of $\text{C}_{10}\text{H}_6\text{N}_2\text{O}$, [186°] Hydroxylamine hydrochloride in presence of HOAc forms a salt $\text{C}_{10}\text{H}_6\text{NO}_2(\text{NH}_2\text{OH})$ [141°], which on boiling with HOAc yields nitro hydro-(\beta) naphthoquinone (Zaerthling, *B* 23, 179)

Nitro-(\gamma)-naphthoquinone [208°] Formed by oxidising nitro acenaphthene (Quincke, *B* 21, 1460) Yellowish red needles yielding with aniline $\text{C}_{10}\text{H}_6\text{O}_2(\text{NO}_2)(\text{NHPH})$ [128°], and, with diphenylamine, $\text{C}_{10}\text{H}_6\text{O}_2(\text{NO}_2)(\text{NPh}_2)$

NITRO-(\alpha\alpha)-DINAPHTHYL $\text{C}_{12}\text{H}_8\text{C}_2\text{H}_4\text{NO}_2$

[188°] Formed from dinaphthyl, HNO_3 , and HOAc (Julius, *B* 19, 2549) Orange plates

Di-nitro-dinaphthyl $\text{C}_{10}\text{H}_6(\text{NO}_2)_2\text{C}_{10}\text{H}_6(\text{NO}_2)_2$

[280°] Formed by nitrating dinaphthyl (J) Yellow needles, insol alcohol, v sl sol C_6H_6

Tetra-nitro-dinaphthyl $\text{C}_{20}\text{H}_{10}(\text{NO}_2)_4$ Formed from dinaphthyl and fuming HNO_3 (Lossen, *A* 144, 77) Amorphous orange powder

Tetra-nitro-(\beta\beta)-dinaphthyl [150°] Formed from isodinaphthyl and HNO_3 (Staub a Watson

Smith, *C J* 47, 104) Amorphous yellow powder

NITRO-(\alpha)-NAPHTHYL-AMIDO-BENZOIC ACID $\text{C}_{10}\text{H}_7\text{NH C}_6\text{H}_4(\text{NO}_2)\text{CO}_2\text{H}$ [1 3 1] Formed by heating (\alpha) naphthylamine with bromo nitro benzoic acid (Heidenleben, *B* 23, 3158) Yields on reduction the amido-acid [90°] —NaA reddish brown powder

Ethyl ether EtA' [109°]

Nitro-(\beta)-naphthyl-amido-benzoic acid Formed in like manner from (\beta) naphthylamine (H) Brick-red crystals, insol water $\text{—NaA}'$

Ethyl ether EtA' [127 5°]

NITRO-(\alpha) NAPHTHYLAMINE

$\text{C}_{10}\text{H}_6(\text{NO}_2)(\text{NH}_2)$ [2 1] [144°] Formed by saponifying its acetyl derivative, which is produced, together with that of the (4, 1) isomeride [190°], by nitrating the acetyl derivative of (\alpha)-naphthylamine (Lellmann a Remy, *B* 17, 109, 19, 236, 796) Red monoclinic prisms (from alcohol) Converted by boiling alcoholic potash into nitro-naphthol [128°]

Acetyl derivative $\text{C}_{10}\text{H}_6(\text{NO}_2)(\text{NHAc})$ [199°] With 1 mol of the (4, 1)-isomeride it forms a compound [170°]

Di-acetyl derivative $\text{C}_{10}\text{H}_6(\text{NO}_2)(\text{NAc}_2)$ [115°] Got by using Ac_2O at 140°

Benzooyl derivative $\text{C}_{10}\text{H}_6(\text{NO}_2)(\text{NHBz})$ [175°] (Worms, *B* 15, 1814) Yields benzenyl-

naphthylamine [210°] on reduction with tin and HCl (Hubner, *A* 208, 324)

Nitro (α)-naphthylamine

$C_{10}H_7(NO_2)(NH_2)[4\ 1]$ [190°] Formed as above (Liebermann, *A* 183, 232, *L* & *R*) Orange needles (from alcohol) Yields naphthylene-diamine [186°] on reduction

Acetyl derivative [190°]

Di acetyl derivative [144°]

Benzoyl derivative [224°] Yields nitro-naphthol [164°] on boiling with potash

Nitro (α) naphthylamine

$C_{10}H_7(NO_2)(NH_2)[4\ 1]$ [119°] Formed by reduction of 'α' di nitro naphthalene with alcoholic NH_3 and H_2S (Beilstein & Kuhlberg, *A* 169, 81) Small red needles (from water) On elimination of NH_3 it yields (α) nitro naphthalene - $B_2H_5SO_4$ 2aq needles, sl sol cold water

Sulphonic acid

$C_{10}H_7(NO_2)(NH_2)(SO_3H)[4\ 1\ 4]$ Formed from (1, 4) naphthylamine sulphonic acid by acetylation, nitration, and subsequent saponification (Nietzki & Zubelen, *B* 22, 451) Colourless needles Potash yields nitro naphthylamine

Nitro (β) naphthylamine

$C_{10}H_7(NO_2)(NH_2)[1\ 2]$ [127°] (*L* & *J*), [124°] (Meldola, *C* *J* 47, 520) Formed by saponifying its acetyl derivative, which is obtained by nitrating the acetyl derivative of (β) naphthylamine (Liebermann & Jacobson, *B* 14, 806, 1792, *A* 211, 42) Orange needles, sol hot water

Acetyl derivative $C_{10}H_7(NO_2)(NHAc)$ [124°] Reduced by tin and $HOAc$ to ethenyl (αβ) naphthylene diamine (Fischer & Hepp, *B* 20, 2473)

Di nitro (α)-naphthylamine $C_{10}H_6(NO_2)_2(NH_2)[4\ 2\ 1]$ [239°] (Witt, *B* 19, 2032) Obtained from its acetyl derivative, which is formed by nitrating $C_{10}H_7NHAc$ The saponification is effected by alcoholic NH_3 or by H_2SO_4 (Liebermann, *A* 183, 274, Meldola, *B* 19, 2683) Formed also by heating di nitro (α) naphthol with alcoholic NH_3 (Witt) Lemon yellow needles Readily converted by potash into di nitro α naphthol On elimination of NH_3 , it yields di nitro naphthalene [144°]

Acetyl derivative [250.5°] (Ebell, *A* 208, 330), [247°] (*L*) Needles

Benzoyl derivative [252°] Needles

Di nitro-(β) naphthylamine

$C_{10}H_6(NO_2)_2(NH_2)[242°]$ Formed by heating $C_{10}H_7(NO_2)(OEt)$ [144°] with aqueous NH_3 at 140° (Graebe & Drews, *B* 17, 1172, Gaess, *J* *pr* [2] 43, 31) On elimination of NH_3 it yields di nitro naphthalene [161.5°] or [167°] Two isomeric compounds $C_{10}H_6(NO_2)_2NHAc$ [185°] and [235°] are got by nitrating acetyl (β) naphthylamine (Maschke, *C* *C* 1886, 824)

Di-nitro (β) naphthylamine

$C_{10}H_6(NO_2)_2(NH_2)[1\ 1\ 2]$ [223°] Formed from $C_{10}H_7(NO_2)(OEt)$ [215°] (Gaess) Yields di nitro naphthalene [172°]

Di-nitro-naphthylamine obtained from the $C_{10}H_7(NO_2)_2(OEt)$ [215°] of Onufrovitch (*B* 23, 3362) carbonises at 235°-250°

Tri-nitro-(α)-naphthylamine

$C_{10}H_5(NO_2)_3(NH_2)[c\ 264°]$ Formed from $C_{10}H_7(NO_2)_2OEt$ and alcoholic NH_3 at 50° (Staedel, *B* 14, 901, *A* 217, 173) Yellow prisms (from toluene) On elimination of NH_3 it yields tri nitro naphthalene [181°]

Tri-nitro (β) naphthylamine. Formed from $C_{10}H_7(NO_2)_2OEt$ and NH_3 (*S*) Yellow needles Yields tri nitro-naphthalene [181°] on elimination of NH_3

Tetra nitro-(α)-naphthylamine

$C_{10}H_4(NO_2)_4NH_2$ [194°] Formed from $C_{10}H_7Br(NO_2)_3$ and NH_3 (Merz & Weith, *B* 15, 2718) Yellow needles, sl sol alcohol

Tetra-nitro-(β) naphthylamine [202°]

Formed from (β) bromo tetra nitro naphthalene and NH_3 (*M* & *W*) Yellow needles

Nitro-di-(β)-naphthylamine **Benzoyl derivative** $C_{10}H_5NBzC_{10}H_6NO_2$ [168°] Got by nitrating $(C_{10}H_7)_2NBz$ (Ris, *B* 20, 2625) Nodules (from alcohol) or prisms (containing C_6H_5) [95°] (from benzene) Yields, on reduction, benzenyl naphthylene naphthyl-diamine [163°]

Di nitro di (β)-naphthyl-amine

$C_{20}H_{12}(NO_2)_2NH$ [225°] Formed from di (β) naphthylamine, $HOAc$, and HNO_3 (Ris & Weber, *B* 17, 197)

Tetra nitro-di-(β)-naphthyl amine

$C_{20}H_{10}(NO_2)_4NH$ [286°] Formed by mixing di (β) naphthylamine, $HOAc$, and HNO_3 without cooling (*R* & *W*) Granules On further nitration it yields hexa-nitro di naphthyl amine

NITRO NAPHTHYLENE DIAMINE **Di-**

acetyl derivative $C_{10}H_7(NO_2)(NHAc)$ [244°] [*c* 295°] Formed by nitrating $C_{10}H_7(NHAc)$ (Kleemann, *B* 19, 335) Yellow needles, yielding phthalic acid on oxidation

DINITRO-(α) DINAPHTHYLENE-OXIDE

$C_{20}H_{10}(NO_2)_2O$ [270°] Prepared by nitration of (α) dinaphthylene oxide (Knecht & Unzeitig, *B* 13, 1725) Yellow needles

Dinitro (β) dinaphthylene-oxide

$C_{20}H_{10}(NO_2)_2O$ [221°] Prepared by nitration of (β) dinaphthylene oxide (*K* & *U*) Orange-red needles

TETRA NITRO DI NAPHTHYL METHANE

$CH_2(C_{10}H_7(NO_2)_2)_2$ Formed from di (α) naphthyl methane and HNO_3 (Grabowski, *B* 7, 1605) Colourless crystals, decomposing at 260°-270°

NITRO-(α) NAPHTHYL PIPERIDINE

$C_{10}H_5(NO_2)NC_5H_{10}$ [77°] Formed by heating (1, 4) bromo nitro naphthalene with piperidine (Lellmann & Buttner, *B* 23, 1387) Yellow needles Weak base

DI NITRO DI-(α) NAPHTHYL DISULPHIDE

$S(C_{10}H_6NO_2)_2$ [186°] Formed from nitro naphthalene sulphonic chloride and HI (Cleve, *B* 23, 960) Greenish yellow scales

NITRO-NITROSO-ANTHRONE $C_{14}H_9NO_3$

[263°] Formed by boiling hydroanthracene nitrite (vol 1 p 277) with alkalis (Liebermann & Landshoff, *B* 14, 470) Yellow needles, yielding anthraquinone on oxidation

NITRO-OCTANE $C_8H_{17}NO_2$ (205°-212°)

Formed by distilling octyl iodide with $AgNO_3$ (Eichler, *B* 12, 1883)

NITRO-OCTOIC ACID $C_8H_{15}(NO_2)_2O_2$ *S* *G*

A 1093 Formed by boiling the non volatile acids of cocoa nut oil for a long time with HNO_3 (Wirz, *A* 104, 289) Oil - AgA' - EtA' *S* *G* *A* 1031 Oil

NITRO-OCTYL-BENZENE $C_8H_7(NO_2)C_6H_5$

The three isomerides may be got by nitrating octyl benzene (Ahrens, *B* 19, 2721) *o* Nitro octyl benzene carbonises at 130°, the *m* isomeride melts at 124°, the *p*-variety melts at

204° They yield the corresponding nitro benzene acids on oxidation by KMnO_4 .

Di-nitro-octyl-benzene $\text{C}_8\text{H}_7(\text{NO}_2)_2\text{C}_8\text{H}_{17}$, [226°] Formed by nitration of octyl benzene

NITRO-OPICIANIC ACID ν **OPICIANIC ACID**

NITRO OREIN $\text{C}_8\text{H}_7\text{NO}_2$, ν

$\text{C}_8\text{H}_7\text{Me}(\text{NO}_2)(\text{OH})_2$ Two nitro oreins [120°] and [115°] are formed, together with an azo compound, by the action of a cold mixture of nitrous and nitric acids on an ethereal solution of orein (Weselsky, *B* 7, 439) They may be separated by steam distillation, the compound melting at 120° being volatile The nitro orein [120°] forms $\text{Ba}(\text{HA}'')_2$, the isomeride [115°] yields $\text{Ba}(\text{HA}'')_2 \cdot 8\text{aq}$ and $\text{BaA}'' \cdot 3\text{aq}$ By the action of HNO_3 and HNO_2 on the ethyl derivative of orein there are formed two compounds $\text{C}_8\text{H}_7\text{Me}(\text{NO}_2)(\text{OH})(\text{OEt})$, melting at 64° and 103° respectively (Weselsky a Benedikt, *M* 2, 871)

Di-nitro-orein $\text{C}_8\text{H}_7\text{Me}(\text{NO}_2)_2(\text{OH})_2$, [164 5°] *S* (alcohol) 5.5 at 15° Formed from di nitroso-orein and cold dilute HNO_3 (*S* G 1 3) (Stenhouse a Groves, *A* 188, 358) Yellow plates, m sol boiling water May be sublimed — $\text{Ba}(\text{HA}'')_2 \cdot \text{aq}$

Di-nitro-orein [110°] Obtained by saturating toluene with NO_2 (Leeds, *A C J* 2, 416, *B* 14, 483) Golden needles (from alcohol) Dyes silk yellow

Tri-nitro-orein $\text{C}_8\text{H}_7\text{Me}(\text{NO}_2)_3(\text{OH})_2$, [163°] Formed from orein, H_2SO_4 , and HNO_3 (Stenhouse, *Pr* 19, 410, Merz a Zetter, *B* 12, 681) Yellow needles, v sol hot water Explodes above 163° Dyes the skin yellow — $\text{K}_2\text{A}''$ orange needles — $\text{BaA}'' \cdot 3\text{aq}$ — PhA'' — $\text{Ag}_2\text{A}''$ amorphous orange red pp

Methyl ether MeA'' , [69 5°].

Ethyl ether EtA'' , [61 5°]

NITRO-OXINDOL $\text{C}_8\text{H}_7\text{ON}(\text{NO}_2)$ Prepared by adding powdered KNO_3 to a solution of oxindol in H_2SO_4 (Baeyer, *B* 12, 1313) Yellow needles, sol alcohol Begins to decompose at 175°

p NITRO-o-XY-ACETOPHENONE

$\text{C}_8\text{H}_7(\text{NO}_2)\text{COCH}_2\text{OH}$, [121°] Formed by boiling bromo nitro acetophenone with NaOAc and HOAc , diluting with water, and boiling again (Engler a Zulke, *B* 22, 204) Sol hot alkalis

Nitro-di-oxy-acetophenone

$\text{C}_8\text{H}_7(\text{NO}_2)(\text{OH})_2\text{COCH}_3$, [142°] Formed by nitration (Nencki a Sieber, *J pr* [2] 23, 537)

TETRA NITRO OXY AMIDO ANTHRAQUINONE $\text{C}_{14}\text{H}_7\text{N}_4\text{O}_{11}$, ν $\text{C}_{14}\text{H}_7(\text{NO}_2)_4(\text{NH}_2)(\text{OH})_2$

Chrysammidic acid Formed by boiling chrysammic acid $\text{C}_{14}\text{H}_7(\text{NO}_2)_4(\text{OH})_2$, with ammonia (Schunck, *A* 65, 236, Graebe a Liebermann, *A Suppl* 7, 310) Olive green needles, forming a dark purple aqueous solution Its salts explode when heated

NITRO OXY AMIDO BENZOIC ACID

$\text{C}_8\text{H}_7\text{N}_2\text{O}_5$, ν $\text{C}_8\text{H}_7(\text{NO}_2)(\text{OH})(\text{NH}_2)\text{CO}_2\text{H}$ [5 2 3 1] [220°] Got by reducing di nitro o oxy benzoic acid (Hubner a Babcock, *B* 12, 1345) Crystals

NITRO-OXY-AMIDO DIPHENYL *Acetyl*

derivative $\text{C}_8\text{H}_7(\text{NHAc})\text{C}_6\text{H}_5(\text{NO}_2)(\text{OH})$ [264°] (Schmidt a Schultz, *A* 207, 347) Yellow needles

NITRO-DI-OXY-DI AMIDO TRI PHENYL

METHANE *Di methyl-ether*

$\text{C}_8\text{H}_7(\text{NO}_2)(\text{OH})(\text{C}_6\text{H}_5(\text{OMe})\text{NH}_2)_2$, [108°] Formed by heating *p* nitro benzoic aldehyde with anis-

idine sulphate and H_2SO_4 (Fischer, *B* 15, 680) Golden needles (containing C_6H_5) *n* Anisidine yields an isomeric body [189°] (Kock, *B* 20, 1562)

NITRO DI OXY AMIDO QUINONE $\text{C}_{12}\text{H}_7\text{N}_2\text{O}_5$, ν $\text{C}_6(\text{NO}_2)(\text{OH})_2(\text{NH}_2)_2\text{O}_2$ The salt $\text{C}_6\text{H}_7\text{K}_2\text{N}_2\text{O}_5$ is precipitated on exposing a solution of $\text{C}_6(\text{NO}_2)(\text{NH}_2)(\text{OH})$, in aqueous K_2CO_3 to the air (Nietzki a Benckiser, *B* 18, 499) It forms coppery needles

NITRO OXY ANTHRAQUINONE *Ethyl*

ether $\text{C}_6\text{H}_4\text{<CO>C}_6\text{H}_4(\text{NO}_2)(\text{OEt})$ [1 2 3]

[243°] Formed by oxidation of the nitroso anthrone of the ethyl ether of nitro anthrol (Liebermann a Hagen, *B* 15, 1795) Colourless needles, sl sol alcohol

Nitro-di-oxy-anthraquinone $\text{C}_{14}\text{H}_7(\text{NO}_2)_2\text{O}_4$

(a) **Nitro alizarin** [196°] Prepared by nitration of the di acetyl derivative of alizarin (Schunck a Romer, *B* 12, 687) Less soluble in alcohol and HOAc than the (β) isomeride Oxidised by HNO_3 to phthalic acid Yields purpurin on warming with H_2SO_4

Nitro-di-oxy-anthraquinone (β) **Nitro alizarin** *Alizarin orange* [244°] Formed by nitration of alizarin dissolved in HOAc (*S* a *R*, Rosenstiel, *Bl* [2] 26, 63) Formed also by boiling di nitro oxy anthraquinone with dilute (20 p c) NaOH (Simon, *B* 15, 692) Orange needles (from benzene), sl sol KOH aq Dyes alumina mordants red and iron mordants reddish-violet

Di-acetyl derivative [218°] Needles

Nitro-tri-oxy-anthraquinone $\text{C}_{14}\text{H}_7(\text{NO}_2)_3\text{O}_4$ Formed from alizarin and fuming HNO_3 (Strecker, *Z* 1868, 264) Red crystalline powder — $\text{K}_2\text{A}''$

Di-nitro-oxy-anthraquinone

$\text{C}_6\text{H}_4\text{C}_6\text{O}_2\text{C}_6\text{H}_4(\text{NO}_2)_2(\text{OH})$ [1 3 2] [270°] Formed from oxy anthraquinone and fuming HNO_3 (Simon, *B* 14, 464, 15, 694) Yellow needles Dyes wool and silk orange — KA' — $\text{CuA}'_2 \cdot 2\text{aq}$ — $\text{MgA}'_2 \cdot 5\text{aq}$ — AgA'

Ethyl ether EtA' , [158°]

Di nitro di oxy anthraquinone

$\text{C}_{14}\text{H}_7(\text{NO}_2)_2\text{O}_4$ *Di nitro-purpuroxanthin* [250°] Formed from purpuroxanthin and cold HNO_3 (*S* G 1 48) (Plath, *B* 9, 1205) Red needles (from HOAc) — $\text{NH}_4\text{HA}''$ — BaA'' red needles

Tetra-nitro di oxy anthraquinone

$\text{C}_{14}\text{H}_7(\text{NO}_2)_4\text{O}_4$ *Tetra nitro anthraflavic acid* *Tetra nitro anthraflavone* Formed by boiling anthraflavic acid with HNO_3 (*S* G 1 4) (Schardinger, *B* 8, 1487) Yellow needles, melting with explosion at 307 6° cor — Salts — $(\text{NH}_4)_2\text{A}''$ — $(\text{NH}_4)_2\text{A}''\text{NH}_3$ — $(\text{NH}_4)_2\text{A}''\text{NH}_3$ — $\text{Ag}_2\text{A}''$ brownish needles

Tetra-nitro-di-oxy-anthraquinone

$\text{C}_{14}\text{H}_7(\text{NO}_2)_4\text{O}_4$ *Tetra-nitro isanthraflavic acid* Formed by nitrating isanthraflavic acid (Roemer a Schwarzer, *B* 15, 1045) Yellow plates (from dilute HNO_3) Its alkaline solution is red Dyes wool and silk orange yellow, but does not combine with mordants — $\text{KA}'' \cdot 2\text{aq}$ silky red needles

Tetra nitro-di-oxy-anthraquinone

$\text{C}_{14}\text{H}_7(\text{NO}_2)_4\text{O}_4$ *Tetra nitro anthrarufin* Formed by nitration of anthrarufin (Liebermann, *B* 12, 188) Small yellow plates (from fuming HNO_3) — $\text{Na}_2\text{A}'' \cdot 4\text{aq}$ — $\text{K}_2\text{A}'' \cdot \text{aq}$; bronze coloured prisms. $\text{MgA}'' \cdot 6\text{aq}$

Tetra-nitro di-oxy-anthraquinone

$C_{14}H_4(NO_2)_4O_4$ *Chrysammic acid* Formed by heating aloes with HNO_3 (S G 137) (Schunck, *A* 39, 1, 65, 235, Stenhouse & Muller, *C J* 19, 319, Tilden, *Ph* [3] 2, 845) Formed also by nitration of chrysazin (di oxy anthraquinone) (Liebemann & Giesel, *B* 8, 1643, 9, 329, *A* 183, 193) Golden plates or monoclinic prisms Explodes when quickly heated Aqueous potassium cyanide at 60° forms chrysocyanmic acid $C_{14}H_4N_2O_{12}$ 6aq, which forms the dark red salts $(NH_4)'A'$ 3aq, K_2A'' 3aq, and CaA'' 3aq

Salts — NaA'' 3aq (Mulder, *A* 68, 339, 72, 285) — K_2A'' 3 S 08 in the cold Flat rhomboidal plates, polarises light, the two rays being differently coloured (Brewster, *P* 69, 552, Hirschwald, *A* 183, 198) — BaA'' 2aq — BaA'' 4aq — MgA'' 5aq — PbA'' 5aq — $PbA''(OH)_2$ — PbA'' 4aq — MnA'' 5aq — CuA'' 4aq

Ethyl ether EtA'' Pale red needles or yellow prisms (Stenhouse)

Benzoyl derivative $C_{14}H_4BzN_4O_{12}$ Yellow, almost insoluble, prisms

NITRO-OXY-BENZENE NITRO PHENOL

Nitro tri oxy benzene Tri methyl derivative $C_6H_3(NO_2)_3(OMe)_3$ [x 123] [100°] Formed from $C_6H_3(OMe)_3$ and conc HNO_3 (Will, *B* 21, 612) Thick prisms with HNO_3 it yields $C_6H_3(NO_2)_3(OMe)_3$ [126°], sl sol alcohol

Di-nitro tetra-oxy-benzene Methylene di nitro apone

$C_6(NO_2)_2(OMe)_2 \begin{smallmatrix} O \\ \diagup \quad \diagdown \end{smallmatrix} CH$ [118°] Formed from apionic acid (4 g), $HOAc$ (50 cc), and HNO_3 (100 cc of SG 14) (Ciamician & Silber, *B* 22, 2489, 23, 2290) Yellow needles, insol water Yields on reduction $C_6H_5O_2(NH)_2$ [119°] With diacetyl it condenses to $C_6H_4N_2O_4$ [176°], and with benzil to $C_6H_5O \begin{smallmatrix} N \\ \diagup \quad \diagdown \end{smallmatrix} \begin{smallmatrix} Ph \\ CPh \end{smallmatrix}$ [222°], both crystallising in needles

NITRO-OXY-BENZOIC ACID

$C_6H_5(NO_2)(OH)(CO_2H)$ [5 21] *Nitro salicylic acid Amiotu acid* [228°] Mol w 183 S 176 at 22° (Hubner), 07 at 155° (Schiff) Formed, together with the isomeride [144°], by the action of nitric acid on salicylic acid and on indigo (Chevreul, *A Ch* [1] 72, 131, Buff, *A Ch* [2] 37, 160, Dumas, *A Ch* [2] 63, 205, [3] 2, 227, Gerhardt, *A Ch* [3] 7, 325, Marchand, *J pr* 26, 385, Piria, *A* 56, 35, Stenhouse, *A* 70, 253, Schiff, *A* 154, 14, Schiff & Masimo, *G* 9, 318, *A* 198, 258, Hubner, *A* 195, 6) Formed also by the action of aqueous $NaNO_2$ and H_2SO_4 on salicylic acid at 50° (Deninger, *J pr* [2] 42, 550), by boiling (e) nitro amido benzoic acid [263°] with potash (Griess, *B* 11, 1730), and by heating *p* nitro phenol with COI and alcoholic potash at 100° (Hasse, *B* 10, 2188) White needles, v sol alcohol Its aqueous solution is coloured red by $FeCl_3$ Distillation with CaO yields *p* nitro phenol [114°]

Salts — KA' — NH_4A' — BaA' 6aq — $BaO_2C_6H_4NO_2$ 2aq — CaA' 6aq — SrA' 5½aq — MgA' 4aq — ZnA' 5aq — AgA'

Methyl ether MeA' [88°] Formed from methyl salicylate and nitrous acid (Deninger, *J pr* [2] 40, 302)

Ethyl ether EtA' [94°] Needles.

Methyl derivative $C_6H_4(NO_2)(OMe)(CO_2H)$ [149°] Formed from

$C_6H_4(OMe)(CO_2H)$ and HNO_3 (Kraut, *A* 150, 6, Salkowski, *A* 173, 41) Needles Gives no colour with $FeCl_3$

Ethyl derivative $C_6H_4(NO_2)(OEt)CO_2H$ [163°] (P), [161°] (K) Formed from ethoxy benzoic aldehyde $C_6H_4(OEt)CHO$ and conc HNO_3 (Perkin, *A* 145, 311) Formed also from $C_6H_4(NO_2)(OH)CO_2Me$, EtI , and KOH , the resulting $C_6H_4(NO_2)(OEt)CO_2Me$ being saponified (Kraut) Plates (from boiling water) — BaA' 2aq — EtA' [98°] (Hubner)

Amide $C_6H_4(NO_2)(OH)CONH_2$ [225°] Needles Behaves as an acid, decomposing carbonates, and forming the salts $C_6H_4KN_2O_4$ aq — CaA' 4aq — BaA' 4aq — PbA' 4aq

Amide $C_6H_4(NO_2)(OH)CONHPh$ [224°] Formed by nitrating $C_6H_4(OH)CONHPh$ (Mensching, *A* 210, 343) Needles

Nitro-oxy-benzoic acid

$C_6H_4(NO_2)(OH)CO_2H$ [3 21] [144°] S 13 at 155° (Schiff) Formed as above, and also by adding H_2SO_4 to a solution of salicylic acid and $NaNO_2$ at 100° (Deninger, *J pr* [2] 42, 551), and by the action of nitrous acid on oil of wintergreen (Smith & Knapp, *Am* 8, 100) Obtained also by heating *o* nitro phenol with CCl_4 and alcoholic potash at 100° (Hasse) Needles (containing aq) Melts at 125° when hydrated, 144° when anhydrous $FeCl_3$ colours its solution red Yields *o* nitro phenol on heating with dilute H_2SO_4 at 150°

Salts — NaA' — KA' — BaA' Needles — $BaC_6H_4NO_2$ 1½aq — MgA' 2aq — $PbC_6H_4NO_2$ — AgA'

Ethyl ether EtA' [118°]

Amide $C_6H_4(NO_2)(OH)CONH_2$ [146°] Needles Yields the salts $Ca(C_6H_4NO_2)_2$ 4aq — BaA' 2aq — $Pb(OH)A'$ 2aq

Nitro-oxy benzoic acid $C_6H_4NO_2$ 12

$C_6H_3(NO_2)(OH)(CO_2H)$ [6 21] *Amide of the methyl derivative* $C_6H_3(NO_2)(OMe)CONH_2$ [195°] Formed by boiling its nitrile with baryta water (Lobry de Bruyn, *R T C* 2, 217) Yellowish needles, m sol acetone

Nitrile of the methyl derivative $C_6H_3(NO_2)(OMe)CN$ [171°] Formed by boiling *m* di nitro benzene with KCy and $MeOH$ (Lobry de Bruyn, *R T C* 2, 212) Needles (from alcohol) Boiling alcoholic potash converts it into $C_6H_4(OH)_2CO_2H$

Amide of the ethyl derivative $C_6H_4(NO_2)(OEt)CONH_2$ [197°] Needles

Nitrile of the ethyl derivative $C_6H_4(NO_2)(OEt)CN$ [137°] Formed from *m* di-nitro benzene and alcoholic KCy Tables

***a* Nitro-m oxy benzoic acid**

$C_6H_4(NO_2)(OH)CO_2H$ [6 31] [169°] Formed by boiling (a) nitro amido benzoic acid with potash (Griess, *B* 11, 1733) Light yellow crystals containing aq, v sol water — $BaC_6H_4NO_2$ 6aq orange prisms

Methyl derivative

$C_6H_4(NO_2)(OMe)(CO_2H)$ [188°] Formed by oxidising $C_6H_4(NO_2)(OMe)CHO$ with $KMnO_4$ (Rieche, *B* 22, 2354)

***β* Nitro-m oxy benzoic acid**

$C_6H_4(NO_2)(OH)CO_2H$ [4 31] [230°] Formed by boiling the corresponding nitro amido-benzoic acid with KOH aq and, together with the (2,3,1) and (5,3,1) isomerides, by nitration of a boiling aqueous solution of *m* oxy-benzoic acid by HNO_3

(Griess, *B* 5, 856, 20, 403). Yellow plates (from hot water) — $\text{BaC}_2\text{H}_3\text{NO}_3$, aq

Nitrile $\text{C}_6\text{H}_3(\text{NO}_2)(\text{OH})\text{CN}$ [183°] From *m* oxy benzoic nitrile and HNO_3 (Smith, *J pr* [2] 16, 227) Coloured dark red by aqueous FeCl_3 . Belongs perhaps to one of the isomeric nitro-*m*-oxy-benzoic acids

(*γ*)-Nitro-*m*-oxy-benzoic acid
 $\text{C}_6\text{H}_3(\text{NO}_2)(\text{OH})\text{CO}_2\text{H}$ [2 3 1] [178°] Formed by boiling (2,3,1)-nitro amido benzoic acid with potash, and also by nitrating *m* oxy benzoic acid (Griess, *B* 11, 1734, 20, 403) Yellow plates or prisms (containing aq) Tastes sweet — BaA'_2 , 13aq

Methyl derivative $\text{C}_6\text{H}_3(\text{NO}_2)(\text{OMe})\text{CO}_2\text{H}$ [251°] Formed from its aldehyde and Ag_2O (Rieche, *B* 22, 2352) White plates — AgA'

(*δ*)-Nitro-*m*-oxy-benzoic acid
 $\text{C}_6\text{H}_3(\text{NO}_2)(\text{OH})\text{CO}_2\text{H}$ [5 3 1] [167°] Formed by nitrating *m*-oxy benzoic acid (*o supra*) Crystals (containing aq) — BaA'_2 , 6aq prisms

Methyl derivative $\text{C}_6\text{H}_3(\text{NO}_2)(\text{OMe})\text{CO}_2\text{H}$ [233°] Long white needles (Rieche, *B* 22, 2356)

δ-Nitro-*p*-oxy-benzoic acid
 $\text{C}_6\text{H}_3(\text{NO}_2)(\text{OH})\text{CO}_2\text{H}$ [3 4 1] [185°]

Formation — 1 By dissolving *p* oxy-benzoic acid in dilute nitric acid (Barth, *Z* 1866, 647, Griess, *B* 20, 408) — 2 By boiling (3,4,1)-nitro-amido benzoic acid with potash (Griess, *B* 5, 856) — 3 By the action of aqueous NaNO_2 and H_2SO_4 on *p*-oxy benzoic acid at 40° (Deninger, *J pr* [2] 42, 552) — 4 In small quantity in the action of CCl_4 and alcoholic potash on *o* nitro phenol (H)

Properties — Needles, sl sol hot water Gives no colour with FeCl_3

Salt — $\text{BaC}_2\text{H}_3\text{NO}_3$, aq red nodules

Methyl derivative $\text{C}_6\text{H}_3(\text{NO}_2)(\text{OMe})\text{CO}_2\text{H}$
Nitransic acid [187°] Formed by boiling anisic acid or oil of anise with nitric acid (Laurer, *B* 23, 416, Cahours, *A Ch* 41, 71, Engelhardt, *A* 108, 243, Salkowski, *A* 163, 6, *B* 10, 1254) Also from $\text{C}_6\text{H}_3(\text{NO}_2)(\text{OMe})\text{CH}_2\text{NHAc}$ by oxidation with $\text{K}_2\text{Cr}_2\text{O}_7$ (Goldschmidt a Polonowska, *B* 20, 2410) Small needles (from water) Yields *o*-nitro phenol on heating with water at 220° — NaA'_2 aq — KA'_2 aq — CaA'_2 , 4aq — SrA'_2 , 4aq — BaA'_2 , flocculent pp — PbA'_2 — AgA'

Methyl ether of the methyl derivative
 $\text{C}_6\text{H}_3(\text{NO}_2)(\text{OMe})\text{CO}_2\text{Me}$ [108°] Plates

Ethyl ether of the methyl derivative
 $\text{C}_6\text{H}_3(\text{NO}_2)(\text{OMe})\text{CO}_2\text{Et}$ [100°] Tables

Nitrile of the methyl derivative
 $\text{C}_6\text{H}_3(\text{NO}_2)(\text{OMe})\text{CN}$ [150°] Formed from $\text{C}_6\text{H}_3(\text{OMe})\text{CN}$ and HNO_3 (Henry, *B* 2, 668) Small needles (from alcohol)

Nitro di-oxy-benzoic acid Di methyl derivative $\text{C}_6\text{H}_2(\text{NO}_2)(\text{OMe})_2\text{CO}_2\text{H}$ [4 5 3 1] [225°] Formed from $\text{C}_6\text{H}_2(\text{OMe})_2\text{CO}_2\text{H}$ and dilute HNO_3 (H Meyer, *M* 8, 431) Needles (from water) Yields on reduction, $\text{C}_6\text{H}_2(\text{NH}_2)(\text{OMe})_2\text{CO}_2\text{H}$ [182°] — NaA'_2 , 23aq — PbA'_2 — AgA' needles

Nitro-di-oxy-benzoic acid Methyl derivative $\text{C}_6\text{H}_2(\text{NO}_2)(\text{OH})(\text{OMe})\text{CO}_2\text{H}$ [x 4 3 1]
Nitro-xanthic acid White needles (Matsumoto, *B* 11, 122)

Acetyl derivative
 $\text{C}_6\text{H}_2(\text{NO}_2)(\text{OAc})(\text{OMe})\text{CO}_2\text{H}$ [182°] Needles

Di-methyl derivative
 $\text{C}_6\text{H}_2(\text{NO}_2)(\text{OMe})_2\text{CO}_2\text{H}$ *Nitro veratric acid*.

Yellow needles (containing 3aq) Its methyl ether melts at 144° and its ethyl ether at 100°

Di-nitro-oxy-benzoic acid

$\text{C}_6\text{H}_2(\text{NO}_2)_2(\text{OH})\text{CO}_2\text{H}$ [5 3 2 1] *Di nitro-salicylic acid* [173°] Formed by nitration of salicylic acid (Cahours, *A Ch* [8] 25, 11, Stenhouse, *A* 78, 1, Hübner, *A* 195, 45) Tables or needles (containing aq) After fusion it melts at 158° FeCl_3 colours its solution red — $\text{NH}_4\text{A}'$ — KA' sl sol water — $\text{K}_2\text{C}_2\text{H}_3\text{N}_2\text{O}$, aq m sol water — NaA' — BaA'_2 — $\text{BaC}_2\text{H}_3\text{N}_2\text{O}$, 3aq needles — CaA'_2 , 2aq — $\text{PbC}_2\text{H}_3\text{N}_2\text{O}$, 3aq — AgA' sparingly soluble granules

Methyl ether MeA' [127°] Yellowish scales Yields $\text{NH}_4\text{C}_2\text{H}_3\text{N}_2\text{O}$, and $\text{AgC}_2\text{H}_3\text{N}_2\text{O}$,

Ethyl ether EtA' [99°] Yields the salts $\text{NH}_4\text{C}_2\text{H}_3\text{N}_2\text{O}$, and $\text{AgC}_2\text{H}_3\text{N}_2\text{O}$,

Methyl derivative of the methyl ether $\text{C}_6\text{H}_2(\text{NO}_2)_2(\text{OMe})\text{CO}_2\text{Me}$ [69°] Prisms (Salkowski, *A* 173, 43)

Methyl derivative of the ethyl ether $\text{C}_6\text{H}_2(\text{NO}_2)_2(\text{OMe})\text{CO}_2\text{Et}$ [47°] Tables

Ethyl derivative of the methyl ether $\text{C}_6\text{H}_2(\text{NO}_2)_2(\text{OEt})\text{CO}_2\text{Me}$ [80°] Prisms

Ethyl derivative of the ethyl ether $\text{C}_6\text{H}_2(\text{NO}_2)_2(\text{OEt})\text{CO}_2\text{Et}$ [49°] Prisms

Di nitro p oxy benzoic acid

$\text{C}_6\text{H}_2(\text{NO}_2)_2(\text{OH})\text{CO}_2\text{H}$ [5 3 4 1] [237°] Formed by boiling di nitro-*p*-amido benzoic acid with potash (Salkowski, *A* 173, 36) Thin tables — KA' S 7 at 16 5° — $\text{K}_2\text{C}_2\text{H}_3\text{N}_2\text{O}$, 2aq S 32 at 16 5° — $\text{BaC}_2\text{H}_3\text{N}_2\text{O}$, 5aq — $\text{BaC}_2\text{H}_3\text{N}_2\text{O}$, 33aq — AgA' — $\text{AgC}_2\text{H}_3\text{N}_2\text{O}$,

Ethyl ether EtA' [87°] Yields $\text{KC}_2\text{H}_3\text{N}_2\text{O}$, S 9, and $\text{AgC}_2\text{H}_3\text{N}_2\text{O}$,

Methyl derivative $\text{C}_6\text{H}_2(\text{NO}_2)_2(\text{OMe})\text{CO}_2\text{H}$
Dinitransic acid [182°] Formed by nitration of nitro anisic acid (Salkowski a Rudolph, *B* 10, 1254, Stöhr, *A* 225, 86) — KA' aq — AgA'

Ethyl ether EtA' [79°]

Ethyl derivative of the ethyl ether $\text{C}_6\text{H}_2(\text{NO}_2)_2(\text{OEt})\text{CO}_2\text{Et}$ [59°] Needles

Tri nitro m oxy benzoic acid

$\text{C}_6\text{H}(\text{NO}_2)_3(\text{OH})\text{CO}_2\text{H}$ Formed by warming *m* diazoamide benzoic acid with HNO_3 (Griess, *A* 117, 28), and by the action of fuming HNO_3 on *m* amido benzoic acid (Beilstein a Geitner, *A* 139, 11) Large crystals (from conc HNO_3) Explodes when heated ($\text{NH}_4\text{C}_2\text{H}_3\text{N}_2\text{O}$, 2aq — $\text{BaC}_2\text{H}_3\text{N}_2\text{O}$, 3aq — $\text{AgC}_2\text{H}_3\text{N}_2\text{O}$,

Tri-nitro oxy benzoic acid [105°] Formed from *m*-oxybenzoic acid by successive treatment with dilute H_2SO_4 and HNO_3 (Schardinger, *B* 8, 1490) Tables and prisms (containing aq) — $\text{BaC}_2\text{H}_3\text{N}_2\text{O}$, 2aq — $\text{CaC}_2\text{H}_3\text{N}_2\text{O}$, 5aq green needles, exploding at 237°

NITRO-OXY BENZOIC ALDEHYDE

$\text{C}_6\text{H}_3(\text{NO}_2)(\text{OH})\text{CHO}$ [3 2 1] [109°] Formed together with the (5,2,1) isomeride, by nitrating salicylic aldehyde (Mazzara, *B* 6, 460, Von Miller, *B* 20, 1927, 22, 1709, Taegge, *B* 20, 2109) Its compound with NaHSO_3 does not crystallise Sodium amalgam forms brown amorphous $\{\text{C}_6\text{H}_3(\text{OH})(\text{CHO})\}_n\text{N}_2\text{H}_4$ (Briegleb, *A* 135, 169) — $\text{NaC}_2\text{H}_3\text{NO}$ — $\text{BaC}_2\text{H}_3\text{NO}$, 2aq orange crystals

Acetyl derivative [110°]

Phenyl-hydraside [165°]

Methyl derivative $\text{C}_6\text{H}_3\text{NO}_4$ [102°]

Nitro-oxy-benzoic aldehyde

$\text{C}_6\text{H}_3(\text{NO}_2)(\text{OH})\text{CHO}$ [5.2.1] [125°] Formed as

above Fo ms a solid compound with NaHSO_4 — NaA' 2aq needles — BaA' 6aq — BaA' 8aq

Acetyl derivative [119°]

Phenyl-hydrazide [186°]

Methyl derivative $\text{C}_6\text{H}_5(\text{NO}_2)(\text{OMe})\text{CHO}$ [90°] Formed by nitrating $\text{C}_6\text{H}_5(\text{OMe})\text{CHO}$ (Schnell, *B* 17, 1382, cf Voswinkel, *B* 15, 2027) Needles (from water)

Nitro-m-oxy-benzoic aldehyde

$\text{C}_6\text{H}_5(\text{NO}_2)(\text{OH})(\text{CHO})$ [2 3 1] [128°] Formed, together with the isomeride [166°], by nitration of *m*-oxy benzoic aldehyde (Tiemann & Ludwig, *B* 15, 2052, 3052) Yellow plates, sol benzene

Methyl derivative

$\text{C}_6\text{H}_5(\text{NO}_2)(\text{OMe})\text{CHO}$ [107°] (T & L), [102°] (R) Formed from the aldehyde, KOH, and MeI (T & L) Formed also, together with the (6, 3, 1) and (5, 3, 1) isomerides from $\text{C}_6\text{H}_5(\text{OMe})\text{CHO}$ and HNO_3 at 0° (Rieche, *B* 22, 2350) Forms a very soluble compound with NaHSO_4 With acetone and NaOH it condenses to di methoxy indigo $\text{C}_6\text{H}_5(\text{NO}_2)(\text{OMe})\text{CH NOH}$, its oxim, melts at 170°, and its phenyl hydrazide $\text{C}_6\text{H}_5(\text{NO}_2)(\text{OMe})\text{CH N}_2\text{HPH}$ at 134°

Nitro-m-oxy-benzoic aldehyde

$\text{C}_6\text{H}_5(\text{NO})_2(\text{OH})\text{CHO}$ [6 3 1] [166°] Formed by nitrating *m*-oxy benzoic aldehyde (T & L) Needles, v sl sol benzene

Methyl derivative [83°] Plates, vola-

tile with steam Its compound with NaHSO_4 is m sol water With acetone and KOH it yields a substance resembling indigo Its oxim $\text{C}_6\text{H}_5(\text{NO}_2)(\text{OH})\text{CH NOH}$ melts at 152°, and the corresponding phenyl hydrazide at 154°

Nitro-m-oxy-benzoic aldehyde *Methyl*

derivative $\text{C}_6\text{H}_5(\text{NO}_2)(\text{OMe})(\text{CHO})$ [5 3 1] [97°] (U), [104°] (R) A product of the nitration of [3 1] $\text{C}_6\text{H}_5(\text{OMe})\text{CHO}$ (Ulrich, *B* 18, 2572, Rieche, *B* 22, 2354) Does not give the indigo reaction Forms an oxim [148°], and a phenyl hydrazide [126°]

Nitro p oxy benzoic aldehyde

$\text{C}_6\text{H}_5(\text{NO})_2(\text{OH})\text{CHO}$ [3 4 1] [140°] Formed by nitrating *p*-oxy benzoic aldehyde (Mazzara, *G* 7, 285, Herzfeld, *B* 10, 1269) Needles, slightly volatile with steam — $\text{C}_6\text{H}_5(\text{NO}_2)(\text{OK})\text{CHO}$ aq — AgA' canary yellow pp

Methyl derivative *Nitro anisic alde-*

hyde [84°] Formed from anisic aldehyde, HNO_3 , and H_2SO_4 (Einhorn & Grabfield, *A* 243, 370) Yellow needles Forms a phenyl hydrazide [131°]

Di nitro m-oxy benzoic aldehyde *Methyl*

derivative $\text{C}_6\text{H}_5(\text{NO}_2)_2(\text{OMe})\text{CHO}$ [110°] Formed, together with an isomeride [155°], by treating [3 1] $\text{C}_6\text{H}_5(\text{OMe})\text{CHO}$ with H_2SO_4 and KNO_3 (Tiemann & Ludwig, *B* 15, 2055)

Nitro-di oxy benzoic aldehyde *Ethyl*

derivative $\text{C}_6\text{H}_5(\text{NO}_2)(\text{OEt})(\text{OH})\text{CHO}$ [5 2 1] [180°] Formed from $\text{C}_6\text{H}_5(\text{OEt})(\text{OH})\text{CHO}$ and dilute HNO_3 (Hantzsch, *J pr* [2] 22, 472) Scarcely volatile with steam

TETRA-NITRO o-OXY BENZYL ANILINE

$\text{C}_{10}\text{H}_5(\text{OH})(\text{NO}_2)_4\text{N}$ [86°] Formed from *o*-oxybenzyl aniline, HNO_3 , and H_2SO_4 (Emmerich, *A* 241, 345) Yellow needles (from benzene)

DI-NITRO-DI-OXY DI BENZYL UREA

Di-ethyl derivative $\text{CO}(\text{NH CH}(\text{OEt})\text{C}_6\text{H}_5(\text{NO}_2))_2$ [170°] Formed from urea, nitro-benzoic aldehyde, alcohol, and H_2SO_4 (Lüdy, *M* 10, 305).

NITRO-OXY-CINNAMIC ACID *o*. **NITRO-COUMARIC ACID**

NITRO-OXY-CUMINIC ACID *Methyl*

derivative $\text{C}_6\text{H}_3(\text{C}_6\text{H}_5)(\text{NO}_2)(\text{OMe})\text{CO}_2\text{H}$ [146°] Formed from the methyl derivative of cymophenol and dilute HNO_3 (Paterno & Canzoneri, *G* 10, 233) Yellowish crystals — BaA' 2½aq

Isomeride *v* **NITRO OXY ISOPROPYL BENZOIC ACID**

NITRO DI OXY ETHYL PYRIMIDINE

$\text{CO} \begin{smallmatrix} \text{Net CH} \\ \text{NH CO} \end{smallmatrix} \text{C}(\text{NO}_2)_2$ aq *Nitro-ethyl uracil* [194 5°] Formed from potassium nitro uracil and EtBr at 150° (Lehmann, *A* 253, 84) White needles — KA' — AgA' needles, v sl sol cold aq

DI NITRO OXY-ETHYL QUINOLINE

TETRAHYDRIDE *Ethyl derivative*

$\text{C}_{10}\text{H}_7\text{N}_2\text{O}_2$ *vs* $\text{C}_6\text{H}_5\text{Et}(\text{OEt})(\text{NO}_2)_2\text{N}$ [77°] Formed by nitrating $\text{C}_6\text{H}_5\text{Et}(\text{OEt})\text{N}$ (Kohn, *C J* 49, 509, *B* 19, 1048) Yellow prisms (from alcohol), with basic properties

NITRO OXY MALEIC IMIDE $\text{C}_6\text{H}_5\text{N}_2\text{O}_2$

The salt $\text{NH} \begin{smallmatrix} \text{CO CNO} \\ \text{CO COK} \end{smallmatrix}$ separates as light-yellow crystals when a solution of di chloro maleic imide is warmed with KNO_3 (Ciamician & Silber, *Rend Accad Linc* [4] 4, 447, *B* 22, 33, 2490). It is almost insol water

TETRA NITRO-DI-OXY METHYL-ANTHRAQUINONE $\text{C}_{14}\text{H}_2\text{Me}(\text{OH})_2(\text{NO}_2)_4\text{O}_2$ *Tetra nitro chrysophanic acid*

Formed by heating chrysophanic acid with fuming HNO_3 (Liebermann & Giesel, *A* 183, 175) Yellow plates or needles

NITRO-DI OXY METHYL-ETHYL-PYRIMIDINE

$\text{CO} \begin{smallmatrix} \text{NMe CH} \\ \text{Net CO} \end{smallmatrix} \text{C}(\text{NO}_2)_2$ [73°] *Nitro-methyl ethyl uracil* Formed from silver nitro-ethyl uracil and MeI at 130° (Lehmann, *A* 253, 86) Pearly rhombohedra (containing aq)

Nitro-di oxy methyl ethyl pyrimidine

$\text{CO} \begin{smallmatrix} \text{Net CH} \\ \text{NMe CO} \end{smallmatrix} \text{C}(\text{NO}_2)_2$ [109°] Formed from silver nitro methyl uracil and EtBr at 150° (L) Needles (containing aq)

NITRO-OXY-TRI METHYL-PYRIDINE

$\text{C}_6\text{H}_5(\text{NO}_2)\text{NO}$ *Nitro methyl ψ lutidostyryl* [161°] Formed by nitrating oxy tri methyl-pyridine (Hantzsch, *B* 17, 1032) Yellow needles or prisms, sol hot alcohol

NITRO-DI OXY METHYL PYRIMIDINE

$\text{C}_6\text{H}_5\text{N}_2\text{O}_2$ *vs* $\text{CO} \begin{smallmatrix} \text{NH CH} \\ \text{NMe CO} \end{smallmatrix} \text{CNO}_2$ *Nitro-methyl-uracil* [255°] S 714 at 20° Formed from potassium nitro-uracil and MeI at 140° (Hoffmann, *A* 253, 77) Needles (containing aq), v sol hot water Baryta water at 160° liberates methylamine and NH_3 — KA' needles — BaA' — AgA' minute white needles (from hot water)

Nitro-di-oxy-di-methyl pyrimidine

$\text{CO} \begin{smallmatrix} \text{NMe CH} \\ \text{NMe CO} \end{smallmatrix} \text{C}(\text{NO}_2)_2$ [154 5°] From the silver salt of nitro di oxy methyl pyrimidine and MeI at 120° (Lehmann, *A* 253, 82) Needles (containing aq) Yields NMeH , when heated with baryta at 180°

NITRO-OXY METHYL-QUINAZOLINE

$\text{C}_6\text{H}_5(\text{NO}_2) \begin{smallmatrix} \text{C(OH) N} \\ \text{N} \text{---} \text{CMe} \end{smallmatrix}$ Formed from oxy-methyl-quinazoline [288°] and conc. HNO_3 (De

hoff, *J pr* [2] 42, 348) Yellow crystalline powder, sol alcohol Does not melt below 280°

Methyl derivative [165°] Needles

(Py 2) **NITRO (Py 1) OXY (Py 3)-METHYL-QUINOLINE** $C_8H_7 \begin{smallmatrix} \diagup C(OH)C(NO_2) \\ \diagdown N=CMe \end{smallmatrix}$ Formed by nitrating oxy (Py 3) methyl quinoline (Conrad a Lumpach, B 20, 950) Needles, v sl sol boiling alcohol Does not melt at 270°

(B 3) Nitro (B 4) oxy (B 1) methyl quinoline $\begin{smallmatrix} CHCMe \\ C(NO_2)C(OH)CNOH \end{smallmatrix}$ [206°] This body

is formed by oxidising the corresponding nitroso oxy-methyl quinoline with alkaline K_2FeC_4 (Noelting a Trautmann, B 23, 3667) In the same way may be formed (B 4) nitro (B 1) oxy- (B 2)-methyl quinoline, (B 1) nitro (B 4) oxy- (B 3) methyl quinoline [193°], and (B 2) nitro- (B 1) oxy- (B 4) methyl quinoline [182°]

NITRO-OXY-TRI-METHYL QUINOLINE CARBOXYLIC ACID $C_8H_7N_3O_6$ Formed by warming (Py 1) oxy (B 1, 2, 4, Py 3) tetra methyl-quinoline with HNO_3 (S G 14) (Conrad a Lumpach, B 21, 529) White insoluble powder —NaA' aq yellowish needles

NITRO-DI OXY-NAPHTHALENE v NITRO-HYDRONAPHTHOQUINONE

Di-nitro di-oxy-naphthalene *Di ethyl-derivative* $C_{10}H_6(NO_2)_2(OEt)_2$ [229°] Formed from (e)-di chloro di nitro naphthalene and alcoholic potash (Alén, B [2] 36, 435) Yellow needles

NITRO-OXY-NAPHTHOIC ACID $C_{10}H_5(NO_2)(OH)CO_2H$ [242°] Formed by dissolving its anhydride in alkalis (Ekstrand, *J pr* [2] 38, 282) Plates — $CaC_2H_5NO_5$ aq yellowish-red needles

Anhydride $C_{10}H_5(NO_2) \begin{smallmatrix} O \\ \diagdown CO \end{smallmatrix}$ [242°]

Formed by nitrating *peroxy* oxy naphthoic acid Yellow needles (from HOAc)

Nitro-oxy-naphthoic acid [202°] (Schmidt a Burkard, B 20, 2700) Yields (β)-nitro (α) naphthol on distillation with lime

NITRO-OXY-(α)-NAPHTHOQUINONE

$C_{10}H_6NO_4$ is $C_6H_5 \begin{smallmatrix} COCNO_2 \\ COCOH \end{smallmatrix}$ *Nitronaphthalic acid* [157°] Formed from oxy (α)-naphthoquinone, H_2SO_4 , and HNO_3 , the yield being 85 p c of the theoretical (Diehl a Merz, B 11, 1317) Formed also from di chloro (α)-naphthoquinone and alcoholic $NaNO_2$ at 100° (Kehrmann, B 21, 1780, *J pr* [2] 40, 180) Yellow leaflets or tables — NH_4A' — NaA' aq — $KA'A'$ — BaA'_2 — PbA'_2 aq — PbA'_2 — AgA'_2

Sulphonic acid

$C_{10}H_6(SO_3H)(NO_2)(OH)O_2$ [2 3 2 4 1] Formed from di chloro (α) naphthoquinone sulphonic acid [229°] and $NaNO_2$ (K) — $K_2C_8H_5N_2SO_6$ (dried at 100°) slender yellow needles

DI-NITRO-DI OXY (ββ) DINAPHTHYL DISULFONIC ACID $C_{20}H_{12}(NO_2)_2(OH)_2(SO_3H)_2$ Got by nitrating $C_{20}H_{12}(OH)_2(SO_3)_2Ba$ (Julius, *Chem Ind* 10, 97) Yellow needles (containing 3aq)

o NITRO-p OXY-DIPHENYL

[4 1] $C_6H_4(OH)C_6H_5$ [1 2] [138°] Formed by the action of nitrous acid on the corresponding nitro amido diphenyl (Schultz a Strasser, B 14, 614, A 207, 351) Yellow needles

p Nitro p oxy-diphenyl

[4 1] $C_6H_4(OH)C_6H_4NO_2$ [1 4] [170°] Formed from p nitro p amido diphenyl (S a 3)

Nitro oxy-diphenyl $C_{12}H_9(NO_2)(OH)$ [67°] Formed by nitration of oxydiphenyl (Latschhoff, *J R* 5, 52) Lemon yellow prisms (from ether)

Di-nitro-oxy-diphenyl $C_{12}H_7(NO_2)_2(OH)$ [154°] Formed at the same time as the preceding (L) Golden plates (from alcohol) — KA' , 2aq sparingly soluble plates

Di nitro di-oxy-diphenyl

[3 4 1] $C_6H_4(NO_2)(OH)C_6H_4(NO_2)(OH)$ [1 3 4] [272°] (K), [280°] (S) Formed from *pp* di-oxy diphenyl and HNO_3 (S G 1 45) (Kunze, B 21, 3331, Schutz, B 21, 3531) Yellow needles, insol alcohol

Di-acetyl derivative [215°] Needles

Di-benzoyl derivative [206°] Plates

Ethyl ether [193°] Formed by nitrating the ethyl ether of di oxy diphenyl (Husch, B 22, 336)

Di-nitro-di-oxy-diphenyl $C_{12}H_7N_2O_6$ [184°]

Formed by oxidising o nitro phenol with aqueous $KMnO_4$ (Goldstein, *J R* 6, 193, 10, 318) Yel low needles (from benzene)

Di-benzoyl derivative [191°] Needles

Tetra-nitro-di-oxy-diphenyl

[4 3 5 1] $C_6H_4(OH)(NO_2)C_6H_4(NO_2)(OH)$ [1 3 5 4] [220°] (K), [225°] (S) From di-oxy diphenyl, HOAc, and HNO_3 (Kunze, B 21, 3333, Schutz, B 21, 3532) Yellow needles — NaA'' — $NaHA''$ brownish red needles

Di-acetyl derivative [236°] Needles

Hexa-nitro tetra-oxy-diphenyl $C_6H_4N_6O_{10}$ *Hexa nitro duresorein* Formed by warming tetra acetyl duresorein with fuming HNO_3 (Bene dikt a Julius, M 5, 178) Yellow crystals, exploding at 230°, v e sol water

m NITRO α OXY PHENYL ACETIC ACID v NITRO MANDELIC ACID

NITRO OXY PHENYL AMIDO BENZOIC ACID $C_8H_5(OH)NH C_6H_4(NO_2)CO H$ [4 3 1] [261°] Formed from bromo nitro benzoic acid, alcohol, and amido phenol at 120° (Schopff, B 22, 3248) Small needles, m sol water

DI NITRO α OXY DIPHENYLAMINE

[2 1] $C_6H_4(OH)NH C_6H_4(NO_2)$ [1 2 4] [199°] Got from $C_6H_5Br(NO_2)_2$ and o amido phenol (Schopff, B 22, 900) Orange crystals (from alcohol)

Acetyl derivative [150°] Needles

Ethyl derivative [164°] Red needles.

Methyl derivative [161°] Needles

Di-nitro p oxy-diphenylamine *Di benzoyl derivative* $C_{12}H_7(NO_2)_2(OBz)NBz$ [195°] Got by nitrating $C_{12}H_7(OBz)NBz$ (Philip a Calm, B 17, 2437) Small crystals, sl sol alcohol

Di-nitro-di-oxy-diphenylamine *Diethyl derivative* $C_{12}H_9NHC_6H_4(NO_2)_2(OEt)_2$ [133°]

Formed by heating aniline with the diethyl derivative of di nitro hydroquinone (Dietzki, A. 215, 157) Red needles (from alcohol)

NITRO-OXY-PHENYL-ANGELIC-(β) LACTONE

[4 1] $C_6H_4(NO_2)CHCH \begin{smallmatrix} CH_2 \\ O \end{smallmatrix} CO$ [111°] Formed by adding soda to a cold solution of $C_6H_4(NO_2)C_6H_4BrCH \begin{smallmatrix} CH_2 \\ O \end{smallmatrix} CO$ (Einhorn a Gehrenbeck, B 22, 47, A 253, 870)

o-NITRO-β-OXY PHENYL BUTYLENE DI-CARBOXYLIC ACID

[1 2] $C_6H_4(NO_2)CHCHCH(OH)CH(CO_2H)_2$

[269°] Formed by heating malonic acid (10 g) with *o*-nitro cinnamic aldehyde (15 g) at 125° (Einhorn) *A* 253, 375) Stellate needles

NITRO-OXY-PHENYL-CARBAMIC ETHER

Ethyl derivative $C_6H_4(NO_2)(OEt)NHCO_2Et$ [71°] Formed, as well as two di-nitro-derivatives [141°] and [121°] and a tri-nitro derivative [212°] by the action of nitric acid on [4 1] $C_6H_4(OEt)NHCO_2Et$ (Köhler, *J pr* [2] 29, 261) All four compounds crystallise from alcohol in needles

NITRO DI OXY PHENYL CROTONIC ACID

Anhydride $C_6H(NO)(OH)<\overset{CMe}{O}-CO$ Nitro (8) methyl umbelliferone Formed, as well as the di-nitro derivative [220°] by nitration of (8) methyl umbelliferone dissolved in HOAc (Fechmann & Cohen, *B* 17, 2136) Both compounds crystallise in yellow needles, sol alcohol

NITRO-OXY-PHENYL-ETHYLENE

NITRO- α -STYRENE

o-NITRO- β -OXY-PHENYL ETHYL METHYL KETONE

$C_{10}H_{11}NO_3$ *ie* $C_6H_4(NO_2)CH(OH)CH_2COCH_3$ [69°] Formed from *o*-nitro benzoic aldehyde, acetone, and dilute aqueous NaOH (Baeyer & Drewsen, *B* 15, 2857) Prisms

p-Nitro- β -oxy-phenyl-ethyl methyl ketone [58°] Formed in like manner from *p*-nitro benzoic aldehyde (Baeyer & Becker, *B* 16, 1969) Crystals Yields nitro styryl methyl ketone on boiling with AcO Boiling potash forms a compound ($C_{10}H_9NO_3$)_n [254°]

p-NITRO- β -OXY-PHENYL (Py 3) ETHYL-QUINOLINE $C_6H_4(NO_2)CH(OH)CH_2(CNCH_2)_3$ [160°] Formed by heating (Py 3) methyl quinoline with *p*-nitro benzoic aldehyde at 120° (Bulach, *B* 20 2016) Silky needles (from alcohol) —B' H_2PtCl_6 —B' HNO_3 white needles

DI-NITRO DI OXY DI PHENYL-HYDRAZINE

Di-ethyl derivative $\{C_6H_4(NO_2)(OEt)\}_2N_2H_2$ [202°] Formed by reduction of the corresponding azo compound [285°] by alcoholic ammonium sulphide (Andree, *J pr* [2] 21, 325) Yellow prisms, insol cold alcohol Hot HClAq converts it into nitro amido phenol and $\{C_6H_4(NO)(OEt)\}_2N_2$

NITRO OXY-PHENYL-METHYL PYRAZOLE

$C_{10}H_9N_3O_2$ *ie* $C_6H_4N<\overset{COCH(NO_2)}{N}CMe$ [127°-130°] Formed by the action of nitrous acid on oxy phenyl methyl pyrazole, and of nitric acid on the oxim thereof (Knorr, *A* 238, 187) Prisms (from alcohol), insol acids

NITRO-OXY-PHENYL-PROPIOLIC ACID

Methyl derivative $C_6H_4(NO_2)(OMe)CCO_2H$ [135°] Formed from $C_6H_4(NO_2)(OMe)CHBrCHBrCO_2H$ and alcoholic potash (Einhorn & Grabheld, *A* 243, 377) White needles, sol water

NITRO-OXY-PHENYL-PROPIONIC ACID

[8 4 1] $C_6H_4(NO_2)(OH)CH_2CH_2CO_2H$ Nitro-hydro-*p*-coumaric acid [91°] Formed by nitrating *p*-oxy phenyl propionic acid (Stöhr, *A* 225, 57) Orange needles (from water)

Methyl ether MeA' [64°] Needles

Ethyl ether EtA' [38°] Needles

Nitro- α -oxy-phenyl-propionic acid

$C_6H_4(NO_2)CH_2CH(OH)CO_2H$ A mixture of the *o*- and *p*- isomerides is formed by the action of fuming HNO_3 at —5° on α -oxy phenyl pro-

piotic acid (Erlenmeyer & Lupp, *A* 219, 228) The nitrate $C_6H_4(NO_2)CH_2CH(ONO_2)CO_2H$ of the *p* isomeride crystallises from hot water in needles, leaving that of the *o* compound in solution

o-Nitro- β -oxy-phenyl-propionic acid $C_6H_4NO_3$ *ie* [2 1] $C_6H_4(NO_2)CH(OH)CH_2CO_2H$ [126°] Formed by oxidation of the product of condensation of *o*-nitro benzoic aldehyde with acetic aldehyde (Baeyer & Drewsen, *B* 16, 2206) Formed also from β bromo *o*-nitro phenyl propionic acid and hot Na_2CO_3 Aq (Einhorn, *B* 16, 2214, 17, 1660, 2013) Monoclinic prisms (from water) Dilute H_2SO_4 at 190° converts it into *o*-nitro cinnamic acid — BaA_2 2aq needles

Methyl ether MeA' [51°]

β Lactone $C_6H_4(NO_2)CH<\overset{CH_2}{O}>CO$

[124°] Formed from $C_6H_4(NO_2)CHBrCH_2CO_2H$ and cold aqueous Na_2CO_3 Yellow monoclinic crystals (from chloroform) Split up by boiling with water into *o*-nitro styrene and CO_2 Boiling HOAc yields indigo

Amide $C_6H_4NO_3N_2O$ [197°] Formed from β bromo *o*-nitro phenyl propionic acid and ammonia Formed also from the lactone and NH_3 (Einhorn, *B* 16, 2646, Basler, *B* 17, 1494) Needles, v sol water Yields with AcO an acetyl derivative $C_6H_4AcN_2O$ [142°], a compound $C_6H_4N_2O_3$ [c 80°], and the acetyl derivative thereof $C_6H_4AcN_2O_3$ [172°]

m-Nitro- β -oxy-phenyl propionic acid

[3 1] $C_6H_4(NO)CH(OH)CH_2CO_2H$ [105°] Formed by boiling β bromo *m*-nitro phenyl propionic acid with water (Prausnitz, *B* 17, 596, 1660) Plates (from water)

Ethyl ether EtA' [56°] Crystals

Lactone $C_6H_4NO_3$ [98°] Deposited from a cold solution of the sodium salt Yields *m*-nitro styrene on boiling with water

p-Nitro- β -oxy phenyl-propionic acid

$C_6H_4NO_3$ [132°] Formed by the action of alkalis on β bromo *p*-nitro phenyl propionic acid, its lactone being the intermediate product (Basler, *B* 16, 3004, 17, 1494) Needles, m sol cold Aq

Methyl ether MeA' [74°] Prisms

Ethyl ether EtA' [46°] Crystalline

Lactone $C_6H_4NO_3$ [92°] Yields *p*-nitro styrene when boiled with HOAc

Amide $C_6H_4NO_3N_2O$ [166°] Prisms (from alcohol) Forms with Ac_2O an acetyl derivative [146°-150°]

Anilide $C_6H_4N_2O$ [176°] Plates

p-Nitro- α -di-oxy-phenyl-propionic acid

$C_6H_4NO_3$ *ie* $C_6H_4(NO_2)CH(OH)CH(OH)CO_2H$ [168°] Formed from *p*-nitro phenyl glycidic acid and diluted H_2SO_4 (Lupp, *B* 19, 2645) Plates, sl sol cold water

o-Nitro- m -di-oxy phenyl propionic acid

Methyl derivative

[2 5 1] $C_6H_4(NO_2)(OMe)CH(OH)CH_2CO_2H$ [106°] Got from $C_6H_4Cl(NO)CH(OH)CH_2CO_2H$ and NaOMe (Eichengrün & Einhorn, *B* 23, 1491) Colourless plates (from water)

ap-Di-nitro- β -oxy-phenyl-propionic acid.

Methyl derivative

$C_6H_4(NO_2)CH(OMe)CH(ONO_2)CO_2H$ Methyl ether MeA' [118°] Formed from methyl *ap*-di-nitro cinnamate by boiling with MeOH (Friedlander & Mahly, *B* 16, 851, *A* 229, 210).

Q Q

Ethyl ether EtA' [77°]. Formed in like manner, using EtOH

Ethyl derivative

$C_6H_4(NO_2)CH(OEt)CH(NO_2)CO_2H$ Methyl ether MeA' [110°] Formed by boiling *o*-nitro-cinnamic ether with alcohol Monoclinic prisms $a b c = 849 \text{ 1 517}$, $\beta = 87^\circ 25'$ Sol potash Yields the salts $Ba(C_{12}H_{11}N_2O_7)_2$ and $AgC_{12}H_{11}N_2O_7$. Ethyl ether EtA' [52°]. Monoclinic crystals

Di-nitro-oxy-phenyl-propionic acid $C_6H_4N_2O_7$, te [5 3 4 1] $C_6H_4(NO_2)_2(OH)CH_2CH_2CO_2H$ Di-nitro hydro *p*-coumaric acid [138°] Formed by nitrating oxy phenyl propionic acid (Stöhr, A 225, 68) Trimetric prisms (from HOAc) — NH_4A' [230°] — $(NH_4)_2A'$ — $AgHA'$ — AgA' dark red needles

Methyl ether MeA' [87°] Needles Yields $AgMeA'$ on adding Ag_2CO_3 to its ethereal solution

Ethyl ether EtA' [175°] Yields red needles of $AgEtA'$

Methyl derivative

$C_6H_4(NO_2)_2(OMe)CH_2CH_2CO_2H$ [124°] Got by saponifying its methyl ether with H_2SO_4 and HOAc Needles or plates (from dilute alcohol) Methyl ether $C_6H_4(NO_2)_2(OMe)C_6H_4CO_2Me$ [53°] From the basic silver salt and MeI Ethyl ether EtA' [71°] Needles, v sl sol Aq

Ethyl derivative

$C_6H_4(NO_2)_2(OEt)CH_2CH_2CO_2H$ [126°] Needles Methyl ether MeA' [36°] Ethyl ether EtA' [50°] Needles or plates

Di-nitro oxy-phenyl-propionic acid $C_6H_4N_2O_7$, *Dinitromeliotic acid* [155°] Formed by nitrating meliotic acid (Zwenger, A Suppl 5, 118) Prisms (from alcohol) — BaA' aq — Ag_2A'

Two isomeric acids are obtained by nitration of phloretic acid (Hlasiwetz, A 102, 155) Both form yellow crystals

o-NITRO- β -OXY-PHENYL PROPIONIC ALDEHYDE $C_6H_4NO_3$, te

[1 2] $C_6H_4(NO_2)CH(OH)CH_2CHO$ Formed from *o*-nitro benzoic aldehyde, acetic aldehyde, and 2 p c aqueous NaOH (Baeyer & Drewsen, B 16, 2205) Crystalline Forms a compound with aldehyde $C_6H_4NO_3(C_2H_5O)$ [125°], which yields indigo on treatment with alkalis Similar compounds of the formula $C_6H_4NO_3(C_2H_5O)$ are formed by the action of *m*- and *p*-nitro benzoic aldehydes on aldehyde and NaOH aq The *m* compound gives off aldehyde at 100°, the *p* compound melts at about 115° (Göhring, B 18, 372, 720)

m-NITRO- β -OXY-PHENYL-PYROTARTARIC ACID Lactone $C_{11}H_7NO_7$, te

$C_6H_4(NO_2)CH \begin{smallmatrix} O \\ \diagup \diagdown \\ CO \\ \diagdown \diagup \end{smallmatrix} CH_2$ Nitro-phenyl-paraconic acid [171°] Formed by heating *m*-nitro benzoic aldehyde with sodium succinate and Ac_2O at 125° (Salomonson, R T C 6, 1) Crystalline Boiling with NH_4Aq and $BaCl_2$ ppts $C_{11}H_7BaNO_7$ Phenylhydrazine at 130° forms $C_{11}H_7NO_7(N_2HPh)$ [132°] — $Cu(C_{11}H_7NO_7)_2$ — PbA'_2 needles (from hot water)

Methyl ether $MeC_{11}H_7NO_7$ Oil

p-Nitro- β -oxy-phenyl-pyrotartaric acid. Lactones [163°] (S), [155°] (Erdmann, B 18, 2742) Formed in like manner from *p*-nitro-benzoic aldehyde. Boiling with NH_4Aq and $BaCl_2$ ppts. $BaC_{11}H_7NO_7$ — CuA'_2 — AgA'_2

NITRO-*p*-OXY-PHENYL-QUINOLINE

$C_{13}H_9N_2O_2$ [151°] Formed in small quantities when *p*-amido (Py 8) phenyl-quinoline is heated with KNO_3 (Weidel, M 8, 138) Yellow plates, m sol alcohol

m Nitro-(B 2) oxy (Py 1) phenyl-quinoline

Methyl derivative $MeO \begin{smallmatrix} C \\ \diagup \diagdown \\ CH \\ \diagdown \diagup \end{smallmatrix} CHC \begin{smallmatrix} C \\ \diagup \diagdown \\ CH \\ \diagdown \diagup \end{smallmatrix} (C_6H_4NO_2) \begin{smallmatrix} CH \\ \diagup \diagdown \\ CH \\ \diagdown \diagup \end{smallmatrix} [130°]$ Formed

by heating *m* nitro cinnamic aldehyde with *p* anisidine and $HClAq$ (Miller & Kinkeln, B 20, 1919) Needles (from benzene), sl sol alcohol. (Py 1,4,2)-Nitro oxy phenyl isouquinoline

$C_{13}H_{11}N_2O_2$, te $C_6H_4 \begin{smallmatrix} C(NO_2) \\ \diagup \diagdown \\ C(OH) \end{smallmatrix} CPh$ [c 245°].

Formed by passing nitrous acid gas through a solution of oxy phenyl isouquinoline in HOAc (Gabriel, B 19, 831) Small yellow crystals

Methyl derivative $C_{13}H_{11}MeN_2O_2$ [169°]

DI NITRO DI OXY-DI-PHENYL SULPHONE

$C_{12}H_8N_2SO_4$, te $SO_2 \{C_6H_4(NO_2)(OH)\}_2$ Formed by nitrating di oxy di phenyl sulphone (Glutz, A 147, 59) Scales, insol water, sol alkalis Yields $C_6H_4Na_2N_2SO_4$ and $C_6H_4Ag_2N_2SO_4$, a dimethyl derivative $C_6H_4Me_2N_2SO_4$ [215°], a diethyl derivative [192°], and a diisomethyl derivative [151°] With aniline it forms the compound $C_{12}H_8(NHPh)(NO_2)_2SO_4$ crystallising from aniline in red prisms

Tetra nitro di-oxy-di phenyl sulphone

$SO_2 \{C_6H_4(NO_2)_2(OH)\}_2$ [253°] Formed by further nitration of the preceding body (Annahme, B 11, 1668) Long yellow needles — K_2A' — NaA' octahedra

TETRA NITRO DI OXY DIPHENYL DISULPHONIC ACID $C_{12}H_8(NO_2)_4(OH)_2SO_4H_2$ Formed by nitration (Limpriht, A 261, 336) — NaA' aq K_2A' yellow crystals

m NITRO *p* OXY-DI-PHENYL THIO-UREA

[3 1] $C_6H_4(NO_2)NHCSNH C_6H_4(OH)$ [1 4] [152°] Formed from *m* nitro phenyl-thiocarbamide and *p* amido phenol (Staudemann, B 16, 2335) Needles, sl sol ether

DI NITRO OXY PHENYL UREA $C_6H_4N_2O_3$, te

$C_6H_4(NO_2)_2(OH)NHCO NH_2$ Uranidic di-nitro phenol Formed by heating urea with di nitro amido phenol (picramic acid) (Griess, J pr [2] 5, 1) Plates (from water) — AgA' pp

NITRO DI-OXY-PHTHALIC ACID Methyl derivative $CH(OMe)(OH)(NO_2)(CO_2H)_2$ [4 3 6 2 1] Normethylnitrohemipic acid [220°] Formed by boiling its imide with KOH (Elbel, B 19, 2310) White needles, v sol water

Imide $C_6H(OMe)(OH)(NO_2) \begin{smallmatrix} C(NH) \\ \diagup \diagdown \\ CO \end{smallmatrix} O$

[252°] Formed by boiling the oxim $C_6H(OMe)(OH)(NO_2)(CO_2H)CHNOH$ with HOAc Yellow needles, sol hot water and alkalis

Dimethyl derivative v Nitro HEMIPIC ACID

Di-nitro oxy phthalic acid

$C_6H(NO_2)_2(OH)(CO_2H)_2$ Juglonic acid Formed by oxidation of juglone or its acetyl derivative by boiling with HNO_3

Salts — $A'(NH_4)_2$ reddish yellow tables or thin needles, v sol water — $A'HK$ — $A'Ba'$ yellow tables (Berntsen & Semper, B 18, 210)

NITRO-OXY-ISOPROPYL-BENZOIC ACID

$CM_2(OH)C_6H_4(NO_2)CO_2H$ [4 8 1]. [191°] Formed by oxidising nitro-cuminic acid or nitro-cuminol with $KMnO_4$ (Widman, B 15, 2549; 16,

2567, 2., 2292) Long needles (from water) — $\text{NH}_4\text{A}'2\text{aq} - \text{CaA}' - \text{BaA}', 6\text{aq} - \text{PbA}', 5\text{aq} - \text{CuA}', 1\frac{1}{2}\text{aq} - \text{AgA}'\frac{1}{2}\text{aq}$ crystals, m sol hot Aq

Acetyl derivative [138°]

Ethylether EtA' [96°]

Nitro oxy isopropyl benzoic acid

$\text{CMe}_2(\text{OH})\text{C}_6\text{H}_4(\text{NO}_2)\text{CO}_2\text{H}$ [4 2 1] [168°]

Formed by oxidising nitro cymene or the acid $\text{C}_6\text{H}_4\text{Pr}(\text{NO}_2)\text{C}_6\text{H}_4\text{CO}_2\text{H}$ with alkaline KMnO_4 (Widman, *B* 19, 270, Söderbaum, *B* 21, 2128) Tables (from ether), m sol hot water

Isomeride *v* Nitro oxy-cuminic acid

Di nitro-oxy propyl benzoic acid *Lactone*

$\text{C}_6\text{H}_4\langle\frac{\text{C}(\text{NO}_2)(\text{CHMeNO}_2)}{\text{CO}}\rangle$. [90°] Formed

from ethylidene phthalide and NO_2 (Gabriel, *B* 19, 838) Colourless needles (from alcohol)

NITRO- β -OXY-*p*-ISOPROPYL-PHENYL-PROPIONIC ACID

[4 2 1] $\text{C}_6\text{H}_4\text{Pr}(\text{NO}_2)\text{CH}(\text{OH})\text{CH}_2\text{CO}_2\text{H}$ [120°] Formed by boiling $\text{C}_6\text{H}_4\text{Pr}(\text{NO}_2)\text{CHBrCH}_2\text{CO}_2\text{H}$ with aqueous Na_2CO_3 (Einhorn & Hess, *B* 17, 2024) Silvery plates

Amide $\text{C}_{11}\text{H}_{14}\text{N}_2\text{O}_4$ [150°]

Anhydride $\text{C}_6\text{H}_4\text{Pr}(\text{NO}_2)\text{CH}\langle\frac{\text{CH}_2}{\text{O}}\rangle\text{CO}$

[73°] Formed by the action of cold aqueous Na_2CO_3 on bromo nitro cumyl propionic acid Crystals, v sol alcohol

NITRO DI OXY PYRIMIDINE $\text{C}_4\text{H}_2\text{N}_4\text{O}_4$ *tc*

$\text{CO}\langle\frac{\text{NH CH}}{\text{NH CO}}\rangle\text{CNO}_2$. Nitro uracil Formed

by heating the K salt of its carboxylic acid at 130° (Behrend, *A* 229, 85, 240, 8) Yellow needles, which explode on heating Yields isobarbituric acid on reduction Urea forms crystalline $\text{C}_4\text{H}_2\text{N}_4\text{O}_3$ Guanidine gives a similar salt $\text{C}_4\text{H}_2\text{N}_4\text{O}_3\text{aq} - \text{KA}'\text{aq}$ prisms, sl sol water — $\text{CaA}', 6\text{aq} - \text{BaA}', 5\text{aq} - \text{ZnA}', 3\frac{1}{2}\text{aq} - \text{CuA}', 7\text{CuO}$

Nitro-di oxy pyrimidine carboxylic acid

$\text{C}_4\text{H}_2\text{N}_4\text{O}_4\text{tc}$ $\text{CO}\langle\frac{\text{NH C}(\text{CO}_2\text{H})}{\text{NH CO}}\rangle\text{CNO}_2$. Formed

by warming di oxy methyl pyrimidine (methyl uracil) with H_2SO_4 and HNO_3 at 80° (Behrend, *A* 229, 32, 240, 4, Köhler, *A* 236, 32) Yellow crystals (containing 2aq) — $\text{KHA}'\text{aq}$ plates, sl sol water — $\text{BaA}'\frac{1}{2}\text{aq} - \text{AgA}'\text{aq} - \text{PbA}'\frac{1}{2}\text{aq}$

Ethyl ether EtHA' [250°] Prisms

(a) **NITRO (Py 3)-OXY-QUINOLINE**

$\text{C}_8\text{H}_5\text{N}_3\text{O}_4$, *a'*-Nitro-carbostyryl Formed by heating *a'*-nitro *o* amido cinnamic acid with HClAq at 150° (Friedländer & Lazarus, *A* 229, 243) Needles (from alcohol) Does not melt below 220°

(β) Nitro (Py 3)-oxy quinoline [260°]

Formed in like manner from β' nitro amido-cinnamic acid (F & L) Needles (from HOAc)

(γ) Nitro-(Py 3) oxy quinoline [280°]

Formed by nitrating carbostyryl (F. & L) Needles (from HOAc)

Methyl derivative [181°]. From the silver salt and MeI (Feer & Königs, *E* 18, 2398)

(B 4)-Nitro-(Py 3)-oxy quinoline

$\text{CH CH} - \text{C OH CH}$ [168°]. Formed from

$\text{CH C}(\text{NO}_2)\text{O} - \text{N COH}$ Formed from di-methyl *o*-nitro-coumarinate and alcoholic NH_3 , the resulting (8, 2, 1)-nitro amido-cinnamic amide being heated with HClAq at 140° (Miller

& Kinkelin, *B* 22, 1711) Prisms, v sol hot water

(B 4, 2) Nitro-oxy quinoline

$\text{C}(\text{OH})\text{CH}\langle\frac{\text{C OH CH}}{\text{C}(\text{NO}_2)\text{O N CH}}\rangle$ [186°] Formed by nitra

ting (B 2) oxy-quinoline (Skraup, *M* 3, 551) and by the action of nitric acid on (B 2)-oxy-quinoline carboxylic acid (Schmidt & Altschul, *B* 20, 2697, 21, 2255) and on nitroso oxy-quinoline (Mathéus, *B* 21, 1642, 1886) Yellow needles — $\text{B'HNO}_3\text{aq}$ orange prisms, v sol hot alcohol

Nitro-(B 4) oxy-quinoline [173°] Formed

by heating its carboxylic acid with glycerin at 200° (Schmitt & Engelmann, *B* 20, 2693) and by the action of HNO_3 (S G 1 38) on nitroso-(B 4)-oxy-quinoline (Von Kostanecki, *B* 24, 154) Needles, sl sol alcohol

Nitro-(B 2) oxy quinoline [255°] Formed

by nitrating *m* oxy-quinoline (Skraup, *M* 3, 564) Yellow plates, decomposed by fusion

Nitro-oxy quinoline Formed by the action

of HNO_3 on a syrupy acid obtained by oxidation of cinchonine (Weidel & Hazura, *M* 3, 773) Crystalline powder, melting far above 300° — $\text{B'H}_2\text{PtCl}_6$ monoclinic prisms

(B 1, 3) Di nitro (B 4) oxy-quinoline

$\text{CH C}(\text{NO}_2) - \text{C CH CH}$ [276°] Formed by the action of HNO_3 on *o* oxy quinoline carb oxylc acid, and on *ana* nitroso *o* oxy quinoline (Schmitt & Engelmann, *B* 20, 2692, Kostanecki, *B* 24, 155, cf Bedall & Fischer, *B* 14, 1368) Plates

NITRO-(B 4) OXY-QUINOLINE CARBOXYLIC ACID $\text{C}_8\text{H}_5(\text{NO}_2)\text{N}(\text{OH})(\text{CO}_2\text{H})$ Formed

by boiling the nitrate of *o* oxy quinoline carb oxylc acid with HOAc (Schmitt & Engelmann, *B* 20, 2693) Needles, sl sol HOAc

NITRO-OXY-QUINONE *Carbonyl deri-*

ivative $\text{C}_8\text{H}_4\text{O}_2(\text{NO}_2)_2\text{CO}$ [260°] Formed by oxidation of nitro amido phenyl carbonate by chromic acid mixture (Lowenberg, *C* C 1886, 390) Pale brown needles

Nitro-di-oxy quinone

$\text{CO}\langle\frac{\text{CH C}(\text{OH})}{\text{C}(\text{OH})\text{C}(\text{NO}_2)}\rangle\text{CO}$ Formed by warming nitro di imido resorcin with dilute (10 p c) NaOHAq (Nietzki & Schmidt, *B* 22, 1659) Golden needles, m sol water — KA' orange needles

Di nitro di-oxy quinone

$\text{CO}\langle\frac{\text{C}(\text{NO}_2)\text{C}(\text{OH})}{\text{C}(\text{OH})\text{C}(\text{NO}_2)}\rangle\text{CO}$ *Nitrilic acid*.

Formation — 1 By the action of nitrous acid

on hydroquinone (Nietzki, *B* 10, 2147) — 2 By the action of a mixture of fuming HNO_3 and conc H_2SO_4 on di acetyl hydroquinone below — 5°, the yield in this case being 65 p c (Nietzki, *B* 16, 2092, 18, 499) — 3 By adding dimethoxyhydroquinone to a cooled mixture of HNO_3 (3 pts) and HOAc (6 pts) (Nietzki, *A* 215, 142) — 4 By boiling di-nitro-di-amido-quinone with dilute potash (Nietzki, *B* 20, 2116) — 5 By the action of fuming HNO_3 on $\text{CO}_2\text{H C}\langle\frac{\text{CO C}(\text{OH})}{\text{C}(\text{OH})\text{CO}}\rangle\text{CO}_2\text{H}$ (Hantzsch, *B* 19, 2398, cf Loewy, *B* 19, 2385)

Preparation — By slowly adding a hot saturated alcoholic solution of chloranil (4 pts) to a concentrated aqueous solution of sodium nitrate (10 pts), heated to 80°–90°, a yellow crystalline

pp of the sodium nitranilate soon separates, and a small quantity which remains in solution is precipitated by NaOH, the whole is recrystallised from hot water (Nef, *B* 20, 2027)

Properties—Golden tables, v sol water and alcohol, insol ether. When anhydrous it explodes at 170° without previous fusion. FeCl₃ gives a green crystalline pp. Chlorine forms oxalic acid and chloropierin (Levy, *A* 249, 66). On reduction it yields tetra oxy di amido benzene which gives p phenylene diamine on distillation with zinc-dust (Nietzki, *B* 19, 2727). Hydroxylamine hydrochloride forms an explosive salt C₆O₄(NO₂)₂(NH₂O)₂ (Nef, *Am* 11, 17).

Salts—Na₂A'' dichroic monoclinic crystals, *a b c* = 946.1 985, β = 87° 51'—K₂A'' yellow needles (from hot water)—(NH₄)₂A''—BaA'' plates, insol water.

Di-nitro-di-oxy-quinone A product of the action of nitrous acid on protocatechuic acid dissolved in ether (Gruber, *B* 12, 519). Greenish-yellow needles, v sol water—Na₂A'' 2aq explodes when heated.

NITRO-OXY STYRENE *Methyl derivative* [1 2 4] C₆H₃(OMe)(NO₂)CH CH₃ [89°] Formed, together with the di nitro derivative C₆H₃(OMe)(NO₂)₂C₆H₅ [163°], by the action of conc HNO₃ on [4 1] C₆H₄(OMe)CH CH CO₂H (Einhorn a Grabfeld, *A* 243, 366). Crystals, volatile with steam. Yields a dibromide [79°]. Forms, on oxidation, nitranisic acid [187°].

NITRO-OXY-STYRYL METHYL KETONE *Methyl derivative* C₆H₃NO₂ [4 3 1] C₆H₃(OMe)(NO₂)CH CH CO CH₃ [159°] Formed by nitration of the ketone, and also by condensation of C₆H₃(OMe)(NO₂)CHO with acetone (Einhorn a Grabfeld, *A* 243, 364). Yellow needles (from water), sol ether.

NITRO-OXY SULPHO BENZOIC ACID C₆H₃NSO₃ *ie* C₆H₃(NO₂)₂(OH)(SO₃H)(CO₂H) Formed from nitro-o oxy benzoic acid and fuming H₂SO₄ (Mandt, *B* 10, 1701)—Ba₂A'' 12aq hair-like needles.

NITRO OXY TOLUENE *v* Nitro Cresol

Nitro-di oxy-toluene *v* Nitro-ORCIN

Nitro-tetra oxy toluene C₆Me(NO₂)(OH)₄ [157°] Formed by the action of HCl and SnCl₄ on nitro-di oxy-toluquinone (Kehrmanna Brasch, *J pr* [2] 89, 382). Black needles, forming a violet powder. Its solution forms HCl, oxalic acid, and other products on boiling.

Di nitro di-oxy-toluene

C₆HMe(NO₂)₂(OH)₂ [1 3 5 2 4] *Di-nitro cres orcin* [90°] Formed from cresorcin and HNO₃ (Von Kostanecki, *B* 20, 3136). Needles, m sol cold water.

NITRO-OXY-*o* TOLUIC ACID C₆H₃NO₂ *ie* C₆H₂Me(OH)(NO₂)CO₂H [1 4 x 2] [172°] Formed by nitration of oxy *o* toluic acid (Kostanecki a Niemientowski, *B* 18, 254). Needles, sol hot Aq.

Nitro oxy-*m*-toluic acid

C₆H₂Me(OH)(NO₂)CO₂H [3 4 5 1]? [87°] Formed by heating (4,8,1)-oxy toluic acid with conc HNO₃ (Mahon, *Am* 4, 186). Yellow needles, sl sol water. Its salts explode when heated—CaA', 4aq—BaA', 4aq orange needles; crimson when anhydrous.

Nitro-oxy-*p*-toluic acid

C₆H₂Me(OH)(NO₂)CO₂H [4 3 x 1] [188°] Formed by the action of nitrous acid on (3,4,1) amido-

toluic acid (Ahrens, *Z* 1869, 105). Golden needles—BaA', 2aq scarlet, sl sol alcohol.

Nitro-oxy toluic acid *Methyl derivative* C₆H₂Me(OMe)(NO₂)CO₂H [175°] Formed from the methyl ether of thymol and dilute HNO₃ (Paterno a Canzoneri, *G* 9, 445). Slender needles, v sol alcohol—BaA', 2aq straw coloured crystals.

Ethyl derivative [162°] Formed, in like manner, from the ethyl ether of thymol. Long slender needles.

Nitro *o* oxy *o* toluic acid

C₆H₃(NO₂)(CH₂OH)CO₂H [4 2 1] [129°] Formed by dissolving nitro phthalide in aqueous KOH (Hoernig, *B* 18, 3451). Minute needles—AgA'.

Tri nitro oxy *m*-toluic acid

C₆HMe(OH)(NO₂)₃CO₂H [1 3 2 4 6 5] *Nitrococcic acid* [170°-180°] Formed by the action of boiling nitric acid (S G 1 37) on cochineal (De la Rue, *A* 64, 23, Liebermann a Dorp, *A* 163, 100) and on (5,3,1) oxy toluic acid (Kostanecki a Niemientowski, *B* 18, 250). Colourless plates (containing aq). On boiling with moist Ag₂O it yields silver tri nitro cresol and CO₂—(NH₄)₂A' 1/2 aq—BaA', aq—AgA' needles.

NITRO OXY *m* TOLUIC ALDEHYDE

C₆H₃NO₂ *ie* C₆H₂Me(NO₂)(OH)CHO [5 3 2 1] [141°] Formed by warming oxy toluic aldehyde with HNO₃ (Schotten, *B* 11, 788). Yellow needles, sl sol hot water.

Nitro oxy *m* toluic aldehyde

C₆H₂Me(NO₂)(OH)CHO [5 3 4 1] [152°] Formed by nitrating (4,3,1) oxy toluic aldehyde (S) Needles, sl sol hot water.

NITRO DI OXY TOLUQUINONE

C₆Me(NO₂)(OH)₂O₂ [2 5 3 6 4 1] *Tolunitranilic acid* [180°] Formed by the action of a dilute alcoholic solution of KNO₃ on tri chloro tolu quinone (Kehrmann, *B* 21 1779, *J pr* [2] 39, 377). Golden needles (containing 2aq). Its aqueous solution decomposes on boiling forming HCl, oxalic acid, and CO₂—K₂A' 3aq yellowish red prisms.

DI NITRO DI OXY DITOLYL C₁₁H₇N₂O₄

[273°] (G), [270°] (D). Formed by boiling tetrato ditolyl sulphate with HNO₃ (Gerber, *B* 21, 750), by the action of nascent nitrous acid on di amido ditolyl (the yield being quantitative), and by heating di oxy ditolyl di carboxylic acid with HNO₃ (Deninger, *J pr* [2] 40, 300, *B* 21, 1639). Yellow needles (from toluene or pyridine).

NITRO - OXY - TOLYLENE ETHERYL

AMIDINE C₆H₃N₂O₄ [256°] Formed by the action of alcoholic ammonium sulphide on the acetyl derivative of di nitro *p* toluindine (Bankievitch, *B* 21, 2404). Lustrous green needles, not affected by HClAq at 200°.

NITRO-PENTANE C₅H₇NO₂ *ie*

Pr CH₂CH₂NO₂ (150°-160°) Formed from isoamyl iodide and AgNO₃ (V Meyer, *B* 5, 203, *A* 171, 43, 175, 135).

Di nitro pentane C₅H₃CH(NO₂)₂ Formed from di amyl ketone and HNO₃ (Chancel, *C R* 94, 399). Heavy oil. Forms *n*-valeric acid on reduction—KC₆H₇N₂O₄—AgA'.

***o* NITRO-PHENOL** C₆H₃NO₂ *ie*

C₆H₂(NO₂)OH [1 2] Mol w 189 [45°] (214°) S V S 107 64 (Schiff).

Formation—1 Together with the *p* isomeride by nitration of phenol (Hofmann, *A* 103, 847; Fritzsche, *A* 110, 150, *J pr* 78, 293, Gold

stein, *B* 11, 1943)—2 By heating *o*-bromonitro benzene [38°] or *o*-chloronitro benzene with aqueous potash in sealed tubes (Zincke & Walker, *B* 5, 117, Engelhardt & Latschinoff, *B* 3, 423)—3 By boiling *o*-di-nitrobenzene with NaOH aq (Laubenheimer, *B* 9, 1828)—4 Together with *p*-nitro phenol by boiling diazobenzene sulphate with nitric acid (Nörling & Wild, *B* 18, 1338)—5 Together with *p*-nitro phenol by adding liquid N_2O to cooled CS_2 containing C_6H_5ONa in suspension (Schall, *B* 16, 1901)—6 By heating diazobenzene nitrate with dry toluene, nitrogen being given off (Remsen & Orndorff, *Am* 9, 390)—7 By the action of $NaNO_2$ and H_2SO_4 on aniline (Deminger, *J pr* [2] 40, 298)

Preparation—Phenol (1 pt) is slowly added to a mixture of HNO_3 (1 pt of SG 1.38) and water (6 pts), cooled to 0°, the product is neutralised with Na_2CO_3 and distilled with water (Neumann, *B* 18, 3320)

Properties—Light yellow prisms or needles, v sol alcohol and ether, sl sol cold water

Reactions—1 Reduced by tin and HCl to *o*-amido phenol—2 Aqueous NH_3 (35 p.c.) at 160°–200° yields *o*-nitro aniline (Merz & Ris, *B* 19, 1749)—3 Phenyl hydrazine dissolved in xylene at 100° produces *o*-amido phenol, benzene, NH_3 , and nitrogen (Barr, *B* 20, 1497)

Salts—The colour of the salts has been examined by Carnelley & Alexander (*C J Proc* 4, 64)— NH_4A' scarlet plates— KA' aq orange red crystals (F)— KA' aq (Post, *B* 8, 1552) S 16 at 6°, 21 at 15°— NaA' scarlet plates, v e sol water— BaA' , S 9 at 6°— SrA' , 3 aq— CaA' , 4 aq plates— CaA' , aq orange needles— AgA' orange red pp S 14 at 15°

Acetyl derivative $C_6H_4(NO_2)OAc$ [41°] (253°) Long colourless needles or prisms, v sol alcohol (Butcher, *B* 16, 1933)

Benzoyl derivative $C_6H_4(NO_2)(OBz)$ [59°] Formed from *o*-nitro phenol and $BzCl$ (Hübner, *A* 210, 386, Schiaparelli, *G* 11, 73, Neumann, *B* 18, 3320, 19, 2018) Prisms or needles Yields on nitration the compound $C_6H_4(NO_2)OCO_2C_6H_4(NO_2)$ [13] [126°], crystallising in needles

Methyl ether $C_6H_4(NO_2)(OMe)$ *o*-Nitro anisole [9°] (277°) at 735 mm Formed, together with the *p* isomeride, by nitration of anisole Formed also by methylation of *o*-nitro phenol (Brunck, *Z* 1867, 204, Mühlhauser, *A* 207, 237) and by boiling *o*-chloronitro benzene with $NaOMe$ in $HOme$ (De Bruyn, *R T C* 9, 200) Oil Converted into *o*-nitro aniline by heating with ammonia (Salkowski, *A* 174, 278)

Ethyl ether $C_6H_4(NO_2)(OEt)$ *o*-Nitrophenetol (267°) Formed by ethylation of *o*-nitro phenol (Groll, *J pr* [2] 12, 207, Seidel, *J pr* [2] 42, 448) and by heating $C_6H_4Cl(NO_2)$ with $NaOEt$ (De Bruyn) Oil When distilled with alcoholic potash it yields $C_6H_4(NH_2)(OEt)$ and no azo compound, but when reduced in alcoholic solution by sodium-amalgam it forms $N_2(C_6H_4OEt)_2$ and $N_2O(C_6H_4OEt)_2$ (Schmitt & Möhlau, *J pr* [2] 18, 200)

Bromo-ethyl ether $C_6H_4(NO_2)OC_2H_4Br$ [44°] Formed from $C_6H_4(NO_2)(ONa)$ and ethylene bromide (Weddige, *J pr* [2] 24, 246). Yellow prisms (from alcohol)

Reactions—1. With an alcoholic solution of

NH_3 it yields $C_6H_4(NO_2)OC_2H_4NH_2$ [78°] and $NH(C_6H_4OC_2H_4NO_2)_2$ [192°]—2 Heated with potassium salicylic ether $C_6H_4(OK)CO_2Et$ in alcoholic solution it yields two products: (a) an ether $C_6H_4(NO_2)OC_2H_4OC_2H_4CO_2Et$ [c 100°] saponified by HCl yielding the corresponding acid [145°], which may be reduced to an amido acid [110°] whose hydrochloride melts at 177°, and (b) a compound of the formula $C_6H_4(NO_2)OC_2H_4OCO_2C_6H_4OH$ [106°] which gives an acetyl derivative [80°] (Wagner, *J pr* [2] 27, 212)—3 Heated with [41] $C_6H_4(OK)CO_2Et$ in alcoholic solution it forms *o*-nitro phenoxy ethyl *p*-oxy benzoic ether [103°] which is saponified by HCl at 140°, forming an acid $C_6H_4(NO_2)OC_2H_4OC_6H_4CO_2H$ [c 206°] The corresponding amido acid melts at 185° (Wagner)—4 Potassium benzoate at 140° yields $C_6H_4(NO_2)OC_2H_4OBz$ [77°]

Amido ethyl ether
 $C_6H_4(NO_2)OC_2H_4NH_2$ [73°] Formed as above Small plates (from water) Yields a benzoyl derivative [95°] and a dibenzoyl derivative $C_6H_4(NO_2)OC_2H_4NBz_2$ [122°] The benzoyl derivative reduced by tin and hydrochloric acid yields $C_6H_4<NH CPh>N$ [151°] (Weddige, *J pr* [2] 24, 250)

Ethylene ether $C_6H_4(OC_2H_4NO_2)_2$ [163°] Formed from $C_6H_4(NO)ONa$ and C_2H_5Br

Isobutyl ether (275°–280°) SG 22 1.136 (Riess, *B* 3, 780)

Benzyl ether $C_6H_4CH_2OC_6H_4(NO_2)$ [29°] From the K salt and CH_2Cl (Kumpf, *A* 224, 121)

***p*-Nitro benzyl ether**
 $C_6H_4(NO_2)CH_2OC_6H_4NO_2$ [129°] Needles (K)

Phenacetyl ether
 $C_6H_4(NO_2)OCOCH_3$ [118°] (194°) Formed from the K salt and ω -bromo acetophenone (Lellmann & Donner, *B* 23, 172) Needles, m sol alcohol $ZnCl_2$ and HCl at 100° convert it into $C_6H_4<OCH N CPh>$ [103°] crystallising in needles, and forming the salts $B'H.PtCl_4$ and $B'HuCl_4$

***m*-Nitro phenol** $C_6H_4(NO)OH$ [13] [96°] (194° at 70 mm) Obtained from *m*-nitroaniline by the diazo reaction (Fittig & Bantlin, *B* 7, 179, 11, 2099, Henriques, *A* 215, 323, Wagner, *J pr* [2] 32, 70) Yellow crystals, sol. hot water, not volatile with steam— KA' 2 aq: orange needles S 12 at 6° (Posta Mehrrens, *B* 8, 1552)— BaA' 2 aq S 17 at 6°— PbA' (OH) S 013 at 15°— AgA' brownish red pp

Benzoyl derivative $C_6H_4(NO_2)OBz$ [95°] Pale yellow crystals (Neumann, *B* 19, 2979) Nitric acid (SG 1.48) converts it into the crystalline *m*-nitro benzoyl derivative $C_6H_4(NO_2)OCO_2C_6H_4(NO_2)$ [129°]

Methyl ether MeA' [38°] (254°). Needles, volatile with steam (Bantlin)

Ethyl ether EtA' [34°] (264°) (Bantlin, Wagner, *J pr* [2] 32, 71)

Bromo-ethyl ether C_6H_4BrA' [39°]. Formed, with the following, from the K salt and ethylene bromide (Weddige, *J pr* [2] 24, 255)

Ethylene ether C_6H_4A' [139°]

***p*-Nitro-phenol** $C_6H_4(NO_2)OH$ [114°]. S.V.S 108.28 (Schiff)

Formation—1 Together with the *o* isomer-

ide, by the nitration of phenol, especially at low temperatures (Fritzsche, *J pr* 75, 257, Goldstein, *J R*, 10, 353)—2 By heating *p* chloro-nitro-benzene [85°] or bromo nitro-benzene [125°] with aqueous potash at 130° (Engelhardt & Latschnoff, *B* 3, 423, Richter, *B* 4, 460)—3 From acetanilide by nitrating, and heating the [4] $\text{C}_6\text{H}_4(\text{NO}_2)(\text{NHAc})$ with conc NaOH aq (Wagner, *B* 7, 76)—4 From *p*-nitro-aniline by the diazo-reaction (Fittig, *B* 7, 280)—5 By dry distillation of nitro *o* oxy-benzoic acid (Schmidt & Cook, *K* 3, 41)—6 By oxidation of nitroso phenol (Bayer, *B* 7, 965)—7 Together with the *o* isomeride, by adding liquid N_2O , to cooled CS_2 containing $\text{C}_6\text{H}_5\text{ONa}$ in suspension (Schall, *B* 16, 1901)—8 By the action of NaNO_2 (in excess) and H_2SO_4 on aniline, *o* nitro phenol being also formed (Deming, *J pr* [2] 40, 298)

Preparation—The mixture of *o*- and *p* nitro-phenols is distilled with steam. The residue is dissolved in hot water and mixed with excess of Na_2CO_3 when $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{ONa}$ crystallises out on cooling (Salkowski, *A* 174, 280)

Properties—Slender colourless needles (from water Dimorphous (Lehmann, *Z K* 1, 45) V sol alcohol Not volatile with steam Sol hot HCl aq (Kollrepp, *A* 234, 1)

Reactions—1 Distillation with PCl_5 yields *p* chloro-nitro benzene [85°]—2 Aqueous ammonia at 160° to 200° forms *p* nitro aniline (Merz & Ris, *B* 19, 1749)—3 Phenyl hydrazine at 100° produces *p*-amido phenol, benzene, NH_3 , and nitrogen (Barr, *B* 20, 1499)

Salts—The colour of the salts has been examined by Carnelley (*G J Proc* 4, 64)— NaA' aq yellow tables— NaA' 2aq Heat of neutralisation 12,840 (Alexejeff & Werner, *Bi* [3] 2, 718)— NaHA' 2aq orange red prisms (from water)— KA' 2aq S 4.5 at 6° (Post & Mehrrens, *B* 8, 1552)— KHA' 2aq— BAa' 8aq monochlonic prisms S 1 at 6°, 13 at 15°— BaHA' 4aq— SrA' 7aq yellow needles— CaA' 4aq— CaHA' 8aq— MgA' 8aq— $\text{PbO}_2\text{A}'$ — $\text{Pb}_2\text{HA}'$ — AgA' aq scarlet pp, changing to orange prisms— AgA' 2aq S 3 at 15°— AgHA' aq yellowish green laminae— $\text{Ag}_2\text{HA}'$ purple needles

Benzoyl derivative $\text{C}_6\text{H}_4(\text{NO}_2)\text{OBz}$ [142.5°] Colourless efflorescent needles (from alcohol) (Schiaparelli, *G* 11, 73, Neumann, *B* 19, 2020) On treatment with HNO_3 (S G 148) it forms [4] $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OCO C}_6\text{H}_4\text{NO}_2$ [13] [185.5°]

Methyl ether MeA' . [51°] (259°) Formed by methylation of *p* nitro phenol (Brunck, *Z* [2] 3, 202, Willgerodt & Ferko, *J pr* [2] 83, 152, Skraup, *M* 6, 761) Formed also by heating *p* chloro nitro benzene with KOH dissolved in MeOH (Willgerodt, *B* 14, 2632, 15, 1004) Prisms

Ethyl ether EtA' [58°] (283°) (Andreas, *J pr* [2] 21, 831) Formed from the Ag salt and EtH (Fritzsche) and also from $\text{C}_2\text{H}_5\text{OEt}$ and fuming HNO_3 (Hallock, *Am* 1, 271) Obtained also by boiling *p*-chloro-nitro benzene with KOH and dilute (60 *p c*) alcohol (Willgerodt, *B* 15, 1002) Prepared by heating $\text{C}_6\text{H}_4(\text{NO}_2)(\text{OK})$ with KEtSO_4 in alcoholic solution for 8 hours (Willgerodt & Ferko, *J pr* [2] 83, 158). Prisms Sodium amalgam rapidly re-

duces it, in alcoholic solution, to $\text{N}_2(\text{C}_6\text{H}_4\text{OEt})$. (Schmitt & Möhlau, *J pr* [2] 18, 199)

Ethylene ether $\text{C}_2\text{H}_4\text{A}'$. [143°] Formed, at the same time as the bromo ethyl ether, by heating the Na compound with ethylene bromide at 140° (Weddige, *J pr* [2] 21, 127, 24, 254)

Bromo-ethyl-ether $\text{CH}_3\text{Br C}_6\text{H}_4\text{A}'$ [63°] Yellowish plates, insol water, v sol alcohol

Reactions—1 Heated with potassium *p* oxy benzoic ether [4] $\text{C}_6\text{H}_4(\text{OK})\text{CO}_2\text{Et}$ it forms $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OCO C}_6\text{H}_4\text{CO}_2\text{Et}$ [181°], which when saponified by HCl aq at 130° yields the corresponding acid [218°] (Wagner, *J pr* [2] 27, 224)—2 Alcoholic [2] $\text{C}_6\text{H}_4(\text{OK})\text{CO}_2\text{Et}$ forms $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OCO C}_6\text{H}_4\text{OH}$ [181°] and the ether $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OCO C}_6\text{H}_4\text{CO}_2\text{Et}$ [c 81°] whence HCl liberates the acid [132°]

Amido ethyl ether $\text{C}_6\text{H}_4(\text{NH}_2)\text{A}'$ [103°] Formed by heating the bromo ethyl ether with alcoholic ammonia Yellow scales (from water)

Isobutyl ether (285°–290°) S G 20 1105

Benzylether $\text{C}_6\text{H}_5\text{CH}_2\text{A}'$ [106°] Prisms. Yields on nitration a compound melting at 163° (Kumpf, *A* 224, 123)

***p* Nitro-benzyl ether** [183°] Needles Fourth and fifth nitro-phenols have been described by Fittica (*J pr* [2] 24, 5, *B* 13, 711) but their existence has not been confirmed by other chemists (Natanson, *B* 13, 415)

(a)-**Di nitro-phenol** $\text{C}_6\text{H}_3(\text{NO}_2)_2(\text{OH})$ [4 2 1] [114°] S 4.8 at 100°, 5 at 18°, 0.14 at 0° (Gruner, *J pr* 102, 222)

Formation—1 By nitration of phenol, *o* nitro phenol, and *p* nitro phenol (Laurent, *A Ch* [3] 3, 212, Körner, *Z* [2] 2, 662, 731)—2 From anisole by nitration and saponification (Cahours, *A Ch* [3] 25, 22)—3 From di nitro amido phenol (picramic acid) by elimination of NH_2 (Griess, *A* 113, 210)—4 From chloro di-nitro benzene [50°] and bromo di nitro benzene [72°] by heating with potash (Clemm, *J pr* [2] 1, 145, Engelhardt & Latschnoff, *B* 3, 97)—5 By boiling *i tri* nitro benzene with aqueous Na_2CO_3 or by heating it with water at 160° (De Bruyn, *R T C* 9, 191)—6 By boiling *i di* nitro aniline with KOH aq for a long time (Willgerodt, *B* 9, 979)—7 From di nitro di ethyl aniline and dilute potash (Van Romburgh, *R T C* 2, 35)

Properties—Yellowish rectangular plates (from water), v sol cold alcohol

Reactions—1 Yields only picric acid on further nitration (Hübner & Schneider)—2 Aqueous KCy at 70° forms potassium meta purpurate $\text{C}_6\text{H}_3\text{KN}_3\text{O}_6$ which separates as brownish red crystals with green lustre (Pfaunder & Oppenheim, *Z* 1866, 470, Sommaruga, *A* 157, 335)

Salts— KA' aq S 1.4 at 6° (Post & Mehrrens, *B* 8, 1554), 1.6 at 7° (Hübner & Schneider, *A* 167, 92)— KA' aq (Romburgh)— NaA' aq— BAa' 7aq golden needles— BaA' 6aq yellow needles— BAa' 5aq orange prisms S 3 at 7°— BaA' 4aq monochlonic crystals (*P a M*)— MgA' 8aq— MgA' 12aq— $\text{Pb}(\text{OH})\text{A}'$ 2aq S .09 at 15°— MnA' 5aq— NiA' 8aq— AgA' aq. S 4 at 15°

Benzoyl derivative $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OBz}$. Plates (from alcohol) (Laurent & Gerhardt, *A* 75, 77).

m-Nitro-benzoyl derivative [161°] Formed by nitration of $2[1]C_6H_4(NO_2)OBz$ or of the *o*-isomeride (Neumann, *B* 18, 3322, 19, 2021) Needles

*Methyl ether MeA' [88°] Formed by boiling anisole or anisic acid with fuming HNO_3 (Cahours, *A* 69, 236) or by heating (4, 2, 1)-chloro di nitro benzene or 1 tri nitro benzene with KOH dissolved in MeOH (Willgerodt, *B* 12, 762, De Bruyn, *R T C* 9, 190) Needles

Ethyl ether EtA' [86°] Formed by the action of HNO_3 on C_6H_5OEt (Cahours, *A* 74, 315) and on $N_2(C_6H_4OEt)[14]$, (Andrem, *J pr* [2] 21, 335), and also in the same way as the methyl ether (W) Needles

Allyl ether C_6H_5A' [47°]

Di-oxy propyl ether $C_6H_4(OH)_2A'$ [o 83°] Formed from $[124]C_6H_4Cl(NO_2)_2$, glycerin, and KOH (W)

Phenyl ether C_6H_5A' [71°] (W)

Benzyl ether $C_6H_5CH_2A'$ [149°]

p-Nitro benzyl ether [201°] (Kumpf)

(*g*) Di nitro phenol $C_6H_3(NO_2)_2(OH)$ [431] [64°] Formed, together with the (*a*) isomeride, by nitrating *o*-nitro phenol (Hubner & Schneider, *A* 167, 89, Salkowski, *A* 174, 270, Körner, *G* 4, 325) Pale yellow needles (from water), sl sol water, v e sol hot alcohol. Somewhat volatile with steam—NaA' 3aq red needles—KA' S 13 at 6°—BaA', aq golden needles S 18 at 7°—MgA', 6aq—Pb₂OA', S 037 at 15°—AgA' S 3 at 6° (Post & Mehrrens, *B* 8, 1552)—AgA' aq red needles

m-Nitro benzoyl derivative [149°] Got from $[31]C_6H_4(NO_2)OBz$ and HNO_3 (Neumann, *B* 19, 2930) Yellow needles, sl sol ether

Methyl ether MeA' [117°] Converted by NH_4Aq at 130° into di nitro aniline [138°]

Ethyl ether EtA' [58°] Needles

Benzyl ether PhCH₂A' [76°] Prisms

Yields $[41]C_6H_4(NO_2)_2CH_2OC_6H_4(NO_2)_2$ [136] [137°] on nitration (Kumpf, *A* 224, 130)

(*γ*) Di nitro phenol $C_6H_3(NO_2)_2(OH)$ [531] [104°] (B), [122°] (De B) Formed, together with the (*g*) and (*e*) isomerides, by nitration of *m*-nitro phenol (Bantlin, *B* 11, 2103, *A* 215, 324) Obtained also by heating its methyl ether with conc HClAq at 180° (De Bruyn, *R T C* 9, 208) Needles, volatile with steam—KA' 2aq—BaA', 3aq—BaA', 2aq red feathery crystals

Methyl ether MeA' [96°] (B), [105°] (De B) (above 360°) Formed by heating *s*-trinitro benzene with NaOMe in HOMe for 24 hours (De Bruyn) Needles

(*g*) Di-nitro phenol $C_6H_3(NO_2)_2(OH)$ [431] [134°] Prepared, together with the (*e*)- and (*γ*)-isomerides, by nitration of *m*-nitro phenol (Bantlin, *B* 11, 2104) Colourless needles, not volatile with steam—BaA', 3aq brown prisms

Methyl ether MeA' [70°] Needles

(*e*) Di-nitro-phenol $C_6H_3N_2O_5$ s.e.

$C_6H_3(NO_2)_2(OH)$ [821] [144°] Prepared, together with the (*γ*) and (*g*) isomerides, by nitration of *m*-nitro phenol (Bantlin, *B* 11, 2104) Yellow needles (from water)—KA' 2aq yellow needles—BaA', brown needles

Methyl ether MeA' [118°] Tables

Tri-nitro-phenol $C_6H_2(NO_2)_3OH$ [6421]

Picric acid Mol. w 229 [122°] S 626 at 5°; 1225 at 20°; 389 at 77° (Marchand, *J*

pr 64, 91) R_∞ 7862 in a 174 p.c. benzene solution (Kanonnikoff, *J pr* [2] 31, 348)

Formation—1 By the action of hot nitric acid on phenol, tri-bromo phenol, *o* and *p*-nitro-phenols, (*α*)- and (*β*)-di nitro phenols, saligenin, salicylic aldehyde, salicin, salicylic acid, phlorizin, indigo, coumarin, aloes, gum benzoïn, balsam of Peru, and from the resin of *Xanthorrhoea hastilis* (Hausmann, *Journ de Phys et de Chimie*, 1788, Welter, *A Ch* 29, 301, Liebig, *P* 13, 191, 14, 466, *A* 9, 80, 39, 350, Dumas, *A Ch* [2] 53, 178, [3] 2, 228, Laurent, *A Ch* [3] 3, 221, *A* 43, 219, Perra, *D P J* 165, 386, Piria, *A* 56, 63, Stenhouse, *A* 57, 88, 66, 243, Carey Lea, *Am S* [2] 26, 279, E Kopp, *A Ch* [3] 13, 233, Delalande, *A* 45, 337, Marchand, *A* 48, 336, 52, 345, Schunck, *A* 39, 6, 65, 234)—2 By boiling (1,2,4,6)-chloro-tri nitro-benzene (picryl chloride) with aqueous Na₂CO₃ (Engelhardt & Latschmoff, *B* 3, 98, Clemm, *J pr* [2] 1, 145)—3 By oxidation of *s*-tri nitro benzene with K₂FeCy₄ in slightly alkaline solution (Hepp, *B* 13, 2346)—4 By heating iodo benzene with silver nitrite at 150° (Geuther, *A* 245, 100)

Preparation—Phenol (1 mol) is dissolved in conc H₂SO₄ and the resulting phenol sulphonic acid treated with HNO₃ (4½ mols of SG 135) (Schmitt & Glutz, *B* 2, 52)

Properties—Light yellow laminae (from water) or trimetric prisms (from ether) May be sublimed if slowly heated, but if a few milligrammes be dropped into a red hot tube violent detonation occurs, with a larger quantity a less violent decomposition occurs (Berthelot, *A Ch* [6] 16, 21) Sl sol water, a 01 p.c. solution being distinctly yellow V sol alcohol and ether Tastes bitter Dyes silk and wool yellow It is poisonous Forms crystalline compounds with aromatic hydrocarbons (Fritzsche, *J pr* 73, 212, *A* 109, 247)

Reactions—1 Bleaching powder yields, on boiling, chloropiericn KCl₂NO₂ and tetra chloro quinone A mixture of KClO₃ and HCl acts in like manner—2 Distillation with aqueous NaOBr forms bromopiericn (Stenhouse, *P M* [4] 8, 363) 3 Ferrous sulphate and lime reduce it to di nitro amido phenol (picramic acid) (Girard, *C R* 36, 421)—4 Tin and HClAq reduce it to tri amido phenol (picramine) (Roussin, *Bl* 1861, 60, Beilstein, *A* 130, 244) Iodide of phosphorus acts in like manner on its aqueous solution (Lautemann, *A* 125, 1)—5 Hot conc KCyAq forms a blood red solution of potassium isopurpurate which crystallises in brownish-red scales with green luster having either the formula $KC_6H_3N_3O_5$ (Hlasiwetz, *A* 110, 289) or $KC_6H_3N_3O_5$ (Baeyer, *J* 1859 458) Ammonium chloride converts this salt into an ammonium salt $NH_4C_6H_3N_3O_5$, greatly resembling murexide—6 PCl₅ yields $C_6H_3Cl(NO)_3$ (Pisani, *C R* 39, 852)

Salts—Explodes when struck or when strongly heated—NH₄A' trimetric prisms (Laurent, *Rev Scient* 9, 26)—LiA' SG 12 1716 slender yellow prisms (Beamer & Clarke, *Am* 1, 153)—NaA' S about 8 at 15° S (alcohol) 125 in the cold (Hager, *Pharm Centr* 22, 225)—KA' Trimetric prisms, *a b c* = 1270 188 (Laurent, *Rev Scient* 10, 26) S 4 at 15°, 7 at 100° S (alcohol) 04 (Hager) Explodes when struck, giving off CO₂, CO, and

nitrogen, with smaller quantities of hydrogen and CH_4 , and leaving a residue of K_2CO_3 , carbon, and K_2CO_3 (Sartana a Vieille, *C R* 93, 61) — BaA'_2 , 5aq monoclinic crystals — BaA'_2 , 6aq S (of BaA'_2) 12 at 17° (Tscheltzoff, *A Ch* [6] 8, 233) — BaA'_2 , 4aq S 6 at 6° (Post a Mehrtens) — CaA'_2 , 5aq S (of CaA'_2) 50 at 20° — SrA'_2 , 5aq yellow crystals (Marchand) S (of CaA'_2) 14 at 20° — MgA'_2 , 8aq S (of MgA'_2) 10 at 22° — MgA'_2 , 3 NaA'_2 , 9aq (Muller, *Z* 1865, 189) — CdA'_2 , 7aq — CdA'_2 , 3 NH_3 (Carey Lea, *Am S* [2] 81, 78) — CdA'_2 , 6 NaA'_2 , 12aq — CuA'_2 , 5aq green needles — CuA'_2 , 8aq — CuA'_2 , 10aq — CuA'_2 , 4 NH_3 — CoA'_2 , 5aq brown needles — CoA'_2 , 4 NH_3 — CoA'_2 , 6 NaA'_2 , 12aq — NiA'_2 , 8aq — NiA'_2 , 6 NaA'_2 , 12aq — FeA'_2 , 5aq yellow crystals — FeA'_2 , 6 NaA'_2 , 12aq — FeA'_2 , (OH) 8aq — MnA'_2 , 5aq (Muller) — MnA'_2 , 8aq (Marchand) — PbA'_2 , 9aq (E. Kopp, *A Ch* [3] 13, 233) S 9 at 15° — PbA'_2 , 2aq — PbA'_2 , (OH) — PbA'_2 , A_2O , 3aq — PbA'_2 , A_2O — $\text{Pb}(\text{OAc})\text{A}'_2$, 4aq — HgA'_2 , A_2O — AlA'_2 , (OH) 8aq — ZnA'_2 , 8aq S (of ZnA'_2) 125 in the cold — ZnA'_2 , 3 NH_3 — ZnA'_2 , 6 NaA'_2 , 12aq — AgA'_2 , aq S 9 at 15° — AgA'_2 , 2 NH_3 . Compounds of picric acid with organic bases and with aromatic hydrocarbons, are described under those bases and hydrocarbons.

Acetyl derivative $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OAc}$ [76°] Yellow crystals (Tommasi a David, *C R* 77, 207)

Benzoyl derivative $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OBz}$

Methyl ether MeA' *Trinitroanisole* [60°] (O), [64°] (Post a Mehrtens, *B* 8, 1552) Formed by nitrating PhOMe (Cahours, *A* 69, 238) and by methylation of picric acid. Monoclinic tables (Friedlander, *J* 1879, 514)

Ethyl ether EtA' [78°] Long needles (Stenhouse a Muller, *A* 141, 80, Willgerodt, *B* 12, 1277)

Iodoethyl ether $\text{C}_2\text{H}_5\text{IA}'$ [70°] From the Ag salt and $\text{C}_2\text{H}_5\text{I}$ (Andrews, *B* 13, 244)

Phenyl ether PhA' Formed from KOPh and $\text{C}_6\text{H}_5\text{Cl}(\text{NO}_2)_2$. Needles (W)

***o* Nitro phenyl ether** $\text{C}_6\text{H}_4(\text{NO}_2)\text{A}'$ [173°]

***p*-Nitro-phenyl ether** $\text{C}_6\text{H}_4(\text{NO}_2)\text{A}'$ [153°] Plates (from alcohol) (Willgerodt, *B* 17, 1766)

Benzyl ether $\text{C}_6\text{H}_5\text{CH}_2\text{A}'$ [147°] Yellow prisms (Kumpf, *A* 224, 131)

***p*-Nitro-benzyl ether** $\text{C}_6\text{H}_4(\text{NO}_2)\text{CH}_2\text{A}'$ [108°] Formed from silver picrate and *p* nitro-benzyl iodide (K)

(β)-**Tri-nitro-phenol** $\text{C}_6\text{H}_2(\text{NO}_2)_3\text{OH}$ [6431] [96°] Formed, together with the (γ) isomeride and tri nitro resorcin (styphnic acid) by the action of conc HNO_3 on (γ) di nitro phenol (Henriques, *A* 215, 325, cf Bantlin, *B* 8, 21) Needles, v e sol alcohol and ether, m sol hot water. Forms with naphthalene a compound [72°] — KA' violet needles, insol alcohol — BaA'_2 , 4aq red prisms

(γ)-**Tri-nitro-phenol** $\text{C}_6\text{H}_2(\text{NO}_2)_3\text{OH}$ [6321] [118°] Formed by nitrating (ϵ) dinitrophenol, and also, together with the (δ) isomeride, by nitrating (γ - or (δ) dinitrophenol (Henriques) White needles. Readily converted into styphnic acid by boiling with fuming HNO_3 . Forms with naphthalene a compound crystallising in yellow needles [100°] — KHA' red needles, insol alcohol — BaA'_2 , golden yellow scales.

References — **Chloro**, **Bromo**, and **Iodo-nitro-phenol**

***o* Nitro-phenol sulphonic acid**

$\text{C}_6\text{H}_3(\text{OH})(\text{NO}_2)(\text{SO}_3\text{H})$ [124] [122°] Formed by sulphonating *o* nitro phenol (Kekulé, *Z* 1867, 641, Armstrong, *Z* 1871, 321, Armstrong a Brown, *B* 7, 923) Formed also by nitration of phenol *p* sulphonic acid (Schmitt a Glutz, *B* 2, 51, Körner, *G* 2, 444, Kolbe a Gauhe, *A* 147, 71) and by boiling (1, 2, 4) bromo nitro benzene sulphonic acid (Goslich, *A* 180, 105) Needles (containing 8aq) Melts at 52° when hydrated, 122° when anhydrous — $\text{NH}_4\text{A}'$ — NaA' , 3aq — $\text{Na}_2\text{C}_6\text{H}_3\text{NSO}_3$, 3aq — KA' — KA' , aq — KA' , 2aq — BaA'_2 , aq — BaA'_2 , 2aq red crystals, sl sol water

***p* Nitro-phenol sulphonic acid** $\text{C}_6\text{H}_4\text{NSO}_3$, *tc* $\text{C}_6\text{H}_3(\text{OH})(\text{NO}_2)(\text{SO}_3\text{H})$ [142] Formed from *p*-nitro phenol and fuming H_2SO_4 (Körner a Post, *B* 5, 852, 1055, 6, 395, 7, 163, *A* 205, 38) Formed also by nitrating phenol *o* sulphonic acid (Stuckenbergh, *A* 205, 45) Crystals (containing 3aq), beginning to decompose when heated at 110° . Give a brown pp with FeCl_3 — KA' monoclinic crystals, $a b c = 1.704 \ 1.1524$, $\beta = 117^\circ 59'$ — KA' , aq — NaA' , 2aq — $\text{Na}_2\text{A}'$, 2aq — CaA' , 3aq — CaA' , 2½aq — BaA' , aq — BaA' , 2aq — PhA' , 1½aq — CuA'

Nitro-phenol disulphonic acid $\text{C}_6\text{H}_3\text{NS}_2\text{O}_6$, *tc* $\text{C}_6\text{H}_2(\text{OH})(\text{NO}_2)(\text{SO}_3\text{H})_2$ Formed from di nitro benzene disulphonic acid by reduction to nitro amido benzene disulphonic acid and displacement of NH_2 by OH (Lamprecht, *B* 8, 289) Minute needles — BaA' , 2aq crystalline

Di-nitro-phenol sulphonic acid $\text{C}_6\text{H}_3\text{NSO}_3$, *tc* $\text{C}_6\text{H}_2(\text{OH})(\text{NO}_2)_2(\text{SO}_3\text{H})$ Formed by the action of nitrous acid on *s* di phenyl hydrazine disulphonic acid (Balentine, *A* 202, 358) Prisms (from alcohol), v sol water. Decomposes at 160° — KA' , ½aq — KA' , 2aq — BaA' , 3½aq (Bertram, *P Beibl* 6, 779)

Tri-nitro-phenol sulphonic acid $\text{C}_6\text{H}_2(\text{OH})(\text{NO}_2)_3(\text{SO}_3\text{H})$ [64213] Formed from phenol *m* sulphonic acid and HNO_3 (Beandsen, *A* 177, 92) — KA' , aq prisms, exploding when heated — BaA' , 3aq crystals, m sol water

***o* Nitro-diphenyl** $\text{C}_{12}\text{H}_9\text{NO}_2$, *tc* $\text{C}_6\text{H}_4\text{C}_6\text{H}_4\text{NO}_2$ [12] [37°] (c 320°) Formed, together with the *p* isomeride, by nitration of diphenyl (Luddens, *B* 8, 870, Hubner, *A* 209, 341, Schultz, *A* 207, 352) Thick plates (from dilute alcohol)

***p* Nitro-diphenyl** [113°] (340° 1 v) Formed as above (Schultz, *A* 174, 210, Hubner, Zimmermann, *B* 13, 1960) Long needles (from alcohol) Yields *p* nitro benzoic acid on oxidation

***oo* Di-nitro-diphenyl** $\text{C}_{12}\text{H}_8\text{N}_2\text{O}_4$, *tc* [21] $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{C}_6\text{H}_4\text{NO}_2$ [12] [124°] Formed from di nitro *p* amido diphenyl by elimination of amidogen (Tauber, *B* 24, 197) Straw-yellow needles, sl sol cold alcohol

***op* Di-nitro-diphenyl** $\text{C}_{12}\text{H}_8\text{N}_2\text{O}_4$, *tc* [21] $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{C}_6\text{H}_4(\text{NO}_2)$ [14] [935°] Formed, together with the *pp* isomeride, by heating diphenyl with HNO_3 and H_2SO_4 (Fittig, *A* 124, 275, Schultz, Schmidt a Strasser, *A* 207, 349). Golden monoclinic needles, $a b c = 1.08 \ 1.91$; $\beta = 87^\circ 30'$ More sol alcohol than the *pp* isomeride.

mm-Di-nitro diphenyl

[8 1] $C_6H_4(NO_2)_2$ $C_6H_4(NO_2)_2$ [1 8] [198°] Formed from di-nitro di *p* amido diphenyl by elimination of the NH_2 groups (Brunner & Witt, *B* 20, 1028) Small yellow needles Yields di-*m*-amido diphenyl on reduction

pp Di-nitro-diphenyl

[4 1] $C_6H_4(NO_2)_2$ $C_6H_4(NO_2)_2$ [1 4] [233°] (Schultz, *A* 174, 221) Formed as above Needles Yields benzidine on reduction

Tetra-nitro-diphenyl $C_6H_4(NO_2)_4$ [140°]

Formed from diphenyl, H_2SO_4 , and HNO_3 (Losanitsch, *B* 4, 404) Amorphous mass, sl sol alcohol

o NITRO-PHENYL-ACETIC ACID $C_6H_4NO_2$

o $C_6H_4(NO_2)CHCOH$ [138°] (*B*), [141°] (*S*) Formed, in small quantity, in the preparation of the *p* isomeride by nitration of phenyl-acetic acid (Radziszewski, *B* 3, 648, Bedson, *C J* 37, 93) Obtained also by saponifying its nitrile (Gabriel & Borgmann, *B* 16, 2066) Needles (from water) or monoclinic plates (from alcohol) Yields *o* nitro benzoic acid on oxidation with $KMnO_4$ Gives oxindole on reduction — BaA' , 2aq

Nitrile $C_6H_4(NO_2)CH_2CN$ *o* Nitro benzyl cyanide [83°] (*B*), [84°] (*S*) Formed in small quantity, together with the *m* and *p* isomerides, by nitration of benzyl cyanide (H Salkowski, *B* 17, 507) Formed also, together with a larger quantity of the compound $C_6H_4(NO_2)CH_2CH_2C_6H_4NO_2$ [111°] and small quantities of $C_6H_4N_2O_2$ [c 238°] and $C_6H_4N_2O_2$ [191°], by boiling *o* nitro benzyl chloride with alcoholic KCy (Bamberger, *B* 19, 2635) Tri metric prisms

m-Nitro-phenyl-acetic acid

[3 1] $C_6H_4(NO_2)CH_2CO_2H$ [117°] (*G* & *B*), [120°] (*S*) Formed from the nitrile which is obtained from *m* nitro benzyl chloride and KCy Needles — AgA' silky needles

Nitrile [61°] Monoclinic crystals

p Nitro-phenyl-acetic acid

[4 1] $C_6H_4(NO_2)CH_2CO_2H$ [152°] Formed as above (Radziszewski, *B* 2, 209, Maxwell, *B* 12, 1765, Gabriel, *B* 14, 2342, 15, 834, Bedson, *C J* 37, 92) Silky needles With *o* nitro-phenyl acetic acid it forms a molecular compound [114°] Yields *p* oxy benzoic acid on oxidation Sodium amalgam yields $N_2(C_6H_4CH_2CO_2H)_2$ [above 300°] (Wittenberg, *Bl* [2] 43, 111) — NaA' , 2aq — BaA'_2 — BaA'_2 , 7aq — ZnA'_2 , aq — AgA' needles

Methyl ether MeA' [55°] NeedlesEthyl ether EtA' [65°] Plates

Amide [192°] Long prisms

Nitrile [116°] Plates Alcohol KOH forms a crimson solution in which diazobenzene chloride ppts $C_6H_4N_2O_2$ [202°] (Czumpelik, *B* 3, 474, Perkin, *C J* 43, 111)

Di-nitro-phenyl-acetic acid $C_6H_4N_2O_2$ *o*

[4 2 1] $C_6H_4(NO_2)_2CH_2CO_2H$ [160°] Formed from phenylacetic acid, H_2SO_4 , and fuming HNO_3 (*R*, Gabriel & Meyer, *B* 14, 823) Formed also by boiling di nitro phenyl-acetoacetic ether with dilute H_2SO_4 (Heckmann, *A* 220, 129) Pale yellow needles, sol hot water Decomposed by heat into di-nitro toluene [71°] and CO_2

Methyl ether MeA' Forms with diazobenzene chloride $C_6H_4(NO_2)_2C(NNHPh).CO_2Me$

[183°] (*V* Meyer, *B* 22, 319) Diazotoluene forms the homologous tolyl hydrazide of methyl di nitro phenyl glyoxylate [168°] crystallising in red needles (Hausknecht, *B* 22, 325) The corresponding derivatives of diaoxylylene and di azonaphthalene melt at 159° and 94° respectively

Ethyl ether EtA' [55°] Needles When heated with alcoholic potash it yields $C_6H_4N_2O_2$ [151°], which forms the salt $K_2C_6H_4N_2O_2$, crystallising in golden plates

Tetra-nitro-di-phenyl-acetic ether

$\{C_6H_4(NO_2)_2\}_2CHCO_2Et$ [154°] Formed from sodium di nitro phenyl acetoacetic ether and biomo di nitro benzene (Von Richter, *B* 21, 2470) Crystalline — $C_{10}H_{10}N_4O_{10}$ [80°] very hygroscopic plates

p-NITRO-PHENYL-ACETIC ALDEHYDE

$C_6H_4(NO_2)CHCHO$ [86°] Formed by boiling the barium salt of $C_6H_4(NO_2)CHClCH(OH)CO_2H$ with water (Lipp, *B* 19, 2645, cf Foirer, *B* 17, 984) Needles, sl sol cold water

DI-NITRO-PHENYL-ACETOACETIC ETHER

[4 2 1] $C_6H_4(NO_2)_2CHAcCO_2Et$ [94°] Formed from $C_6H_4Br(NO_2)_2$ acetoacetic ether, and $NaOEt$ (Heckmann, *A* 220, 128) Plates

Tri-nitro-phenyl-acetoacetic ether

$C_6H_4(NO_2)_3CHAcCO_2Et$ [98°] Formed, together with $\{C_6H_4(NO_2)_2\}_2CHAcCO_2Et$ [205°], from picryl chloride $C_6H_2Cl(NO_2)_3$ and sodium acetoacetic ether (Dittrich, *B* 23, 2720) Crystals, v sol hot alcohol

p-NITRO PHENYL ACETURIC ACID

$C_6H_4(NO_2)CH_2CO_2NHCH_2CO_2H$ [173°] Got by nitrating phenylacetic acid (Hotter, *J p* [2] 38, 110) Needles, decomposed by boiling $HClAq$ into glycooll and *p* nitro phenyl acetic acid — ZnA'_2 , 2aq — AgA' needles, v sl sol cold water

o NITRO PHENYL ACETYLENE $C_6H_4NO_2$

o $C_6H_4(NO_2)CCH$ [82°] Formed by boiling *o* nitro phenyl propionic acid with water (Baeyer, *B* 13, 2259) Needles, sol hot water Gives pps with ammoniacal $AgNO_3$ and Cu_2Cl_2

p-Nitro phenyl acetylene [149°] (*M*), [152°] (*D*)

Formed by boiling *p* nitro phenyl propionic acid with water (Drewson, *A* 212, 158) Formed also from $C_6H_4(NO_2)CHBrCHBrCO_2Et$ and alcoholic potash (*C L* Muller, *A* 212, 133) Needles (from hot water) Gives a red pp with ammoniacal Cu_2Cl_2 and a greenish yellow pp with ammoniacal $AgNO_3$

Di *p* nitro di phenyl-acetylene $C_6H_4N_2O_2$ *o* $C_6H_4(NO_2)_2C \equiv C C_6H_4NO_2$ [288°] Formed from $C_6H_4(NO_2)CHBrCHBrC_6H_4NO_2$ by heating with soda lime at 180° (Elbs & Bauer, *J p* [2] 34, 346) Yellow needles (by sublimation)

o Nitro di phenyl di acetylene $C_6H_4NO_2$ *o* $C_6H_4(NO_2)_2C \equiv C C_6H_4NO_2$ [155°] Formed by the action of K_2FeCy_4 on a mixture of the cuprous salts of phenyl acetylene and *o* nitro phenyl-acetylene (Baeyer & Landsberg, *B* 15, 57). Yellow plates, sol alcohol

Di-*o*-nitro di phenyl diacetylene

$C_6H_4(NO_2)_2C \equiv C C \equiv C C_6H_4(NO_2)_2$ [212°] Formed by the action of an alkaline solution of K_2FeCy_4 on the cuprous salt of *o* nitro-phenyl acetylene (Baeyer, *B* 15, 51) Golden needles, sol chloroform Fuming H_2SO_4 converts it into the isomeric disatogen

NITRO-PHENYL-ACRYLIC ACID *o*. NITRO-CINNAMIC ACID

NITRO-DI-PHENYL-ACRYLIC ACID

Nitrile Theo.-m., and p., varieties, melting at 128°, 134°, and 118° respectively, are formed by the action of the corresponding nitro-benzoic aldehydes on phenyl-acetic nitrile (benzyl cyanide) in presence of alcoholic NaOEt (Frost, *A* 250, 180)

***o*-NITRO-PHENYL-ALLENYL-MALONIC ACID** $C_6H_4(NO_2)CH=CHCH(CO_2H)_2$ [213°] Formed by heating *o* nitro cinnamic aldehyde with malonic acid and HOAc at 100° (Einhorn, *A* 253, 374) Needles—CuA''—AgA'' yellowish plates

***p*-Nitro-phenyl-allenyl-malonic acid** [208°] Formed from *p* nitro-cinnamic aldehyde, malonic acid, and HOAc (Einhorn a Gehrenbeck, *B* 22, 45) Yellow needles (from HOAc) Br forms $C_6H_4(NO_2)CHBrCHBrCBrC(CO_2H)_2$ [206°] crystallising in plates—(NH₄)₂A''—CuA''—AgA'' flocculent pp

Ethyl ether EtA'' [105°] Needles

***o*-NITRO-PHENYL-AMIDO-ACETIC ACID** $C_6H_4(NO_2)NHCH_2CO_2H$ [193°] Formed from bromo-acetic acid and *o* nitro aniline at 125° (Plöchl, *B* 19, 6) Dark red prisms, sl sol ether Yields oxy-quinoxaline dihydride on reduction—NH₄A' flat orange prisms

***p*-NITRO-PHENYL- ω -AMIDO-ACETO-PHENONE** $C_6H_4COCH_2NHCH_2NO_2$ [167°] Formed by heating its nitrosamine with HCl Möhlau, *B* 15, 2474) Golden needles (from HOAc) Yields acetophenone and *p* phenylene-diamine on reduction

Nitrosamine $C_6H_4NH_2NO$, ω

$C_6H_4COCH_2N(NO)C_6H_4NO_2$ Formed from phenyl-amido-acetophenone, HOAc, and nitrous acid gas (M) Plates, decomposing at 135°–145°

Di-nitro-phenyl- ω -amido-acetophenone $C_6H_4COCH_2NHCH_2(NO_2)_2$ [172°] Formed by nitration of phenyl amido acetophenone (Möhlau, *B* 15, 2479) Golden prisms (from HOAc) Yields tri amido benzene on reduction

***s*-TRI-NITRO-TRI-PHENYL-TRI-AMIDO-BENZENE** $C_6(NHPh)(NO_2)_3$ [238°] Formed from $C_6Br_3(NO_2)_3$ and aniline (Jackson a Wing, *Am* 10, 283) Orange powder, insol water

***m*-NITRO-PHENYL-*p*-AMIDO-BENZOIC ACID** $C_6H_4(NO_2)(NHPh)CO_2H$ [3 4 1] [254°] Formed by heating (4,3,1) bromo nitro benzoic acid with aniline (Schöpf, *B* 22, 3281) Garnet red needles Yields an amido-acid [153°]—NaA'—NaA'aq—BaA', 3aq—AgA' orange plates

Ethyl ether EtA' [123°] Hexagonal

Anilide $C_6H_4(NO_2)(NHPh)(NHPh)CO$ [216°] Formed by heating aniline with bromo nitro-benzoyl chloride (Grohmann, *B* 23, 3448) Blood red leaflets from HOAc

Nitrile $C_6H_4(NH_2)(NHPh)CN$ [126°] Formed from bromo nitro benzonitrile and aniline (Schöpf, *B* 23, 3444)

***o*-Nitro-phenyl-*m*-amido-benzoic acid** $C_6H_4(NO_2)(NHPh)CO_2H$ [2 3 1] [248°] Formed from (3,2,1)-bromo-nitro-benzoic acid and aniline (Schöpf, *B* 23, 3440) Yellow needles—NaA' 2aq—BaA', 5aq

Ethyl ether EtA' [112°] Needles

***m*-Nitro-phenyl-*o*-amido-benzoic acid** *Nitrile* $C_6H_4(NO_2)(NHPh)CN$ [5 2 1] [170°] Formed from (2,5,1)-bromo-nitro-benzonitrile and aniline (S.) Lemon-yellow needles

Di-nitro phenyl-*o*-amido-benzoic acid

$C_6H_4N_2O_6$, ω [4 2 1] $C_6H_4(NO_2)_2NHCH_2CO_2H$ [264°] Formed by warming *o* amido benzoic acid with $C_6H_4Cl(NO_2)_2$ (Jourdan, *B* 18, 1448) Orange needles, almost insol. water—BaA', dark red crystalline powder

NITRO-PHENYL-AMIDO-NAPHTHOQUINONE *v* (a)-NAPHTHOQUINONE, *Reactions* 16 and 17

DI-NITRO-PHENYL-DI-AMIDO-DIPHENYL

$C_6H_4N_2O_6$, ω $C_6H_4(NH_2)C_6H_4NHCH_2C_6H_4(NO_2)_2$ [245°] Formed by boiling benzidine with alcohol and (1,2,4) chloro di nitro-benzene (Willgerodt, *B* 9, 981) Long needles (from HOAc)

Di *o* nitro-di-phenyl diamido diphenyl

$C_6H_4(NO_2)_2NHCH_2C_6H_4NHCH_2C_6H_4(NO_2)_2$ [240°] Formed by boiling benzidine with *o* chloro nitrobenzene and alcohol (Schöpf, *B* 22, 904) Needles (from HOAc)

Tetra-nitro di-phenyl-di-amido-diphenyl

$C_6H_4(NO_2)_2NHCH_2C_6H_4NHCH_2C_6H_4(NO_2)_2$ [above 330°] Formed from [1 2 4] $C_6H_4Cl(NO_2)_2$, alcohol, and benzidine at 120° (W) Yellow powder, sl sol alcohol

DI-NITRO-PHENYL-AMIDO-TOLYL AMINE

[4 2 1] $C_6H_4(NO_2)_2NHCH_2C_6H_4MeNH_2$ [147°] Formed from tolylene *o* diamine and *i* chloro di nitro benzene (Ernst, *B* 23, 3428) Brownish yellow needles Yields with HNO₃ the azimide

$C_6H_4(NO_2)_2N \llcorner \begin{smallmatrix} C_6H_5 \\ N \end{smallmatrix} \gg N$ [186°]

***o* NITRO DI PHENYL AMINE** $C_6H_4N_2O_6$, ω

[2 1] $C_6H_4(NO_2)NHC_6H_5$ [75°] Formed from aniline and *o* chloro nitro benzene or *o* bromo nitro benzene at 100° (Schöpf, *B* 22, 903, 23, 1839) Trimetric crystals (from alcohol), $a \cdot b \cdot c = 468 \cdot 1 \cdot 671$ Yields on reduction the amido compound [80°]

***p* Nitro-diphenylamine**

[4 1] $C_6H_4(NO_2)NHCH_2C_6H_5$ [133°] Formed from benzoyl diphenylamine by nitration and elimination of Bz (Hofmann, *A* 132, 167, Lellmann, *B* 15, 825) Formed also from its nitrosamine by treatment with aniline (Witt, *C* 33, 205) Pale-yellow scales (from dilute alcohol) Colours alcoholic potash scarlet Dyes silk yellow

Benzoyl derivative [129°] Prisms

Nitrosamine $C_6H_4(NO_2)N(NO)C_6H_5$

[134°] Formed by warming diphenylamine with HNO₃, isoamyl nitrite, and alcohol (W) Crystals, sol chloroform

Di-o-nitro diphenylamine $NH(C_6H_4NO_2)_2$

[220°] (L), [212°] (W) Obtained from its benzoyl derivative, and also, together with the *p*-isomeride, by the action of alcohol (50 c c), aniline (25 g) and aniline hydrochloride (30 g) at 100° on the mixed di nitro di phenyl nitrosamines prepared from diphenylamine (17 g), amyl nitrite (48 g), alcohol (50 c c), nitric acid (40 c c of SG 1.424), and HOAc (50 c c) (Witt, *C* J 33, 208) Red felted needles

Benzoyl derivative $NBz(C_6H_4NO_2)_2$

Formed, together with that of the *p* isomeride from benzoyl diphenylamine and fuming HNO₃ (Lellmann, *B* 15, 827)

Di *p*-nitro-diphenylamine $C_6H_4N_2O_6$, ω

$NH(C_6H_4NO_2)_2$ [216°] (L), [214°] (W) Got as above Yellow needles with blue reflex

Benzoyl derivative [224°] Monoclinic crystals, sl sol alcohol.

Di nitro diphenylamine $C_{12}H_8N_2O_4$, *s*.
 $C_6H_5NH C_6H_4(NO_2)_2$ [1 2 4]. [157°]. Formed from $C_6H_5Br(NO_2)$, or $C_6H_5Cl(NO_2)$, and aniline or *p*-phenyl thio urea (Clemm, *B* 3, 128, Willgerodt, *B* 9, 977, 11, 601, cf Hepp, *Bl* [2] 80, 4)

Tri-nitro diphenylamine $C_{12}H_5N_3O_6$, *s*.
 $C_6H_5NH C_6H_4(NO_2)_2$ [1 2 4 6]. [175°]. Formed from $C_6H_5Cl(NO_2)_2$ (picryl chloride) and aniline (Clemm, *B* 3, 126) Scarlet prisms

Tri nitro diphenylamine
 $[3] C_6H_5(NO_2) NH C_6H_4(NO_2)_2$ [1 2 4]. [194°] Formed from $C_6H_5Br(NO_2)$, or $C_6H_5Cl(NO_2)_2$, and *m* nitro aniline (Austen, *B* 7, 1250, Willgerodt, *B* 9, 1178) Short yellow needles (from HOAc)

Tri nitro diphenylamine
 $[4] C_6H_5(NO_2) NH C_6H_4(NO_2)_2$ [181°] Formed from *p* nitro aniline and bromo di nitro benzene (A) Yellow powder, *v* sol HOAc

Tri nitro diphenylamine [135°] Formed by boiling the acetyl derivative of diphenylamine with dilute nitric acid (S G 1029) (Norton & Allen, *B* 18, 1997) Yellow needles, *v* sol alcohol

Tetra nitro diphenylamine $C_{12}H_4N_4O_8$, *s*.
 $[3] C_6H_4(NO_2) NH C_6H_4(NO_2)_2$ [1 2 4 6]. [205°] Formed from *m* nitro aniline and picryl chloride (Austen, *B* 7, 1248) Orange crystals (from HOAc)

Tetra nitro diphenylamine
 $[4] C_6H_4(NO_2) NH C_6H_4(NO_2)_2$ [1 2 4 6]. [216°]. Formed in like manner from *p*-nitro aniline (A)

Tetra nitro diphenylamine $NH[C_6H_4(NO_2)_2]_2$. [180°] Got by heating $C_6H_5(NO_2)_2NH CO_2Et$ with alcoholic potash (Hager, *B* 17, 2629) Reddish brown plates (from alcohol)

Tetra nitro diphenylamine [192°] Formed by nitration of diphenylamine, diphenyl nitrosamine, and diphenyl methylamine $NMePh$. (Gnehm & Wyss, *B* 10, 1318) Yellow crystals (from alcohol) Forms a scarlet solution in $NaOH_{aq}$

Hexa nitro-diphenylamine $[C_6N_2(NO_2)_4]_2NH$ *Dipicrylamine* [238°] (A), [234°] (M) Formed by nitrating diphenylamine, diphenyl methylamine, or tetra nitro diphenylamine [216°] (Austen, *B* 7, 1250, Gnehm, *B* 7, 1399, 9, 1245, 1557, Mertens, *B* 11, 845) Yellow prisms (from acetic acid) Its ammonium salt $NH_4C_{12}H_4N_6O_{12}$ is used as a yellow dye ('aurantia') — $Ba(C_{12}H_4N_6O_{12})_2$ red rhombohedra

Hexa nitro diphenylamine [261°] Formed by nitrating tetra nitro diphenylamine [205°] (A) Small yellow crystals (from HOAc). Explodes when heated.

Nitro tri phenyl amine $(C_6H_5)_3N C_6H_4(NO_2)$ [140°] Formed from triphenylamine, HOAc, and HNO_3 (Herz, *B* 23, 2537) Golden plates

Di nitro-tri phenyl amine $C_6H_5N(C_6H_4NO_2)_2$. [207°] Formed from triphenylamine (2 g), HOAc (35 g) and HNO_3 (2 g) at 60° (Herz, *B* 23, 2538) Yellow needles, *v* sol benzene

Tri nitro-tri-phenylamine $N(C_6H_4NO_2)_3$. [280°] Formed from triphenylamine (2 g), HOAc (35 g), and HNO_3 (4 g) at 100° (Heydrich, *B* 18, 2156, Herz, *B* 23, 2559). Bronze-yellow needles, *v* sol HOAc

***m* Nitro-phenyl-angelic aldehyde** $C_6H_4(NO_2)CH CH_2CHO$ [46°] Formed from *m* nitro benzoic aldehyde, butyric acid, and dilute $NaOH_{aq}$ (Von Miller & Rohde, *B* 23,

1888) Plates, reduced by tin and HCl to amido-ethyl indonaphthene [89°]

Phenyl hydraside [185°] Red needles.

DI NITRO DI-*p*-PHENYL BENZENE

$C_{18}H_{12}(NO_2)_2$. [277°] Formed by nitration of diphenylbenzene (Schmidt & Schultz, *B* 11, 1755, A 203, 125) Yellow monoclinic needles (from nitro benzene)

Tri-nitro-di-*p*-phenyl-benzene $C_{18}H_{11}(NO_2)_3$. [195°] Formed from diphenylbenzene and fuming HNO_3 (S & S) Needles Yields on reduction a base [170°]

Tri-nitro diphenylbenzene [200°] Formed by nitration of isodiphenylbenzene (S & S). Needles Yields on reduction a base [288°]

Tetra-nitro-tri-phenyl-benzene $C_{18}H_9(NO_2)_4$. [above 370°] Formed, together with an isomeride [108°], by nitrating triphenylbenzene (Mel lin, *B* 23, 2535) Both bodies crystalline in needles

NITRO-PHENYL-BENZYLIDENE-AMINE

$C_{11}H_9N_2O_2$, *s*. $C_6H_5CH N C_6H_4NO_2$. [66°] (Lazorenko, *J* 1870, 760), [73°] (Lachovitch, *M* 9, 695) Formed from benzoic aldehyde (or hydrobenzamide) and *m*-nitro aniline Needles

***p* Nitro phenyl-benzylidene-amine** $C_6H_5N C_6H_4(NO_2)$ [1 4]. [115°] Formed from hydrobenzamide by warming with *p* nitro-aniline Yellow needles (Lachovitch)

Isomerides *v* Nitro BENZYLIDENE ANILINE

NITRO-PHENYL BENZYL OXIDE

$C_6H_4(NO_2)OCH_2Ph$ Formed from potassium nitro phenol, alcohol, and benzyl chloride (Kumft, A 224, 121) The *o*- compound melts at 29°, and the *p* compound at 106° Both are crystalline

Di nitro phenyl benzyl oxide $C_{11}H_9N_2O_3$, *s*. $C_6H_4(NO_2)_2OCH_2Ph$ Formed from silver di nitrophenol and benzyl iodide (K) The (4,2,1)-compound melts at 149°, and the (6,2,1) compound at 76° ($OH=1$) Both crystalline from HOAc

Tri nitro phenyl benzyl oxide $C_{11}H_7N_3O_3$, *s*. $[6.4.2.1] C_6H_3(NO_2)_3OCH_2Ph$ [147°] Formed from silver picrate and benzyl iodide (K). Yellowish prisms (from benzene)

TRI-NITRO-DI-PHENYL-BENZYL-PHOSPHINE OXIDE $PO(C_6H_4NO_2)_2(C_6H_5NO_2)$ [206°]. Formed from the oxide, fuming HNO_3 , and conc H_2SO_4 (Dörken, *B* 21, 1505) Crystals, *m* sol HOAc

DI-NITRO-PHENYL-BENZYL SULPHIDE $C_6H_5S C_6H_4(NO_2)_2$ [1 2 4]. [128°] Formed from (1,2,4) chloro di nitro-benzene, alcoholic K_2S , and benzyl chloride (Willgerodt, *B* 18, 331) Yellowish plates

NITRO-PHENYL-BROMO-PROPIONIC ACID

o. BROMO-NITRO PHENYL PROPIONIC ACID

***o*-NITRO-PHENYL-BUTINYL METHYL KETONE** $C_6H_4(NO_2)C_4H_7COCH_3$ [78.5°]. Formed, together with $(C_6H_4(NO_2)O.H_2)CO$ [208.5°], from *o* nitro cinnamic aldehyde, acetone, alcohol, and dilute (2 *p*.) $NaOH_{aq}$ (Diehl & Einhorn, *B* 18, 2327) Broad needles (from alcohol)

o-NITRO-PHENYL-BUTINYL PHENYL-BUTINYL KETONE

$C_6H_4(NO_2)C_4H_7CO C_6H_5C_4H_7$ [186.5°]. Formed from *o* nitro-cinnamic aldehyde, phenyl-butynyl methyl ketone, alcohol, and dilute $NaOH_{aq}$

(Drebl & Einhorn, *B* 18, 2829) Golden crystals (from acetone)

***p*-NITRO-PHENYL-ISOBUTYRIC ACID**

$C_6H_4(NO_2)CH_2CHMeCO_2H$ [121°] Formed from phenyl isobutyric acid and conc HNO_3 (Edeleanu, *C. J* 53, 559) Small prisms, sol alcohol

NITRO-PHENYL-CARBAMIC ETHER

$C_6H_4(NO_2)NHC(O)Et$ The *o* compound [58°] and the *p* compound [129°] are got from the corresponding nitro aniline and $ClCO_2Et$ (Rudolph, *B* 12, 1295; Hager, *B* 17, 2625) The *p* compound is also got by nitration of phenyl carbamic ether (Behrend, *A* 233, 9) Both are crystalline

Di nitro-phenyl carbamic ether $C_6H_4N_2O_5$ *re* [4 2 1] $C_6H_4(NO_2)_2NHCO_2Et$ [111°] Formed by nitration of either *o*- or *p*-nitro phenyl-carbamic ether (Hager, *B* 17, 2629) Needles (from alcohol) An isomeride [210°] is got by the action of HNO_3 on $C_6H_4NHCSOEt$ (Losanitsch, *B* 10, 691)

***Di* nitro *di* phenyl carbamic ether**

$(C_6H_4NO_2)_2NCO_2Et$ The only *o*- compound is formed together with the *p*-compound [134°] by nitration of *di* phenyl carbamic ether (Hager, *B* 18, 2574) The *p*-compound is the less sol. alcohol

***m*-NITRO-TRI-PHENYL-CARBINOL**

$C_6H_4(NO_2)CPh_2OH$ [75°] Formed from $C_6H_4(NO_2)CHPh_2$ by bromination in sunlight, followed by successive treatment with $KOAc$ and KOH (Tschacher, *B* 21, 190) Colourless crystals, sol ligroin

p Nitro-tri phenyl-carbinol [136°] Formed by oxidising *p* nitro tri phenyl methane with CrO_3 in $HOAc$ (Baeyer & Lohr, *B* 23, 1623) Crystals (from dilute $HOAc$)

Tri-*p*-nitro tri phenyl carbinol $C_6H_4(NO_2)_3$ *re* $C_6H_4(NO_2)_3COH$ [172°] Formed by oxidising tri nitro tri-phenyl methane (E & O Fischer, *B* 11, 1079) Colourless crystals (from $HOAc$)

TETRA-NITRO-DI-PHENYL CARBONATE

$(C_6H_4(NO_2)_2)_2CO$ [127°] Formed from *di* phenyl carbonate, HNO_3 , and H_2SO_4 (Kempf, *J pr* [2] 1, 407, Löwenberg, *C C* 1886, 390) Nodules, *v* sl sol ether

NITRO-DIPHENYL CARBOXYLIC ACID

$C_6H_4(NO_2)_2$ [222°] Formed by nitration of diphenyl *o* carboxylic acid (Schmidt, *A* 193, 115) Monoclinic crystals (from alcohol) — BaA' — CaA' nodules, *v* sol water

Nitro diphenyl carboxylic acid $C_6H_4NO_5$ *re* [2 1] $C_6H_4(CO_2H)C_6H_4(NO_2)CO_2H$ [1 4 2] [217°] Formed by oxidation of nitro phenanthraquinone [257°] (Strasburger, *B* 16, 2347) Light yellow needles (from water)

***Di* nitro-diphenyl carboxylic acid**

[4 1] $C_6H_4(NO_2)_2C_6H_4(NO_2)CO_2H$ [1 2 4] [252°] Formed by nitrating diphenyl *p* carboxylic acid (Strasser & Schultz, *A* 210, 192) Needles, *m* sol alcohol

Methyl ether MeA' [156°] Needles

(a) *Di* nitro diphenyl dicarboxylic acid

$C_6H_4(NO_2)_2(CO_2H)_2$ [249°] (*H*), [253°] (Schultz, *A* 196, 26) Formed by oxidation of *di* nitro phenanthraquinone (Struve, *B* 10, 75), and, together with the following acid, by nitration of diphenyl *di* *o* carboxylic acid (Hummel, *A* 193, 129) Pale yellow needles (containing aq) — BaA' , 6aq long prisms.

Methyl ether MeA' [178°] Prisms.

(*β*) *Di*-nitro-diphenyl dicarboxylic acid. [297°] Formed as above (Schultz, *A* 203, 105). Needles (from hot water) — BaA' , 4aq triqmic prisms

Methyl ether MeA' [132°] Tables

NITRO-PHENYL CHLORO-METHYL KETONE

***o* CHLORO NITRO ACETOPHENONE**

***o* NITRO-PHENYL CINNAMIC ACID**

$C_6H_4NO_2$ [196°] Formed from *o* nitro benzoic aldehyde, sodium phenyl acetate, and Ac_2O at 160° (Oghaloro & Rosini, *G* 20, 396) Needles — BaA' 5aq — BaA' 8aq

NITRO-PHENYLENE *o* DIAMINE

$C_6H_4(NO_2)(NH_2)_2$ [4 2 1] [198°] Formed by reducing (4,2,1) *di* nitro aniline with alcoholic ammonium sulphide (Gottlieb, *A* 85, 27; Heum, *B* 21, 2305) Dark red plates (from toluene). Ac_2O at 190° forms $C_6H_4(NO_2)_2NHCM_6$ [216°]. — $B'HCl$ aq — $B'H.PtCl_5$ — $B'HNO_3$ — $B'H_2SO_4$ — $B'H_2C_2O_4$ — $B'H_2PtCl_5$, 5aq

***Di* acetyl derivative [227°] Needles**

Nitro phenylene *m* diamine $C_6H_4N_2O_5$ *re* $C_6H_4(NO_2)(NH_2)_2$ [161°] Obtained from its diacetyl derivative, which is got by nitrating $C_6H_4(NHAc)_2$ (Barbaglia, *B* 7, 1259) Yellowish red prisms, sol water

Di acetyl derivative $C_6H_4(NO_2)(NHAc)_2$ [246°] Slender needles (from alcohol)

***Di* benzoyl derivative**

$C_6H_4(NO_2)(NHBz)_2$ [222°] Got by nitrating dibenzoyl phenylene *m* diamine (Ruhemann, *B* 14, 2652) Needles (from $HOAc$)

Nitro phenylene *p* diamine

$C_6H_4(NO_2)(NH_2)_2$ [2 4 1] [187°] Formed from its diacetyl or dibenzoyl derivatives, got by nitrating the corresponding derivatives of phenylene *p* diamine (Ladenburg, *B* 17, 149; Hinsberg, *A* 254, 255) Dark needles with green lustre

Di acetyl derivative $C_6H_4(NO_2)(NHAc)_2$ [186°] On warming with $NaOHAq$ it yields the mono acetyl derivative crystallising in red needles [195°]

Di* nitro phenylene diamine $C_6H_4N_2O_5$ *re

$C_6H_4(NO_2)_2(NH_2)_2$ [211°] Formed by reducing picramide $C_6H_4(NO_2)_2NH_2$ [6 4 2 1] with ammonium sulphide (Norton & Elliott, *B* 11, 327) Red needles, almost insol cold water

Di acetyl derivative $C_6H_4(NO_2)_2(NHAc)_2$ [246°] Yellow needles

***Di*-nitro-phenylene-*m* diamine**

$C_6H_4(NO_2)_2(NH_2)_2$ [4 2 3 1] [c 250°] Formed by heating *di* nitro resorcin with aqueous ammonia (Barr, *B* 21, 1545) Brownish needles (from $HOAc$), sl sol alcohol

***Di* nitro-phenylene-*m*-diamine**

$C_6H_4(NO_2)_2(NH_2)_2$ [c 300°] Obtained by saponification of its *di* acetyl derivative which is formed by nitration of *di* acetyl *m* phenylene-diamine (Nietzki & Hagenbach, *B* 20, 333). Orange yellow crystals

***Di* acetyl derivative [228°] Needles**

Di nitro-phenylene-*p* diamine *Di* acetyl derivative [258°] Formed by nitrating diacetyl phenylene *p* diamine (Nietzki & Hagenbach, *B* 20, 331) Yellow solid, sl sol alcohol

Tri-nitro-phenylene *m* diamine $C_6H_4N_3O_6$ *re*

$C_6H(NO_2)_3(NH_2)_2$ [6 4 2 3 1] [above 250°] Formed by boiling the *di* ethyl ether of tri-nitro resorcin with alcoholic ammonia (Nöling & Collin, *B* 17, 260; Barr, *B* 21, 1546).

Yellow granules Reduced by SnCl_4 and HCl to unstable penta amido benzene

NITRO-DIPHENYLENE-KETONE $\text{C}_{12}\text{H}_9\text{NO}$, [3 6] $\text{C}_6\text{H}_4(\text{NO}_2) < \frac{\text{O}}{\text{CO}} >$ [220°] Formed, as well as di nitro diphenylene ketone [290°], by nitration of *o* diphenylene ketone (Schultz, *A* 203, 103) The isomeric nitro derivative of iso diphenylene ketone [83°] melts at 220°–230° (Carnelley & Dunn, *B* 21, 2005)

Nitro-diphenylene-ketone carboxylic acid $\text{C}_{12}\text{H}_7(\text{NO}_2)_2\text{O}_4$ [246°] Got by warming di phenylene ketone carboxylic acid with HNO_3 (Fittig & Liepmann, *A* 200, 6) Golden needles (from alcohol) —BaA', 4aq yellow needles, sl sol water

(a) **DI NITRO DIPHENYLENE KETONE OXIDE** $\text{C}_6\text{H}_4(\text{NO}_2) < \frac{\text{O}}{\text{CO}} > \text{C}_6\text{H}_4(\text{NO}_2)$ [190°]

Formed, together with a (β) isomeride [260°], by warming diphenylene ketone oxide with HNO_3 (Wichelhaus & Salzmänn, *B* 10, 1401, *R* Richter, *J pr* [2] 23, 292, *A G* Perkin, *C J* 43, 189, Gracbe, *A* 274, 286) An isomeric body [224°] is formed, together with a di nitro-derivative [235°], by nitration of iso di phenylene ketone oxide [91°] (*R*)

NITRO-PHENYLENE-DI-METHYL-*o* DIAMINE $\text{C}_6\text{H}_3(\text{NO}_2)(\text{NMe}_2)(\text{NH})$ [4 1 2] [63°] Formed by reducing di nitro dimethylaniline by ammonium sulphide (Heim, *B* 21, 2308) Orange needles (from water)

Nitro-phenylene tri methyl diamine $\text{C}_6\text{H}_3(\text{NO}_2)(\text{NMe}_2)(\text{NHMe})$

Acetyl derivative [c 210°]
Nitrosamine $\text{C}_6\text{H}_3(\text{NO}_2)(\text{NMe}_2)(\text{NMeNO})$ [87°] Formed from $\text{C}_6\text{H}_3(\text{NMe}_2)_3$ and HNO in excess (Wurster & Schobig, *B* 12, 1811) Needles

Tri nitro phenylene di methyl diamine $\text{C}_6\text{H}_3\text{N}_3\text{O}_6$ *ie* $\text{C}_6\text{H}(\text{NO}_2)_3(\text{NHMe})_2$ [235°] Formed by heating its nitramine with phenol or aqueous methylamine (Romburgh, *R T C* 7, 6, 8, 279) Golden crystals (from HOAc)

Di nitramine $\text{C}_6\text{H}_3(\text{NO}_2)_2(\text{NMe NO})_2$ Formed by boiling $\text{C}_6\text{H}_3(\text{NMe}_2)_3$ with fuming HNO_3 (Romburgh, *R T C* 6, 252) Pale yellow crystals, decomposing at about 205° A nitramine $\text{C}_6\text{H}_3(\text{NO}_2)(\text{NHMe})(\text{NMeNO})_2$ is got by the action of methylamine on $\text{C}_6\text{H}_3(\text{NO}_2)_3\text{NMeNO}$. It separates from HOAc in golden crystals [192°]

NITRO-PHENYLENE (α) NAPHTHYL *o* DIAMINE $\text{C}_6\text{H}_3(\text{NO}_2)(\text{NH}_2)(\text{NHC}_6\text{H}_4)$ [147°] Formed from (4, 2, 1) di nitro phenyl (α) naphthylamine and ammonium sulphide (Heim, *B* 21, 2302) Needles (from HOAc), v e sol alcohol The isomeric compound from di nitro-phenyl (β) naphthylamine melts at 195°, and yields an acetyl derivative [200°], which is converted by Ac_2O into nitro phenylene (β) naphthyl acetanilide [162°]

DI-NITRO-PHENYLENE NAPHTHYLENE OXIDE $\text{C}_{12}\text{H}_7(\text{NO}_2)_2\text{O}$ [235°] Formed by nitrating (α) phenylene (α) naphthylene oxide (Arr, *A* 209, 141) V sol ether and HOAc

NITRO-PHENYLENE NAPHTHYL-ACET-AMIDINE $\text{C}_6\text{H}_3(\text{NO}_2) < \frac{\text{N}(\text{C}_6\text{H}_4)}{\text{NMe}} >$ [162°]

Formed from nitro amido phenyl (β) naphthylamine and Ac_2O (Heim, *B* 21, 589) Needles.

DI-NITRO-DIPHENYLENE OXIDE $\text{C}_{12}\text{H}_8(\text{NO}_2)_2\text{O}$ [200°] Formed from diphenyl-

one oxide and fuming HNO_3 (Hoffmeister, *A* 159, 211) Crystals (from alcohol)

NITRO PHENYLENE-UREA $\text{C}_6\text{H}_4\text{N}_2\text{O}$, *ie* $\text{C}_6\text{H}_4(\text{NO}_2) < \frac{\text{NH}}{\text{NH}} > \text{CO}$ Formed by heating nitro amido phenyl carbamic ether (Hager, *B* 17, 2630) Colourless needles, not melted at 300°.

NITRO-*o* DI-PHENYL ETHANE $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CHPh CH}_3$ [80°] Formed, together with $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CH CH}_3$ [149°], by nitration of *o* di phenyl ethane (Anschütz & Romig, *B* 18, 664) Long yellow needles

Di-*p* nitro-*s* di phenyl ethane [4 1] $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CH}_2\text{CH}_2\text{C}_6\text{H}_4(\text{NO}_2)$ [1 4] [179°] Formed, together with an isomeride [75°], from dibenzyl and fuming HNO_3 (Stelling & Fittig, *A* 137, 260, Leppert, *B* 9, 15) Formed also from *p* nitro benzyl chloride, SnCl_4 , and NaOHAq (W Roser, *A* 238, 364) Needles, sl sol alcohol.

DI NITRO DI PHENYL ETHANE DICARB OXYLIC ACID $\text{C}_{12}\text{H}_{12}(\text{NO}_2)_4\text{O}_4$ Formed by nitrating *s* di phenyl ethane di *o* carboxylic acid (Dobref, *A* 239, 70) Minute crystals, melting above 300° —CaA''

Ethyl ether LiHA'' [60°]
Di nitro di phenyl ethane dicarboxylic acid [226°] Formed by nitration of *s* di phenyl ethane *aB* dicarboxylic acid (Reimer, *B* 14, 1802) Amorphous (containing aq) An isomeric acid [242°] is got by nitrating *s* di phenyl ethane *aa* dicarboxylic acid Both acids yield *p* nitro benzoic acid on oxidation

***o* NITRO PHENYL-ETHYL CARBONATE** $\text{C}_6\text{H}_3(\text{NO}_2)\text{OCO OEt}$ (275° 285°) Formed by the action of chloro formic ether upon the potassium salt of *o* nitro phenol (Bender, *B* 19, 2268) Heavy yellow oil

NITRO-PHENYL-ETHYLENE *v* Nitro-STERENE

Nitro *s* di phenyl ethylene $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CPh CH}_3$ [36°] Formed from $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CPh}(\text{OH})\text{CH}_3$ and AcCl (Anschutz & Romig, *B* 18, 664) Yellow crystals (from ether)

Di-*o* nitro di phenyl ethylene $(\text{C}_6\text{H}_4(\text{NO}_2))_2\text{CH}$ Formed in two modifications 'cis' [126°] and 'trans' [196°], by the action of alcoholic potash upon *o* nitro benzyl chloride (Bischoff, *B* 21, 2072, 23, 2072) Both crystallise in needles

Di *p* nitro di phenyl ethylene Formed in two modifications, [c 213°] and [c 282°], by the action of alcoholic potash on *p* nitro benzyl chloride (Walden, *B* 23, 1959, cf Strakosch, *B* 6, 328) Both are crystalline

DI-*m*-NITRO-DI-PHENYL-ETHYLENE-DIAMINE $(\text{C}_6\text{H}_4(\text{NO}_2)_2\text{NH})_2\text{C}_6\text{H}_4$ [206°] Obtained by heating *m* nitro aniline with ethylene bromide at 130° (Gattermann & Hager, *B* 17, 778) Reddish yellow crystals, insol alcohol

NITRO-*s*-DI-PHENYL-ETHYLENE CARB OXYLIC ACID *Nitrile*

$\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CCy CHPh}$ [176°] Got from *p*-nitro-phenyl acetonitrile, benzoic aldehyde, and alcoholic NaOEt at 50° (Remse, *B* 23, 8134). Yellow needles By using *o* or *m* nitro benzoic aldehyde, the corresponding di nitro-compounds [4 1] $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CCy CH C}_6\text{H}_4(\text{NO}_2)$ [2 1] [185°] and [4 1] $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CCy CH C}_6\text{H}_3(\text{NO}_2)$ [3 1] [195°] may be made Both crystallise in needles.

***m*-NITRO-PHENYL-ETHYLENE-QUINOLINE** $\text{C}_6\text{NH}_2\text{CH CH C}_6\text{H}_4\text{NO}_2$ [136°] Ob-

tained by heating methyl quinoline (lepidine) with *m* nitro benzoic aldehyde and KHSO_4 at 160° (Heymann & Königs, *B* 21, 1424). Needles

NITRO-PHENYL ETHYL KETONE

$\text{C}_6\text{H}_4(\text{NO}_2)\text{CO.C}_2\text{H}_5$ [100°] Formed, together with a syrupy isomeride, by nitration of phenyl ethyl ketone (Barry, *B* 6, 1007) Prisms

NITRO-PHENYL-FURFURYL-ACRYLIC

ACID *Nitrile* $\text{C}_6\text{H}_4(\text{NO}_2)\text{CO.C}_4\text{H}_3\text{O}$ [173°] Formed from furfuraldehyde, *p* nitro phenyl acetone, and alcoholic NaOEt (Freund & Immerwahr, *B* 23, 2852) Needles

o-NITRO-PHENYL-GLYCIDIC ACID

$\text{C}_6\text{H}_3(\text{NO}_2)\text{CH(OH)CH(OH)CO}_2\text{H}$ *Nitro phenyl oxyacrylic acid* [108°] Formed from $\text{C}_6\text{H}_3(\text{NO}_2)\text{CH(OH)CHClCO}_2\text{H}$ and alcoholic potash (Baeyer, *B* 13, 2262; Morgan, *B* 17, 219; Lipp, *B* 19, 2649) Prisms (containing aq) Melts at 94° when hydrated, 108° when anhydrous Yields indigo and CO_2 on heating $-\text{NH}_4\text{A}'$ aq $-\text{AgA}'$ white crystalline pp

p-Nitro-phenyl-glycidic acid [188°]

Formed in the same way as the *o* isomeride (Lipp), and also by the action of HOCl on sodium *p* nitro cinnamate (Erlenmeyer, *B* 14, 1868) Plates (from hot water) Yields $\text{C}_6\text{H}_3(\text{NO}_2)\text{CH(OH)CH(OH)CO}_2\text{H}$ on boiling with dilute H_2SO_4

NITRO-PHENYL GLYCOCOLL v NITRO

PHENYL AMIDO ACETIC ACID

NITRO-PHENYL GLYCOLLIC ACID v GLYCOLLIC ACID

o-NITRO-PHENYL GLYOXYLIC ACID

$\text{C}_6\text{H}_3(\text{NO}_2)\text{CO.CO}_2\text{H}$ [49°] Formed from its amide, which is got by the action of cold conc HClAq on $\text{C}_6\text{H}_3(\text{NO}_2)\text{CH}_2\text{Cy}$ (Claisen & Shadwell, *B* 12, 352; Fehrlin, *B* 23, 1577) Needles *Amide* [189°] (*C* & *S*), [199°] (*F*) *Nitrile* [54°] Prisms (from ligroin)

Phenyl hydrazide [166°] Changed by dissolving in alcoholic KOH and adding HCl into an isomeride [190°] HNO_3 converts the first phenyl hydrazide into a body melting at $77^\circ-80^\circ$ and the second into one melting at $95^\circ-100^\circ$ Both yield isatin phenyl hydrazide on reduction The ethyl ether of the phenyl-hydrazide $\text{C}_{10}\text{H}_7\text{N}_2\text{O}_2$ crystallises in yellow prisms [128°] (Krause, *B* 23, 3617)

Phenyl-methyl hydrazide [142°]

Oxam of the ethyl ether

$\text{C}_6\text{H}_3(\text{NO}_2)\text{C(OH)CO}_2\text{Et}$ [163°] Needles (from boiling water) (Gabriel, *B* 16, 519)

m-Nitro-phenyl glyoxylic acid [78°] Formed by boiling its amide with alkalis (Claisen & Thompson, *B* 12, 1944, 14, 1187) Prisms — KA' flat prisms — BaA' aq — AgA' nodules

Amide $\text{C}_6\text{H}_3(\text{NO}_2)\text{CO CONH}_2$ [152°]

Formed from *m* nitro phenyl acetone, and also by nitration of phenyl-glyoxylic amide

Nitrile (231° at 145 mm) Oil

Phenyl hydrazide [176°] (*F*) Crystals Successive treatment with KOH and HClAq yields a green compound [285°]

Di-nitro-phenyl-glyoxylic acid

Phenyl-hydrazide of the methyl ether $\text{C}_6\text{H}_3(\text{NO}_2)_2\text{C(OH)CO}_2\text{Me}$ [183°] Formed from methyl-di-nitro-phenyl-acetate and diazobenzene chloride (V. Meyer, *B* 22, 819) Alcoholic potash forms a blue solution, changing to

yellow, and forming $\text{C}_6\text{H}_3(\text{NO}_2)_2\text{C(CO}_2\text{H)NPhN}$ [272°], which yields a methyl ether [132°]

m NITRO-DIPHENYL-GUANIDINE

$\text{NH C(NHPh) NHC}_6\text{H}_4\text{NO}_2$ [182°] Formed from *m* nitro-di-phenyl thio urea, PbO , and alcoholic NH_3 (Brückner, *B* 7, 1236) Cyanogen, followed by dilute HClAq , changes it to the oxalyl derivative [168°], whence hot conc HCl forms nitro di phenyl parabanic acid (Hirsch, *C* 1888, 624)

m-Nitro-tri-phenyl-guanidine [159°]

Formed from *m* nitro-di phenyl thio urea, PbO , and aniline (*B*, Losanitsch, *B* 16, 50) Yellow plates — $\text{B'H}_2\text{P}^+\text{Cl}_4$

m Di-nitro di phenyl-guanidine

$\text{NH C(NH C}_6\text{H}_4\text{NO}_2)_2$ [190°] Formed from *m* nitro aniline and cyanogen chloride (Hofmann, *A* 67, 156), or from di nitro di phenyl thio urea, PbO , and alcoholic NH_3 (*B*) Scales $\text{B'HCl} - \text{B'H}_2\text{P}^+\text{Cl}_4$

Tri *m* nitro tri phenyl guanidine

$\text{C}_6\text{H}_3(\text{NO}_2)_3\text{N C(NH C}_6\text{H}_4\text{NO}_2)_2$ [189°] Formed from di nitro di phenyl thio urea, iodine, and alcohol (*L*) Yellow plates, sol hot alcohol

DI-NITRO DI-PHENYL HEPTANE

$\text{C}_6\text{H}_4(\text{C}_6\text{H}_4\text{NO}_2)_2$ Oil (Auger, *Bl* [2] 47, 42)

o-NITRO-PHENYL-HYDRAZINE

$\text{C}_6\text{H}_3(\text{NO}_2)\text{NH.NH}_2$ [90°] Formed from *o*-nitro diazobenzene chloride, HCl , and SnCl_4 at 0° (Bischler, *B* 22, 240, 2801) Brick red needles from benzene, al sol cold alcohol With benzaldehyde it yields $\text{C}_6\text{H}_3\text{CH.NH C}_6\text{H}_4\text{NO}_2$ [187°] — B'HSnCl_4 prisms — B'HCl needles — $\text{B'H}_2\text{SO}_4$ flesh-coloured needles

Formyl derivative

$\text{C}_6\text{H}_3(\text{NO}_2)\text{NH.NHCHO}$ [177°] Formed from *o* nitro phenyl hydrazine hydrochloride, formic acid, and some Na_2CO_3 Needles, v e sol hot Aq

Acetyl derivative [141°] Needles

Di-acetyl derivative [58°] Prisms

Benzoyl derivative [166°] Needles

Oxalyl derivative

$\text{C}_6\text{O}_2(\text{NH.NH C}_6\text{H}_4\text{NO}_2)_2$ Formed from the hydrazide and oxalic ether Yellow needles, sol hot nitrobenzene.

m-Nitro phenyl-hydrazine

$\text{C}_6\text{H}_3(\text{NO}_2)\text{NH.NH}_2$ [93°] Prepared in the same way as the *o*-compound (Bischler & Brodsky, *B* 22, 2809) Canary yellow needles Reacts with ketonic compounds, yielding their *m* nitro phenyl hydrazides, with the following melting points from aldehyde [98°], from acetone [112°], from benzoic aldehyde [118°], from acetophenone [160°], from benzal [158°]; and from aceto acetic ether [117°]

Salts — $\text{B'HCl} - \text{B'H}_2\text{SO}_4$ yellow crystalline groups, v sol hot water

Acetyl derivative [146°] Plates

Di-acetyl derivative [150°] Tables

(a)-*Acetyl-(B)* benzoyl derivative

$\text{C}_6\text{H}_3(\text{NO}_2)\text{NBz.NHAc}$ [137°] Formed from the acetyl derivative and Bz_2O at 160° Yellowish aggregates of crystals

(a)-*Benzoyl-(B)*-acetyl derivative

$\text{C}_6\text{H}_3(\text{NO}_2)\text{NAc.NHBz}$ [147°] Formed from the benzoyl derivative, Ac_2O , and NaOAc . Needles (by sublimation).

Benzoyl derivative [151°] Needles

Di-benzoyl derivative. [158°]. Plates.

Di-nitro di-phenyl-hydrazine

$C_6H_5NHNH.C_6H_5(NO_2)_2$ [1 2 4] [120°] Formed from $C_6H_5Cl(NO_2)_2$ and phenyl hydrazine (Willgerodt, *J pr* [2] 87, 350, 40, 252, 42, 132) Red plates Converted by shaking with HgO into $C_6H_5.N_2.C_6H_5(NO_2)_2$ [117°] On boiling with alcohol it yields $C_6H_5.N_2.C_6H_5(NO_2)_2$ [178°], boiling $HOAc$ forms $C_6H_5.N_2.C_6H_5(NO_2)_2$ [175°]

Di m nitro s di phenyl hydrazine

$(C_6H_5(NO_2))_2N_2H_2$ [220°] Formed from di-nitro azobenzene and cold alcoholic ammonium sulphide (Lermontoff, *B* 5, 236) Yellow needles

Tri nitro di phenyl-hydrazine

$C_6H_5NH.NHC_6H_5(NO_2)_2$ [1 2 4 6] [185°] Formed from $C_6H_5Cl(NO_2)_2$ and phenyl hydrazine hydrochloride (Willgerodt, *J pr* [2] 87, 346, 40, 264, Fischer, *A* 190, 132, 253, 1) Red crystals, melting at 175°-180° when slowly heated, but 183°-185° when quickly heated By boiling with $MeOH$ it is converted into the compound $C_6H_5.N_2.C_6H_5(NO_2)_2$ [218°] On heating with $HOAc$ it yields $C_6H_5.N_2.C_6H_5(NO_2)_2(NO)$ [248°] (Freund, *B* 22, 1663), which yields a mono sulphonic acid crystallising from water in yellow needles, not melted at 360°

o NITRO PHENYL-HYDRAZINE SULPHONIC ACID $C_6H_5(NO_2)(SO_3H)NHNH_2$ Formed from o nitro diazobenzene sulphonic acid and a well cooled, strongly acid, solution of $SnCl_2$ (Nietzki & Lerch, *B* 21, 3220) — $HA.HCl$

m Nitro phenyl hydrazine sulphonic acid [3 6 1] $C_6H_5(NO_2)(SO_3H)NHNH_2$ Formed from nitro diazobenzene sulphonic acid and cold $SnCl_2$ (Lamprecht, *B* 18, 2194) Yellow needles (containing aq) — KA' 1/4aq — BaA' 1/4aq — PbA' 4aq

NITRO PHENYL HYDROXYLAMINE v HYDROXYLAMINE DERIVATIVES

DI NITRO DI PHENYL HYPOPHOSPHOROUS ACID $(C_6H_5(NO_2))_2PO.OH$ [268°] Formed from Ph_2PO_2H , nitric acid, and H_2SO_4 (Dörken, *B* 21, 1513) Yellow pp, v sol water — NH_4A' [260°] — KA' 2aq — BaA' 6aq — PbA' — AgA'

NITRO PHENYL INDIAZINE CARBOXYLIC ACID $C_6H_5(NO_2) < \begin{smallmatrix} N(C_6H_5) \\ C(CO_2H) \end{smallmatrix} > N$ [272°]

Formed by the action of alcoholic potash on the red needles [183°] formed from di-nitro-phenyl acetic ether and diazobenzene (V Meyer, *B* 22, 319, *A* 264, 149) Sulphur yellow needles, v sl sol alcohol Reduced by $SnCl_2$ to a dihydride [235°]

Methyl ether MeA' [192°] Needles Converted by HNO_3 into $C_6H_5.N_2O_2$ [281°], and by H_2SO_4 into a sulphonic acid (Strassmann, *B* 23, 714)

Ethyl ether EtA' [158°] Needles

NITRO-DI-PHENYL-KETONE v NITRO-BENZOPHENONE

DI-NITRO-PHENYL-MALONIC ETHER $C_6H_5(NO_2)_2CH(CO_2Et)_2$ [51°] Formed from sodium malonic ether and $C_6H_5Br(NO_2)_2$ (Von Richter, *B* 21, 2472) Pale-yellow prisms.

m-NITROPHENYL MERCAPTAN

$C_6H_5(NO_2).SH$ Formed by the action of alcoholic potash on the ether formed from potassium xanthate and m-diazobenzene (Leuckart, *J pr* [2] 41, 197). Yellow liquid with nasty smell

p-Nitro-phenyl mercaptan $C_6H_5(NO_2).SH$ [77°] Formed from $C_6H_5Cl(NO_2)$ and alcoholic KSH (Willgerodt, *B* 18, 881). Crystals

Di nitro-phenyl mercaptan $C_6H_5(NO_2)_2SH$ [4 2 1] [181°] Formed from $C_6H_5Cl(NO_2)_2$ and KSH (Willgerodt, *J* 1884, 974) Needles

Ethers (Willgerodt, *B* 18, 330) MeA' [126°] — EtA' [113°] — PrA' [94°] — $PrOH_2A'$ [72°] — $C_6H_5CH_2A'$ [128°] — BzA' [113°]

Di-nitro-phenyl-mercaptan [195°] Formed by heating $C_6H_5(NO_2).SCN$ with conc H_2SO_4 (Austen & Smith, *Am* 8, 90) Yellow powder

Tri-nitro phenyl mercaptan $C_6H_5(NO_2)_3SH$ [6 4 2 1] [114°] Formed from $C_6H_5Cl(NO_2)_3$ and alcoholic KSH (W) Small yellowish needles Explodes at 115° — KA' brown needles

o-NITRO PHENYL-METHACRYLIC ACID $C_6H_5(NO_2).CH.CMe.CO_2H$ [165°] Formed by hydrolysis of its ether, which is prepared from methyl phenyl methacrylate and HNO_3 Formed also from phenyl isobutyric acid and HNO_3 (Edeleanu, *B* 20, 621, *C J* 53, 559) Crystalline powder, yielding o nitro benzoic acid on oxidation

m Nitro phenyl-methacrylic acid [197°] Formed from m nitro benzoic aldehyde by heating with sodium propionate and propionic anhydride (Von Miller, *B* 23, 1900) White powder, v sol hot alcohol

p-Nitro phenyl methacrylic acid [208°] Formed in the same way as the o isomeride (E) Crystals — AgA' white needles

Methyl ether MeA' [115°] Plates

m NITRO-PHENYL METHACRYLIC ALDEHYDE $C_6H_5(NO_2).CH.CMe.CHO$ [83°] Formed from m nitro-benzoic aldehyde, propionic aldehyde, and dilute (10 p c) aqueous $NaOH$ (Miller & Kinkelin, *B* 19, 530) Thin prisms (from alcohol) Aniline yields only $C_6H_5(NO_2).CH.NPh$ and crystalline $C_6H_5(NO_2).CH(NHPh)_2$ [170°] Tin and HCl yield a base $C_6H_5.N$ [98°], whence AcO forms $C_6H_5.AcN$ [145°], and benzoic aldehyde produces $C_6H_5.N(CHPh)$ [73°]

Phenyl hydrazide $C_6H_5.N_2O_2$ [135°] **NITRO DI PHENYL METHANE** $C_6H_5(NO_2)_2$

o compound Formed from o nitro benzyl chloride, benzene, and $AlCl_3$ (Geigy & Koenigs, *B* 18, 2402) Oil

m compound [141°] Formed from m nitro-benzyl alcohol and conc H_2SO_4 (Becker, *B* 15, 2091) Oil, sol alcohol

p compound [31°] Formed by either of the above methods (Basler, *B* 16, 2716, Manns, *C C* 1888, 1363) Prisms, v sol alcohol

m-Nitro-tri-phenyl methane

$CHPh_2.C_6H_5.NO_2$ [90°] Formed from m-nitro benzoic aldehyde, benzene, and H_2SO_4 (Tscha cher, *B* 19, 2463, 21, 188) Crystals (from ligroin)

p-Nitro-tri-phenyl-methane [93°] Formed in like manner (Baeyer & Löhr, *B* 23, 1622)

Di-nitro-di-phenyl-methane $C_6H_5(NO_2)_2$ By nitrating diphenyl methane Doer (*B* 5, 795) obtained two compounds of this formula, melting at 183° and 172°, while Staedel (*A* 194, 863) obtained the (a)- compound [188°] and a (β) isomeride [118°] By nitrating m-nitro diphenyl-methane Becker (*B* 15, 2092) obtained a fourth isomeride [94°], while Basler (*B* 16, 2719) got a fifth isomeride [175°] by the nitration of p-nitro-di-phenyl-methane

Tri-nitro-tri-phenyl-methane $CH(C_6H_5.NO_2)_3$ [207°] Formed by nitrating tri phenyl-methane

(E a O Fischer, *A* 194, 254, cf Hemilian, *B*, 7, 1208) Scales (from benzene)

Tetra-nitro-di-phenyl-methane [172°] Formed from CH_2Ph_2 and fuming HNO_3 at 0° (Staedel, *A* 218, 359) Long pointed needles

DI-NITRO-DI-PHENYL-METHYL-AMINE $\text{C}_6\text{H}_4(\text{NO}_2)\text{NMeC}_6\text{H}_5$ [167°] Formed from $\text{C}_6\text{H}_4\text{Cl}(\text{NO}_2)_2$ and $\text{C}_6\text{H}_5\text{NMe}$, or $\text{C}_6\text{H}_5\text{NHMe}$ (Leymann, *B* 15, 1235) Reddish needles

NITRO DI PHENYL-METHYL CARBINOL $\text{C}_6\text{H}_4(\text{NO}_2)\text{CPh}(\text{OH})\text{CH}_3$ [107°] Formed from *u*-di phenyl ethane and HNO_3 (Anschtz a Romig, *B* 18, 664) White prisms, yielding an acetyl derivative [86°]

***o*-NITRO PHENYL TRIMETHYLENE GLYCOL** $\text{C}_6\text{H}_3(\text{NO}_2)\text{CH}(\text{OH})\text{CH}_2\text{CH}_2(\text{OH})$ [109°] Formed from *o* nitro benzoic aldehyde, acetic aldehyde, and an alkali (Baeyer & Drewson, *B* 15, 2861) Colourless needles

NITRO PHENYL TRIMETHYLENYL KE TONE CARBOXYLIC ACID

$\text{C}_6\text{H}_3(\text{NO}_2)\text{CO C}(\text{CO}_2\text{H})\text{CH}_2\text{CH}_2$ [176°] Obtained from its ether, which is produced by the action of ethylene bromide on sodium *p* nitro benzoyl acetic ether (Perkin a Bellinot, *B* 18, 958) Colourless needles—AgA' amorphous *Ethyl ether* EtA' [84°] Prisms

NITRO-PHENYL-METHYL KETONE v NITRO-ACETOPHENONE

***p*-NITRO-PHENYL METHYL-OXAZOLE** $\text{CH C}(\text{C}_6\text{H}_4\text{NO}_2)\text{N}=\text{C}(\text{CH}_3)\text{CH}_2\text{O}$ [157°] Formed by the action of conc HNO_3 on the base obtained from acetamide and bromo acetophenone (Lewy, *B* 21, 926) Yellow needles, v sol hot alcohol Yields an amido-derivative [115°]

***p*-NITRO PHENYL (*β*)-METHYL PIPERIDINE** $\text{C}_6\text{H}_4(\text{NO}_2)\text{NC}_4\text{H}_8\text{Me}$ [61°] Formed by heating (*β*) methyl piperidine with $\text{C}_6\text{H}_4\text{Cl}(\text{NO}_2)_2$ at 150° (Lellmann a Buttner, *B* 23, 1389) Golden plates (from alcohol)—B'HAuCl₂ 2aq

Di-nitro phenyl (*β*) methyl piperidine [67°] Formed in like manner, using the compound $\text{C}_6\text{H}_4\text{Cl}(\text{NO}_2)_2$ [1 2 4] Yellow needles

***o*-NITRO DI PHENYL-METHYL-PYRAZOLE**

$\text{C}_6\text{H}_3(\text{NO}_2)_2\text{C}(\text{CH}_3)\text{N}(\text{NPh})\text{CHMe}$ [95°] or [105°] (285° at 70 mm) Formed by heating its carboxylic acid [218°] (Knorr a Jödicke, *B* 18, 2261) Indescent plates [95°], slender needles, or thick prisms [105°]—B'H₂PtCl₆ [198°]

***p* Nitro di-phenyl-methyl-pyrazole** $\text{C}_6\text{H}_3(\text{NO}_2)_2\text{C}(\text{CH}_3)\text{N}(\text{NPh})\text{CHMe}$ Formed by heating its carboxylic acid [202°] Oil—B'H₂PtCl₆ slender needles

Tri-nitro di phenyl methyl pyrazole $\text{C}_6\text{H}_2(\text{NO}_2)_3\text{C}(\text{CH}_3)\text{N}(\text{N(C}_6\text{H}_4\text{NO}_2)_2)\text{CHMe}$ [178°] Formed by nitrating di-phenyl methyl pyrazole (Knorr a Laubmann, *B* 22, 174) V sl sol alcohol

NITRO-DI-PHENYL-METHYL PYRAZOLE CARBOXYLIC ACID

$\text{C}_6\text{H}_3(\text{NO}_2)_2\text{C}(\text{CH}_3)\text{N}(\text{N(C}_6\text{H}_4\text{NO}_2)_2)\text{CHMe}$. The ethers of the *o*-acid [218°] and of the *p*-acid [202°] are respectively formed by heating *o*- and *p*-nitro-benzoyl-acetoacetic ether with phenyl-hydrazine in HOAc (Knorr a Jödicke, *B* 18, 2267) The *o*-ether melts at 146° and the *p*-ether at 128° Both crystallise well

***m*-NITRO-PHENYL-DI-METHYL-PYRIDINE DICARBOXYLIC ETHER** $\text{C}_{10}\text{H}_8\text{N}_2\text{O}_4$ $\text{C}_6\text{H}_4(\text{NO}_2)\text{C}_2\text{NMe}_2(\text{CO}_2\text{Et})_2$ [65°] Formed from its dihydride and fuming HNO_3 (Lepetit, *G* 17, 461) Colourless crystals (from alcohol)—B'H₂PtCl₆ [202°] Orange yellow needles—B'HNO₃ [130°] Colourless needles

NITRO PHENYL-DI-METHYL PYRIDINE DIHYDRIDE DICARBOXYLIC ETHER $\text{C}_6\text{H}_4(\text{NO}_2)\text{C}_2\text{NHMe}(\text{CO}_2\text{Et})_2$ Formed from nitro benzoic aldehyde, acetoacetic ether, alcohol, and NH₃ (Lepetit, *G* 17, 460, *B* 20, 1841) *o* isomeride [120°] Yellow tables *m*-isomeride [161°] Tables *p* isomeride [118°–122°]

***m* NITRO-PHENYL-METHYL-QUINOLINE**

$\text{C}_6\text{H}_4\text{CH}(\text{CMe})\text{C}_8\text{H}_6\text{N}_2$ [145°] Formed by heating aniline with *m* nitro α methyl cinnamic aldehyde and conc HClAq (Miller a Kinkelin, *B* 19, 531) Small plates, v sol hot alcohol—B'HCl—B'H₂PtCl₆ long needles

DI NITRO PHENYL (*α*) NAPHTHYLAMINE $\text{C}_{10}\text{H}_7\text{N}_2\text{O}_4$ $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{NHC}_6\text{H}_5$ [190 5°] Formed from (*α*) naphthylamine and $\text{C}_6\text{H}_4\text{Br}(\text{NO}_2)_2$ [72°] (Heim, *B* 21, 2302) Orange red needles (from alcohol) Conc H₂SO₄ forms a dark blue solution

Isomeride [77°] Prepared by nitration of phenyl (*α*) naphthylamine (Streiff, *B* 13, 1853) **Di nitro phenyl (*β*) naphthylamine** [169°] (*H*), [179°] (*E*) Prepared by the action of $\text{C}_6\text{H}_4\text{Br}(\text{NO}_2)_2$ or $\text{C}_6\text{H}_4\text{Cl}(\text{NO}_2)_2$ on (*β*) naphthylamine (Heim, *B* 21, 589, Ernst, *B* 23, 4429) Prisms By nitration of phenyl (*β*) naphthylamine Streiff obtained $\text{C}_{10}\text{H}_7(\text{NO}_2)_2\text{N}$ [85°] and $\text{C}_{10}\text{H}_7(\text{NO}_2)_2\text{N}$ [192°–195°], both crystalline

Tetra nitro phenyl naphthylamine $\text{C}_{10}\text{H}_5(\text{NO}_2)_4\text{NHC}_6\text{H}_5$ The (*α*) compound [162 5°] and the (*β*) isomeride [253°] are formed from aniline and the corresponding bromo tetra nitro naphthalenes (Merz a Weith, *B* 15, 2712) Both crystallise from benzene in orange needles (containing benzene)

DI-NITRO-PHENYL-(*α*) NAPHTHYL HYDRAZINE $\text{C}_{10}\text{H}_7\text{N}_4\text{O}_4$ [181°] Formed from (*α*) chloro di nitro benzene and (*α*) naphthyl hydrazine (Willgerodt, *J pr* [2] 43, 184) Rec prisms (*β*)-Naphthylamine yields an isomeride [188°]

Tri-nitro-phenyl-(*α*) naphthyl hydrazine $\text{C}_{10}\text{H}_5\text{N}_5\text{O}_6$ [176°] Got from an alcoholic solution of picryl chloride and (*α*) naphthyl hydrazine (*W*) Occurs in a red stable modification and an unstable yellow one (*β*)-Naphthylamine forms an isomeride [175°] also occurring in two forms

DI-NITRO-PHENYL-(*β*)-NAPHTHYL OXIDE $\text{C}_{10}\text{H}_7\text{O}_2$ $\text{C}_6\text{H}_4(\text{NO}_2)_2$ [1 2 4] [95°] Formed from $\text{C}_6\text{H}_4\text{Cl}(\text{NO}_2)_2$ and (*β*) naphthol (Ernst, *B* 23, 3430) Yellow needles (from alcohol)

NITRO-PHENYL NITRO-BENZYL OXIDE $\text{C}_6\text{H}_4(\text{NO}_2)\text{OCH}_2\text{C}_6\text{H}_4(\text{NO}_2)$ The compound formed from *p* nitro benzyl chloride, *o* nitro phenol, and alcoholic potash melts at 129°, the *pp* isomeride at 183° (Kumpf, *B* 17, 1077, *A* 224, 107)

Di-nitro-phenyl nitro benzyl oxide. [4 2 1] $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OCH}_2\text{C}_6\text{H}_4\text{NO}_2$ [1 4] [201°] an [6 2 1] $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{OCH}_2\text{C}_6\text{H}_4\text{NO}_2$ [1 4] [187°] have been prepared (Staedel, *B* 14, 899, Kumpf

They yield di nitro aniline and *p* nitro benzyl alcohol [91°] on heating with alcoholic ammonia

Tri nitro phenyl nitro benzyl oxide
[6 4 2 1] $C_6H_2(NO_2)_3 \cdot O \cdot CH_2 \cdot C_6H_4 \cdot NO_2$ [1 4] [108°]. Long thin needles (K)

DI NITRO-DI-PHENYL OXIDE
 $C_6H_4(NO_2)_2$ [185°] Formed by nitrating di-phenyl oxide (Hoffmeister, *A* 159, 191) Needles

Di nitro di phenyl oxide $C_6H_4(NO_2)_2 \cdot OC_6H_5$ [71°] Formed from $C_6H_5 \cdot Cl(NO_2)_2$ and $PhOK$ (Willgerodt) Needles

Tri nitro-di phenyl oxide
[4 2 1] $C_6H_4(NO_2)_3 \cdot O \cdot C_6H_4 \cdot NO_2$ [1 x] Formed from $C_6H_5 \cdot Cl(NO_2)_2$ and $C_6H_5 \cdot (NO_2)_2(OK)$ The *o* cern compound ($x=2$) melts at 119°, the *p* isomeride ($x=4$) at 114° (Willgerodt & Häetlin, *B* 17, 1764) Both are v sol benzene

Tetra nitro di-phenyl oxide
[6 4 2 1] $C_6H_2(NO_2)_4 \cdot O \cdot C_6H_4 \cdot (NO_2)_2$ [1 x] Formed from $C_6H_5 \cdot Cl(NO_2)_2$ and $C_6H_5 \cdot (NO_2)_2(OK)$ The *o* compound ($x=2$) melts at 173° and the *p*-isomeride at 153° (W & H) Both are crystalline

Tetra nitro di-phenyl oxide $\{C_6H_4(NO_2)_2\}_2 \cdot O$ [195°] Formed by the action of $C_6H_5 \cdot Cl(NO_2)_2$ on $C_6H_5 \cdot (OK)(NO_2)_2$ (Willgerodt, *B* 13, 887) Thick crystals, almost insol alcohol

NITRO-PHENYL-OXY-ACETIC ACID v NITRO MANDELIC ACID

p NITRO-PHENYL OXY BUTYL KETONE
ANHYDRIDE CARBOXYLIC ACID, $H_{11}(NO_2)_2 \cdot O$.

i.e. $O \leftarrow \begin{matrix} C(C_6H_4NO_2) \\ CH, CH, CH \end{matrix} \rightarrow CO \cdot OH$ Formed from trimethylene bromide and sodium *p* nitro benzyl acetic ether (Peikin, *jun*, *B* 18, 954, *C* *J* 51, 735) Melts at 172° when crystallised from benzene, 183° when crystallised from water — *AgA'* light yellow needles

Ethyl ether EtA' [63°] Lustrous yellow monoclinic crystals, $a \ b \ c = 2 \ 353 \ 1 \ 1 \ 553$, $\beta = 80^\circ 42'$

m-NITRO PHENYL PARACONIC ACID

$C_{11}H_9NO_4$ *i.e.* $C_6H_4(NO_2) \cdot CH \leftarrow \begin{matrix} CH(CO_2H) \\ CO \end{matrix} \rightarrow CH$.

[171°] Formed by heating *m* nitro benzoic aldehyde with Ac_2O and $NaOAc$ at 125° (Salo monson, *B* 18, 2154, *R* *T* *C* 6, 1) Crystals Yields with baryta the salt $C_{11}H_8NO_4 \cdot Ba$ The corresponding *p*-isomeride melts at 163° (*S*, cf Erdmann, *B* 18, 2742)

p NITRO-PHENYL PENTINOIC ACID

$C_9H_7(NO_2) \cdot CH \cdot CH \cdot CH \cdot CO \cdot H$ [271°] Formed from *p* nitro cinnamic aldehyde, Ac_2O , and $NaOAc$ (Einhorn & Gehrenbeck, *B* 22, 45, *A* 253, 357) Formed also by oxidising the ketone $C_9H_7(NO_2) \cdot CH \cdot CH \cdot CH \cdot CO \cdot Me$ with $NaOCl$ Yellowish needles (from alcohol) Forms a tetrabromide [254°] — *AgA'* flocculent pp

Ethyl ether EtA' [118°] Plates

o Nitro-di-phenyl pentinoic acid *Nitrolic* $C_6H_4 \cdot CH \cdot CH \cdot CH \cdot C(C_6H_4 \cdot NO_2) \cdot CN$ [206°] Formed from *p* nitro benzyl cyanide, cinnamic aldehyde, and $NaOEt$ (Rames, *B* 23, 3185) Needles

p NITRO-PHENYL PHENYL-(β)-AMIDO-PROPIONIC ACID

$C_9H_7(NO_2) \cdot CH(NHPh) \cdot CH_2 \cdot CO_2H$ [122°] Formed from aniline and $C_6H_4(NO_2) \cdot CHBr \cdot CH_2 \cdot CO_2H$ (Basler, *B* 17, 1500) Yellow crystals

Ethyl ether EtA' [78°] Crystals

DI-NITRO-PHENYL-PHENYLENE-DI-AMINE $C_6H_4(NO_2)_2 \cdot NH \cdot C_6H_4 \cdot NH_2$ [172°].

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Formed from $C_6H_5 \cdot Cl(NO_2)_2$ [1 2 4] and phenylene-diamine (Leymann, *B* 15, 1237)

NITRO-PHENYL PHOSPHATES The following compounds have been prepared by Rapp (*A* 224, 158), the NO_2 being in the *p*-position: $C_6H_4(NO_2) \cdot O \cdot PO(OH)_2$ [112°], $(C_6H_4(NO_2)_2)_2 \cdot PO \cdot OH$ [133°], $(C_6H_4(NO_2)_2)_2 \cdot PO \cdot OEt$ [135°], and $(C_6H_4(NO_2)_2)_2 \cdot PO$ [155°] Tri *o* nitro tri phenyl phosphate melts at 126° (Engelhardt & Lat schinoff, *Z* 1870, 230)

TRI-NITRO TRI-PHENYL PHOSPHINE OXIDE $OP(C_6H_4NO_2)_3$ [243°] Got by nitrating tri phenyl phosphine hydrate (Michaelis & Soden, *B* 17, 921, *A* 229, 324) It is accompanied by an isomeride [68°]

m NITRO PHENYL PHTHALIMIDE

$C_6H_4 \left\langle \begin{matrix} CO \\ C(C_6H_4NO_2) \end{matrix} \right\rangle$ [243°] Prepared by heating phthalic anhydride with *m* nitro aniline (Gabriel, *B* 11, 2261) Needles, v sol $EtOH$

NITRO PHENYL PIPERIDINE The following compounds are formed by the action of piperidine upon the corresponding halogen derivatives of nitro- and di nitro benzene respectively (Lellmann, *B* 20 680, 21, 2281) [1 4] $C_6H_4(NO_2)_2 \cdot NC_4H_9$ [81°] red prisms, yielding the salts $BHCl$ and $B \cdot H_2PtCl_6$ [1 2] $C_6H_4(NO_2)_2 \cdot NC_4H_9$ [105 5°] yellow plates, yielding the salts $BHCl$ and $B \cdot H_2PtCl_6$ [4 2 1] $C_6H_4(NO_2)_3 \cdot NC_4H_9$ [92°] orange needles

o NITRO PHENYL-PROPIOLIC ACID

$C_6H_4 \cdot NO_2$ *i.e.* $C_6H_4(NO_2) \cdot C \equiv C \cdot CO_2H$ Formed from $C_6H_4(NO_2) \cdot CHBr \cdot CHBr \cdot CO \cdot H$ (or its ether) and aqueous $NaOH$ (Baeyer, *B* 13, 2258, Muller, *A* 212, 127) Needles (from hot water), decomposing at 156° On boiling with water it gives *o* nitro phenyl acetylene Boiling alkalis yield *o*-nitro Reduction with glucose and alkalis produces indigo $FeSO_4$ also reduces it, in alkaline solution, to indigo white Its Na salt boiled with aqueous KCN and glucose yields indigo, even in presence of much HCN (Michael, *J* *pr* [2] 35, 254)

Ethyl ether EtA' [61°] Tables Converted by conc H_2SO_4 into isatogenic ether (*g* v)

p-Nitro-phenyl propiolic acid [181°] (*M*), [198°] (*D*) From $C_6H_4(NO_2) \cdot CHBr \cdot CHBr \cdot CO_2Et$ and alcoholic potash (*C* Muller, *A* 212, 127, Drewson, *A* 212, 154, Perkin, *C* *J* 49, 442). Yields *p* nitro phenyl acetylene and CO_2 on distilling with steam — *AgA'* amorphous powder

Ethyl ether EtA' [126°] Needles

o NITRO β PHENYL PROPIONIC ACID

$C_6H_4(NO_2) \cdot CH \cdot CH \cdot CO_2H$ [112°] Formed from (2, 4, 1) nitro amido phenyl propionic acid by elimination of NH_2 (Gabriel & Zimmermann, *B* 12, 600, 13, 1680) Yellow crystals — *AgA'*

m-Nitro-phenyl propionic acid [118°]

Formed in like manner from the (3, 4, 1)-nitro-amido-phenyl propionic acid (Gabriel, *B* 15, 845) Yellow needles, sl sol water

p Nitro- β -phenyl-propionic acid [164°]

Formed, together with the *o* acid, by nitration of (β) phenyl propionic acid (Glaser & Buchanan, *Z* 1869, 193, Beilstein & Kuhlberg, *A* 163, 132) — $CaA', 2aq$ — $BaA', 2aq$ small needles.

Ethyl ether EtA'

o-Nitro α phenyl-propionic acid

$C_6H_4(NO_2) \cdot CHMe \cdot CO_2H$ [110°] Formed, together with the *p*-isomeride [88°], by nitration

R R

of *o*-phenyl propionic acid (Trinius, *A* 227, 262) The *o* acid forms the salt $\text{CaA}', 2\text{aq}$, the *p*-acid gives $\text{CaA}', 2\text{aq}$ and $\text{BaA}', 2\text{aq}$

Di-nitro-phenyl propionic acid

$\text{C}_6\text{H}_3(\text{NO}_2)_2\text{CH}_2\text{CH}_2\text{CO}_2\text{H}$ [127°] Formed by nitration of β phenyl-propionic acid (G & Z)

Ethyl ether EtA' [32°] Needles

α -NITRO PHENYL PROPYLENE

$\text{C}_6\text{H}_4\text{CH}(\text{NO}_2)\text{CH}_3$ [64°] Formed from benzoic aldehyde, nitro ethane, and ZnCl_2 at 140° (Priests, *A* 225, 354), yellow needles

Di nitro phenyl propylene

$\text{C}_6\text{H}_3(\text{NO}_2)_2\text{CH}(\text{NO}_2)\text{CH}_3$ The *o* compound [77°] is formed, together with the *p* isomeride [116°], by nitrating α nitro phenyl propylene (T)

Di-nitro phenyl-propylene

$\text{C}_6\text{H}_3(\text{NO}_2)_2\text{CH}(\text{CHMe})$ [118°] Formed from $\text{C}_6\text{H}_5\text{CH}(\text{Me})\text{CO}_2\text{H}$ and HNO_3 (Edeleanu, *B* 20, 622) Yellowish needles

m-NITRO-PHENYL PYRIDYL-ETHYLENE

$\text{C}_6\text{H}_3(\text{NO}_2)\text{CH}(\text{CH}_2\text{C}_5\text{H}_4\text{N})$ [120°] Formed from *m* nitro benzoic aldehyde and methyl pyridine (Schuffan, *B* 23, 2716) Plates— $\text{B}'\text{H}_2\text{PtCl}_6$ [240°]—Mercury double salt $\text{B}'\text{HClHgCl}_2$ [211°]—Picrate yellow plates

TETRA-NITRO TETRA-PHENYL PYRROLE

$\text{NH}(\text{C}_6\text{H}_4\text{NO}_2)_4$ Formed from tetra phenyl pyrrole and HNO_3 (Fehrlin, *B* 22, 554) Yellow needles (from HOAc), decomposing at 123°

(α) NITRO (Py 1) PHENYL QUINOLINE

$\text{C}_6\text{H}_4\text{N}_2\text{O}_2$ [187°] Formed together with a smaller quantity of a (β) isomeride [118°] and a little of a (γ) isomeride [135°] by nitration of

(Py 1)-phenyl-quinoline $\text{C}_6\text{H}_4\text{N} \begin{smallmatrix} \text{CPh CH} \\ \text{N=CH} \end{smallmatrix}$ (Königs *a*. Net, *B* 20, 624) They are all crystalline

m-Nitro-(Py 3)-phenyl-quinoline

$\text{C}_6\text{H}_4\text{N} \begin{smallmatrix} \text{CH}_2\text{CH} \\ \text{N} \end{smallmatrix} \text{CO}_2\text{H}$ [124°] Obtained by heating *m* nitro-cinnamic aldehyde with aniline and HClAg at 140° (Miller & Kinkelin, *B* 18, 1900) White needles— $\text{B}'\text{HCl}$ — $\text{B}'\text{H}_2\text{PtCl}_6$

Tetrahydride $\text{C}_6\text{H}_4\text{N} \begin{smallmatrix} \text{CH}_2\text{CH} \\ \text{NH CH} \end{smallmatrix} \text{C}_6\text{H}_4\text{NO}_2$

[101°] Tables Yields a nitrosamine [71°]— $\text{B}'\text{HCl}$ silky needles

Nitro (B 2) phenyl-quinoline $\text{C}_{11}\text{H}_{10}\text{N}_2\text{O}_2$

[173°] Formed, as well as a di nitro derivative [208°], by the nitration of (B 2) phenyl-quinoline $\text{CPh CH} \begin{smallmatrix} \text{CH} \\ \text{CH} \end{smallmatrix} \text{C}_6\text{H}_4\text{N}$ (La Coste & Sorger, *A* 230, 28)

— $\text{B}'\text{H}_2\text{PtCl}_6$ yellow needles

DI-NITRO-DI-PHENYL-SUCCINIC ACID

$\text{C}_{12}\text{H}_8\text{O}_4(\text{NO}_2)_4$ The (α)- compound [226°] and its (β)-isomeride are formed by nitrating (α)- and (β)-di-phenyl succinic acid respectively (Reimer, *B* 14, 1804) Both give *p*-nitro benzoic acid on oxidation

DI-NITRO-DI-PHENYL-SULPHAZIDES

$\text{C}_{12}\text{H}_8(\text{NO}_2)_2\text{N}_2\text{SO}_2$ Compounds formed by the action of SO_2 , nitrous acid, and alcohol on the nitro-anilines (Limprioth, *B* 20, 1241). The *o*-, *m*-, and *p*-compounds melt at 150°, 182°, and 160° respectively They yield nitrogen, nitro benzene, and nitro benzene sulphonic acid on boiling with baryta

DI-NITRO-DI-PHENYL DISULPHIDE

$(\text{C}_6\text{H}_4\text{NO}_2)_2\text{S}_2$ [181°] (W), [170°] (L) Formed by oxidation of *p* nitro phenyl mercaptan (Willgerodt, *B* 16, 888), or by boiling with alcoholic

potash the product of the combination of potassium xanthate with *p* nitro diazobenzene (Leuckart, *J pr* [2] 41, 199) Prisms (from H_2OAc)

Tetra nitro-di-phenyl-sulphide

$\text{S}(\text{C}_6\text{H}_4(\text{NO}_2)_2)_2$ [193°] Obtained from [1° & $\text{C}_6\text{H}_4\text{Cl}(\text{NO}_2)_2$ and alcoholic KSH (Beilstein & Kurbatoff, *B* 11, 2056, Willgerodt, *B* 12, 768) Yellow needles An isomeric body [245°] is formed by the action of H_2SO_4 and fuming HNO_3 on $\text{C}_6\text{H}_3(\text{NO}_2)_3\text{SCN}$ (Austen & Smith, *Am* 8, 91)

Tetra nitro-di phenyl disulphide

$\text{S}_2(\text{C}_6\text{H}_4(\text{NO}_2)_2)_2$ Obtained by oxidising [4 2 1] $\text{C}_6\text{H}_3(\text{NO}_2)_3\text{SH}$ (Willgerodt *Bn* 2 527) Yellow needles, exploding at about 280°

Penta-nitro di-phenyl-sulphide

$\text{C}_6\text{H}_3(\text{NO}_2)_5$, S , $\text{C}_6\text{H}_4(\text{NO}_2)_4$ [217°] Formed from $\text{C}_6\text{H}_4\text{Cl}(\text{NO}_2)_2$, K S, and $\text{C}_6\text{H}_4\text{Cl}(\text{NO}_2)_2$ (W) Thick crystals (from HOAc)

Hexa nitro-di-phenyl sulphide

$\text{S}(\text{C}_6\text{H}_3(\text{NO}_2)_5)_2$ [226°] Formed from picrylchloride and K S (W) Golden plates (from HOAc)

DI NITRO-PHENYL SULPHOCYANIDE

$\text{C}_6\text{H}_3(\text{NO}_2)_2\text{SCN}$ [139°] Formed by heating [1 2 4] $\text{C}_6\text{H}_3\text{Br}(\text{NO}_2)_2$ with potassium sulphocyanide in MeOH (Austen & Smith, *Am* 8, 89) Buff yellow crystals (from chloroform)

NITRO DI PHENYL SULPHONE

$\text{C}_6\text{H}_5\text{SO}_2\text{C}_6\text{H}_4\text{NO}$ [92°] Formed by heating di phenyl sulphone with fuming HNO_3 (Gerike, *A* 100, 208) Minute crystals

Di nitro di phenyl sulphone

$(\text{C}_6\text{H}_4(\text{NO}_2)_2)_2\text{SO}$ [164°] (G), [197°] (S a. N) Formed by nitrating di phenyl sulphone (G) and by the action of SO_2 on nitro benzene (Schmid & Nötting, *B* 9, 79) Small tables

Tetra nitro di phenyl sulphone

$(\text{C}_6\text{H}_3(\text{NO}_2)_4)_2\text{SO}$ [241°] Prepared by oxidising the corresponding sulphide [193°] (Beilstein & Kurbatoff, *A* 197, 78) Yellowish prisms

p NITRO DIPHENYL *p* SULPHONIC ACID

$\text{C}_6\text{H}_4(\text{NO}_2)\text{C}_6\text{H}_4\text{SO}_3\text{H}$ Prepared by sulphonation of *p*-nitro diphenyl or by nitration of diphenyl *p* sulphonic chloride (Gabriel & Damberger, *B* 13, 1408)— NaA' — $\text{CuA}'_2, 4\text{aq}$ — $\text{BaA}'_2, 4\text{aq}$ small needles

Chloride $\text{C}_6\text{H}_4(\text{NO}_2)\text{SO}_2\text{Cl}$ [178°]

Amide $\text{C}_6\text{H}_4(\text{NO}_2)\text{SO}_2\text{NH}_2$ [228°]

Ethyl ether EtA' [169°]

Nitro diphenyl disulphonic acid

The chloride $\text{C}_{12}\text{H}_8(\text{NO}_2)_2(\text{SO}_2\text{Cl})_2$ [130°] is formed, together with $\text{C}_{12}\text{H}_8(\text{NO}_2)_2(\text{SO}_2\text{Cl})_2$ [166°], by nitration of diphenyl disulphonic chloride (G & D).

DI NITRO DI-PHENYL SULPHOXIDE

$(\text{C}_6\text{H}_4(\text{NO}_2)_2)_2\text{SO}$ [116°] Formed from di phenyl sulphoxide, NaNO_2 , and H_2SO_4 (Colby & McLoughlin, *Am* 9, 70, *B* 20, 198) Minute yellow crystals, v sol alcohol

NITRO PHENYL THIO CARBAMIC ACID

$\text{C}_6\text{H}_4(\text{NO}_2)\text{CS OH}$

Methyl ether MeA' The *m* compound [120°] is formed by boiling *m* nitro phenyl thio carbamide with MeOH (Steudemann, *B* 16, 551) Colourless needles

Ethyl ether EtA' The *m* compound [115°] and its *p*-isomeride [176°] are formed by boiling the corresponding nitro anilines with CS_2 , alcohol, and potash (Losantech, *B* 15, 470, 16, 49) Both are crystalline

m NITRO DI-PHENYL THIO-SEMICARBAZIDE

$\text{C}_6\text{H}_4(\text{NO}_2)\text{NH NH CS NHPh}$ [147°] Formed from phenyl thio carbamide and *m*-

nitro-phenyl hydrazine (Bischler & Brodsky, *B* 22, 2815) Dark yellow globular aggregates

m-NITRO-PHENYL THIO CARBIMIDE $C_6H_4(NO_2)NCS$ [61°] (*c* 277°) Formed by heating *m* nitro phenyl thio urea with As_2O_3 (Staudemann, *B* 16, 549, 2331) White needles

DI NITRO PHENYL THIOPHENE $C_6H_4(NO_2)_2C_2SH_2(NO_2)$ [178°] Formed from phenyl thiophene and fuming HNO_3 (Renard, *C R* 109, 699) Amorphous yellow powder

m NITRO PHENYL THIO UREA $C_6H_4N_2SO_2$ \cdot $\frac{1}{2}$ $C_6H_4(NO_2)NHCSNH_2$ [158°] Formed from *m* nitro phenyl thiocarbimide (Staudemann, *B* 16, 550) Lemon yellow crystals

m Nitro di phenyl thio urea $C_6H_4(NO_2)NHCSNHPh$ [156°] Formed from *m* nitro aniline and phenyl thiocarbimide (Bruckner, *B* 7, 1235, Losanitsch, *B* 14, 2365, Gebhardt, *B* 17, 3045) Small needles, sl sol cold alcohol

Di *m* nitro phenyl thio urea $CS(NH C_6H_4NO_2)_2$ [160°] Formed from *m* nitro aniline and *m* nitro phenyl thiocarbimide (Bruckner, *B* 6, 1103, *S*) Yellow crystals

o NITRO PHENYL *p* TOLUIDINE $C_6H_4(NO_2)NH C_6H_4$ [68°] Formed from *p*-toluidine and *o* bromo nitro benzene (Schöpf, *B* 23, 1842) Orange plates

Di nitro phenyl toluidine $[4 \frac{1}{2}] C_6H_4(NO_2)_2NHC_6H_4Me [1 \frac{1}{2}]$ Formed from toluidine and $C_6H_4Cl(NO_2)_2$ (Willgerodt, *B* 9, 980, Leymann, *B* 15, 1236) The *o* compound ($x=2$) melts at 102°, the *p* compound ($x=4$) at 137° The isomeric $PhNH C_6H_4(NO_2)_2Me$ formed from tri nitro toluene and aniline, melts at 142° (Hepp, *A* 215, 369)

DI NITRO PHENYL TOLYLENE DIAMINE $C_6H_4(NO_2)_2NH C_6H_4Me NH_2$ [184°] Formed from tolylene *m* diamine and $[1 \frac{1}{2}] C_6H_4Cl(NO_2)_2$ (Leymann, *B* 15, 1237) Red tables

Formyl derivative [157°]

Acetyl derivative [164°]

NITRO PHENYL TOLYL KETONE $C_6H_4(NO_2)O$ [127°] Formed by the action of HNO_3 on phenyl tolyl ketone and on phenyl *p* tolyl methane (Plascuda & Zinke, *B* 7, 983, Milne, *B* 5, 685) Flat plates (from alcohol)

Di nitro phenyl *m* tolyl ketone $C_6H_4(NO_2)_2O$ [145°] Formed from di nitro phenyl *m* tolyl methane [141°], $HOAc$, and CrO_3 (Senff, *A* 220, 236) Pointed needles (from alcohol) or short prisms (from $HOAc$)

Di nitro phenyl *p* tolyl ketone $C_6H_4(NO_2)_2CO C_6H_4(NO_2)$ [127°] Formed, as well as the tri nitro derivative [165°], by nitration of phenyl *p*-tolyl ketone (*Z* & *P*) Needles

m NITRO-PHENYL DI TOLYL METHANE $(C_6H_4)_2CH C_6H_4NO_2$ [85°] Formed from *m* nitro benzoic aldehyde, toluene, and H_2SO_4 (Tschacher, *B* 19, 2464, 21, 188) Crystals

Di nitro phenyl-tolyl methane $C_6H_4(NO_2)_2$ The three compounds of this composition got by nitrating phenyl *o*-, *m*-, and *p*-tolyl methane melt at 100°, 141°, and 137° respectively (Zinke, *B* 7, 986, Senff, *A* 220, 235) Tetra nitro-phenyl *p* tolyl methane melts at 161°

NITRO-PHENYL-*p*-TOLYL-THIO-UREA $C_6H_4N_2SO_2 \cdot \frac{1}{2}$ $C_6H_4NHCSNH C_6H_4NO_2$ [148°] Formed from (2,4,1) nitro toluidine and phenyl thiocarbimide (Staudemann, *B* 16, 2336) Crystals Melts, after one fusion, at 167°

m-Nitro phenyl-*p* tolyl thio-urea $C_6H_4(NO_2)NHCSNH C_6H_4$ [173°] Formed from *m* nitro phenyl thiocarbimide and *p* toluidine (*S*) Needles, sl sol ether

Di nitro phenyl *p* tolyl thio urea $C_6H_4(NO_2)_2NHCSNH C_6H_4NO_2$ [188°] Formed from *m* nitro phenyl thiocarbimide and (2,4,1)-nitro toluidine (*S*) Crystals, sl sol alcohol

m NITRO PHENYL UREA $C_6H_4N_2O_2 \cdot \frac{1}{2}$ $C_6H_4(NO_2)NHCO NH_2$ Formed from *m* nitro aniline and cyanogen chloride (Hofmann, *A* 67, 156, 70, 137) Yellow needles (from water)

m Nitro di phenyl urea $C_6H_4NHCO NHC_6H_4NO_2$ [197°] (*G*), [187°] (*B*) Formed from *m* nitro di phenyl thio-urea and PbO (Brückner, *B* 7, 1236) Formed also by boiling with benzene the compound $PhNHCO N(C_6H_4NO_2)N NPh$ [104°] obtained by the action of phenyl cyanate on $(C_6H_4NO_2)NH N NPh$ (Goldschmidt, *B* 21, 2578) Yellow needles

p Nitro di phenyl urea [202°] Formed by the action of phenyl cyanate on a solution of *p*-nitro diazoamidobenzene in benzene (*G*) Yellow crystals (from alcohol)

m Nitro tri phenyl-urea $NPh.CO NHC_6H_4NO_2$ [155°] Formed from *m* nitro aniline and $NPh.COCl$ (Lellmann & Bonhöffer, *B* 20, 2121) Yellow needles The isomeride prepared from *p* nitro aniline melts at 176°, and crystallises in bluish green tables

Di *m* nitro di phenyl urea $CO(NH C_6H_4NO_2)_2$ [233°] Formed from the thio urea and PbO (*B*) Yellow needles (Losanitsch, *B* 16, 50)

Tetra nitro di phenyl urea $CO(NH C_6H_4NO_2)_2$ Formed by nitration of di phenyl urea (Losanitsch, *B* 10, 690, 11, 1539) Yellow needles, melting above 200° (*L*) or at 189° (Hentschel, *J pr* [2] 34, 426) The green *K* salt $C_6H_4K_2N_4O_6$ explodes on heating

p NITRO PHENYL VALERIC ACID $C_6H_4(NO_2)CH_2CH_2CH_2CO_2H$ Formed by heating *p* nitro benzyl ethyl malonic ether with aqueous KOH (Lellmann & Schleich, *B* 20, 438) Sparingly soluble powder, carbonising above 300°

p NITRO PHENYL VINYL MALONIC ACID $C_6H_4(NO_2)CH=CHCH(CO_2H)_2$ [208°] Formed from *p* nitro cinnamic aldehyde, malonic acid, and $HOAc$ (Einhorn & Gehrenbeck, *B* 22, 45)

NITRO PHLOROGLUCIN $C_6H_4(NO_2)(OH)$ Formed from phloroglucin and dilute HNO_3 (Hlaswetz & Pfaunder, *A* 119, 199) Reddish yellow scales, sl sol water

Tri nitro phloroglucin $C_6(NO_2)_3(OH)$ [158°] Formed from tri nitroso phloroglucin, HNO_3 , and H_2SO_4 (Benedikt, *B* 11, 1376) Hexagonal crystals (containing aq) Its salts are explosive and dye yellow With KOy it gives the isopurpuric acid reaction $-KHA'' aq - KHA'' - K_2A'''$

NITRO PHTHALIC ACID $C_6H_4(NO_2)(CO_2H)$ [32 1] Mol w 211 [220°] *S* ($HOAc$) 75 at 26° (Aguar, *B* 5, 899)

Formation—1 By boiling naphthalene with HNO_3 (Mangnac, *A* 38, 7, Laurent, *A* 41, 110)—2 By nitration of phthalic acid (Hugo Muller, *Z* 1868, 257, Faust, *A* 160, 57, Milner, *A* 208, 224)—3 By oxidation of nitro-naphthalene or (a) di nitro naphthalene (Beilstein & Kurbatoff, *B* 12, 688, *C O* 1881, 859, *A* 202, 217, Guareschi, *B* 10, 294)

Properties.—Yellow crystals, v sol hot

water. Splits up below its melting point into water and anhydride when slowly heated

Salts — K_2A' aq — KHA'' aq — $(NH_4)_2A''$ — $(NH_4)HA'$ 2aq — BaA'' 2aq — ZnA'' 1½aq — PbA'' 1½aq — Ag_2A'' white powder

Mono ethyl ether EtA'' [111°]

Di-ethyl ether Et_2A'' [45°]

Anhydride $C_6H_4(NO_2)(CO_2H)_2$ [164°] (Graeff, *B* 15, 1127)

Nitro-phthalic acid $C_6H_4(NO_2)(CO_2H)_2$ [4 2 1] [161°]

Formation — 1 Together with the preceding isomeride, by nitration of phthalic acid (O Miller, *A* 208, 224) — 2 By the oxidation of di-nitro-(β)-naphthol (Graebe & Drews, *B* 17, 1171)

Properties — Small pale yellow needles (containing aq) Resolved at 165° into water and its anhydride

Salts — K_2A'' — BaA'' 2aq — Ba_2H_2A'' — Zn_2H_2A'' 2aq — Ag_2A'' long colourless needles

Mono ethyl ether EtHA'' [128°]

Di-ethyl ether Et_2A'' [33°]

Anhydride $C_6H_4(NO_2)_2$ [114°]

Di-nitro phthalic acid $C_6H_4(NO_2)_2(CO_2H)_2$ [5 3 2 1] [226°] Formed by the oxidation of

β - di-nitro naphthalene or tetra nitro (α)-naphthol with dilute HNO_3 (Beilstein & Kurbatoff, *B* 13, 354, *A* 202, 225, Merz & Weith, *B* 15, 2728) Prisms — CaA'' — BaA''

Mono-ethyl ether EtHA'' [187°]

Di-nitro phthalic acid $C_6H_4(NO_2)_2(CO_2H)_2$ [6 3 2 1] [200°] Formed by heating β - bromo

tetra nitro naphthalene with dilute HNO_3 (Meiz & Weith, *B* 15, 2728) Needles — BaA''

Di-nitro phthalic acid $C_6H_4(NO_2)_2(CO_2H)_2$ Formed from nitro anisic acid, HNO_3 , and H_2SO_4 (Engelhardt & Latschnoff, *Z* [2] 7, 262) Tables [from water] — BaA'' nearly insol water

Nitro isophthalic acid $C_6H_4(NO_2)(CO_2H)_2$ [5 3 1] [249°] *S* 146 at 15°, 171 at 16°, 81.1 at 99° Prepared, together with an isomeride [260°] by heating isophthalic acid (100 g)

with fuming HNO_3 (1 kilo) for 24 hours (Beyer, *J pr* [2] 22, 35, 25, 473, cf Storrs & Fittig, *A* 158, 285) Plates (containing 1½ aq)

Salts — K_2A'' 1½aq *S* (80 p.c. alcohol) 744 at 15° — Na_2A'' aq *S* (80 p.c. alcohol)

32 at 15° Explodes above 160° — $(NH_4)HA''$ — MgA'' 5aq *S* 2 16 at 15° — CaA'' 3½aq *S* 72 at 15°

Turned violet by light — SrA'' 4½aq *S* 47 at 15° — BaA'' 2½aq *S* 85 at 15° — ZnA'' aq *S* 55 — CdA'' 2aq *S* 75 at 15° — Pb_2OA'' — Cu_2OA'' — MnA'' 5aq *S* 2 44 at 15°

— Fe_2O_4A'' — CoA'' 4½aq *S* 2 16 at 15° — NaA'' 4½aq *S* 2 74 at 15° — Ag_2A'' Explodes above 100°

Methyl ether Me_2A'' [122°]

Ethyl ether Et_2A'' [84°]

Nitro-isophthalic acid $C_6H_4(NO_2)(CO_2H)_2$ [4 3 1] [246°] (*G*), [259°]. Formed by oxidation of nitro-xylene (Claus, *J pr* [2] 38, 318, cf Wroblewsky, *Bl* [2] 34, 332, Noyes, *Am* 10, 472) White needles (containing 3aq), m sol

cold water — BaA'' 4aq (*G*). — BaA'' 1½aq (*N*) — CaA'' ½aq — MgA'' 6aq — PbA'' 3aq (*W*) — Ag_2A'' 7½aq pearly plates

Di-nitro-isophthalic acid $C_6H_4(NO_2)_2(CO_2H)_2$ [215°] Formed from isophthalic acid and fuming HNO_3 at 180° (Claus) Needles (containing 5aq) — Na_2A'' 2aq — K_2A'' 2aq — BaA'' 7aq — CaA'' 4aq — MgA'' 4aq; needles, v sol water.

NITRO-PHTHALIDE

$[4 \frac{1}{2}] C_6H_4(NO_2) < \begin{smallmatrix} CO \\ CO \end{smallmatrix} > O$ [141°] Formed

by nitration of phthalide (Hoening, *B* 18, 3447) Long needles Aqueous KOH yields $C_6H_4(NO_2)(CH_2OH)(CO_2K)$ CrO_3 in $HOAc$ oxidises it to nitrophthalic acid [161°] The acid of which it is the anhydride melts at 129° An isomeric nitro-phthalide [136°] is formed by oxidising (α) nitro naphthalene

NITRO-DIPHTHALYL $C_{12}H_8NO_4$ 1c

$C_6H_4C_6H_4O_2C_6H_4NO_2$ [270°] Formed by heating nitro phthalide with phthalic anhydride and $NaOAc$ (Graebe & Guye, *A* 233, 244) Yellow needles (from $HOAc$)

NITRO-PIPERIDINE $C_6H_8N(NO_2)$ (245°)

Formed by treating piperyl urea with HNO_3 (*S G* 1 5) at — 10°, pouring upon sodium carbonate and extracting with ether (Franchmont & Klobbie, *R T C* 8, 302) Colourless liquid, solidifying below — 10° Volatile with steam

NITROPODOCARPIC ACID v *PODOCARPIC ACID*

NITRO-PROPANE $C_3H_7NO_2$ 1c *PrNO_2*

(127°) (*M* & *R*), (131°) (*P*) *S G* 1 10108, 2 10023 *MM* 3 819 (Perkin, *C J* 55, 689)

Formed, together with propyl nitrite, by the action of silver nitrite on propyl iodide (V Meyer & Rilliet, *B* 5, 1029, *A* 171, 36, Prubram & Handl, *M* 2, 653, Cahours, *C R* 77, 749) Oil — $NaC_3H_6NO_2$ white powder

Iso nitro propane $PrNO_2$ (c 117°) Formed,

together with an isomeride (44°), by the action of silver nitrite on isopropyl iodide (V Meyer & Locher, *B* 7, 670, *A* 171, 39, Kiesel, *J R* 16, 135, *Bl* [2] 40, 72, *Bn* 1, 225) Oil, decomposed by $HClAq$ at 100°

Di nitro propane $CH_3CH_2CH_2(NO_2)_2$ (189° cor) *S G* 2 1258 Formed from bromo-

nitro propane and KNO_3 (Ter Meer, *A* 181, 19), and by the oxidation of di propyl ketone (Chan cel, *C R* 96, 1466, Kurtz, *A* 161, 208) Oil — KA'' — AgA'' explosive laminae

Iso-di nitro propane $(CH_3)_2C(NO_2)_2$ [58°]

(187° uncor) Formed by oxidation of propyl-

pseudonitrole $(CH_3)_2C(NO_2)(NO_2)$ (Meyer & Locher, *B* 7, 1613) Formed also by the action

of nitric acid on isobutyric and isovaleric acids (Bredt, *B* 15, 2822) White crystals, volatile with steam

NITRO PROPENYL BENZOIC ACID

$C_6H_4(NO_2)(CH_2)CO_2H$ [3 4 1] [155°] Formed

by boiling nitro oxypropyl benzoic acid with aqueous HCl (*S G* 1 10) for a long time (Wid man, *B* 15, 2551, 16, 2569) Short needles — NH_4A'' — CaA'' 2aq *S* 555 at 16° — BaA'' 3½aq

S 425 at 18° — CuA'' aq — AgA'' slender needles

β NITRO-PROPIONIC ACID $C_3H_5NO_4$ 1c

$CH_3(NO_2)CH_2CO_2H$ [67°] Formed from β

iodo propionic acid and $AgNO_3$ (Lewkovitch, *J pr* [2] 20, 165) Scales (from chloroform)

Ethyl ether EtA'' (c 168°) *V D* 4 85 (calc 5 69)

DI-NITRO-PROPYL-ANILINE $C_6H_4N_2O_4$ 1c

$C_6H_4(NO_2)_2NHCH_2CH_2CH_2$ [4 2 1] [95°] Formed from propylamine and $C_6H_5Br(NO_2)$, or by oxidising

the following body (Van Romburgh, *R T C* 4, 191, 8, 252) Yellow needles

Di nitro di propyl aniline

$C_6H_4(NO_2)_2N(C_2H_5)_2$ [4 2 1]. [40°] (Romburgh, *R. T. C.* 6, 252).

Tri-nitro-propyl-aniline $C_6H_4(NO_2)_3.NH.C_3H_7$, [59°] Formed from $NH_4C_3H_7$ and $C_6H_5Cl(NO_2)_3$. Yields with HNO_3 the nitramine $C_6H_4(NO_2)_3.N(NO_2)C_3H_7$, [97°]

NITRO-ISOPROPYL BENZENE v **NITRO-CUMENE**

Di nitro p di propyl-benzene $C_6H_4Pr_2(NO_2)_2$, [65°] Formed from di propyl benzene and fuming HNO_3 (Korner, *B* 11, 1865, *A* 216, 226) Plates, volatile with steam

Tri nitro m di-isopropyl benzene $C_6H_3Pr_3(NO_2)_3$, [111°] Yellow needles (Uhlhorn, *B* 23, 3142)

NITRO-PROPYL-BENZOIC ACID v **NITRO-n CUMINIC ACID**

NITRO ISOPROPYL CINNAMIC ACID [4 2 1] $C_6H_4Pr(NO_2).CH.CH.CO.H$ 'Nitro cumenyl acrylic' acid [157°] Formed by nitration of propyl cinnamic acid (Einhorn *A* Hess, *B* 17, 2016, Widman, *B* 19, 258) Needles, v sol alcohol Yields o nitro cuminic aldehyde on oxidation by $KMnO_4$.

Nitro isopropyl cinnamic acid

[4 3 1] $C_6H_4Pr(NO_2).C_3H_7.CO.H$ [141°] Formed by heating m nitro cuminic aldehyde (4 pts) with Ac_2O (5 pts) and $NaOAc$ (3 pts) at 175° for 4 hours (Widman, *B* 19, 413) Tables Forms a dibromide [184°] — KA' (dried at 100°) — NaA' 3aq — BaA' 54aq — CaA' 3aq *Ethyl ether* EtA' [59°] Tables.

Nitro-n propyl-cinnamic acid

[4 2 1] $C_6H_4Pr(NO_2).C_3H_7.CO.H$ [123°] Formed, in small quantity, in the nitration of isopropyl cinnamic acid (Widman, *B* 19, 273) On oxidation by alkaline $KMnO_4$, it yields nitro n cuminic acid and nitro cuminic acid Bromine forms a di bromide [171°]

DI NITRO PROPYL-THIOPHENE

$C_6HPr_2(NO_2)_2$ S Formed by nitration (Ruff, *B* 20, 1712) Oil

NITRO PROTocatechuic ACID $C_6H_5NO_3$.

Methyl derivative

$C_6H_4(NO_2)(OMe)(OH).CO.H$ [x 3 4 1] Formed by boiling its acetyl derivative [182°] which is produced by the nitration of acetyl vanillic acid $C_6H_3(OMe)(OAc)CO.H$ (Tiemann *A* Matsumoto, *B* 9, 945, 11, 132) Needles (from alcohol)

The isomeric $C_6H_4(NO_2)(OMe)(OH).CO.H$ [5 3 4 1] [202°] is formed by oxidation of acetyl nitro eugenol (Weselsky *A* Benedikt, *M* 3, 392), and crystallises in yellow needles

Methyl derivative

$C_6H_4(NO_2)(OMe)(OH).CO.H$ [6 4 3 1] [173°] Formed from its acetyl derivative [169°], which is got by nitrating acetyl isovanillic acid (T *A* M) Needles

Di methyl derivative $C_6H_4NO_3$ v $C_6H_4(NO_2)(OMe).CO_2H$ Formed by nitration of veratric acid $C_6H_3(OMe)_2.CO_2H$ (T *A* M, Merck, *A* 108, 59) Yellow needles (containing 3aq) Yields the ethers MeA' [144°] and EtA' [100°]

Iso-nitro-protocatechuic acid *Di-methyl ether* $C_6H_4(NO_2)(OMe)_2.CO_2H$ [202°] Formed by methylation of nitro-vanillic acid (T *A* M) Needles Yields MeA' [126°] crystallising in needles

Nitro protocatechuic acid *Methyl-propyl ether* $C_6H_4H_2(NO_2)(OMe)(OPr).CO_2H$ Formed by nitration (Cahours, *Bl* [2] 29, 270)

Nitro-protocatechuic acid *Methylene ether* $CH_2O.C_6H_4(NO_2)CO_2H$. [172°] Formed

from piperonylo acid $CH_2 \begin{smallmatrix} \diagup O \diagdown \end{smallmatrix} C_6H_4.CO_2H$ by nitration (Jobst *A* Hesse, *B* 11, 1031, *A* 199, 70) Needles — KA' 3aq — PbA' 4aq — CuA' 4aq. — AgA' needles or plates

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NITROPYROCATECHIN $C_6H_4NO_3$ v $C_6H_4(NO_2)(OH)$, [4 2 1] [170°] Formed by the action of nitrous acid on pyrocatechin (Benedikt, *B* 11, 362, *J pr* [2] 18, 455) Formed also by boiling the carbonyl derivative of (5,2,1) nitro amido phenol with potash (Von Chelmecki, *J pr* [2] 42, 442) Small yellow needles (from ether) Its aqueous solution is turned purple by alkalis Dyes stuff mordanted with alumina orange (Kostanecki, *B* 22, 1347) — BaA' 3aq dark red plates, with metallic lustre

An isomeric nitropyrocatechin [86°] is formed, together with the preceding, by nitration of pyrocatechin (Weselsky *A* Benedikt, *M* 3, 386) *A di methyl derivative* of nitro pyrocatechin $C_6H_4(NO_2)(OMe)_2$ [96°] is formed by nitrating veratrole (Merck, *A* 108, 60, The mann *A* Matsumoto, *B* 9, 939, 11, 131)

The *methylene derivative*

$C_6H_4(NO_2) \begin{smallmatrix} \diagup O \diagdown \end{smallmatrix} CH$ [148°] is a product of the action of nitric acid on piperonylo acid (Hesse, *A* 199, 73, 341) It crystallises in needles

Di nitro pyrocatechin $C_6H_4(NO_2)_2(OH)_2$ *Methyl derivative* $C_6H_4(NO_2)_2(OMe)(OH)$ *Di nitro guaiacol* [123°] Made by the action of nitrous acid gas upon an ethereal solution of guaiacol at 0° (Herzig, *M* 3, 825) Plates The *di methyl derivative* $C_6H_4(NO_2)_2(OMe)_2$, formed by nitration of veratrol, melts above 100° (M) The *methylene derivative* $C_6H_4(NO_2)_2O.CH_2$, [101°] is formed in the nitration of piperonylo acid (H)

Tri nitro-pyrocatechin *Di methyl-derivative* $C_6H_4(NO_2)_3(OMe)_2$, [145°] Got by nitrating $C_6H_4(NO_2)(OMe)_2$ (T *A* M) Prisms

DI NITRO-PYROCOLL $C_6H_4(NO_2)_2N_2O_2$ Got by nitrating pyrocoll (Ciamian *A* Danesi, *G* 12, 39) Yellow crystals, decomposing before fusion

NITRO-PYROGALLOL $C_6H_2(NO_2)(OH)_3$, [205°] Got by passing nitrous fumes into an ethereal solution of pyrogallol (Barth, *M* 1, 882) Trichmic olive brown prisms (containing aq), $a b c = 2.842.1.493$ $C_6H(NO_2)(OEt)_2(OH)$ [123°] and $C_6H(NO_2)(OEt)(OH)$, [139°] are formed in the same way (Weselsky *A* Benedikt, *M* 2, 214) The compounds $C_6H(NO_2)(OEt)$, [73°] and $C_6(NO_2)_2(OEt)$, [93°] are formed by nitration

NITROPYROMECONIC ACID $C_6H_4(NO_2)O$, Formed by nitrating pyromeconic acid (Ost, *J pr* [2] 19, 192) Crystals (from alcohol) — NaA' — AgA'

DI NITRO PYROMELLITIC ACID

$C_6(NO_2)_4(CO_2H)_2$, [5 2 6 4 3 1] Formed by oxidation of di nitro ψ cuminic acid $C_6Me_2(NO_2)_2.CO_2H$ [205°] (Nef, *C J* 53, 428, *A* 258, 317) Long silky needles — $Ag.A'$ amorphous pp

Methyl ether $Me.A'$ [180 6°]

Ethyl ether $Et.A'$ [180°]

NITRO-PYROMUCIC ACID $C_6H_4NO_3$ v $C_6H_4(NO_2)O.CO.H$ [184°] Formed from dehydromucic acid (1 pt), HNO_3 (10 pts), and conc H_2SO_4 (1 pt) (Klinkhardt, *J pr* [2] 25, 51) Got also from $C_6H_4(NO_2)O.CH_2.CH(NO_2)$ by

oxidation with CrO_3 (Priests, *B* 18, 1862) Yellow plates (from water) Yields succinic acid (and not an amido-acid) on reduction with tin and HCl aq— CaA'_2 — PbA'_2 — AgA'

Ethyl ether EtA'_2 [101°]

DI-NITRO-PYRROLE $\text{C}_4\text{H}_4(\text{NO}_2)_2\text{NH}$ [152°]

Formed by the action of fuming HNO_3 on pyrrol methyl ketone Formed also, together with an isomeride [178°], by the action of fuming HNO_3 on pyrrole carboxylic acid at 0° (Ciamcian a Silber, *B* 18, 1462, 19, 1081, *G* 16, 347) Colourless plates BaA'_2 , yellow needles

NITRO PYRROLE CARBOXYLIC ACID

$\text{C}_4\text{H}_3(\text{NO}_2)\text{NH}(\text{CO}_2\text{H})$ [217°] Formed by saponifying its methyl ether, which is got by nitrating $\text{C}_4\text{H}_4\text{NH}(\text{CO}_2\text{Me})$ (Anderlini, *B* 22, 2505, *Rend Accad Linc* [5] 1, 40) Yellow needles (containing aq), sl sol cold water

Methyl ether MeA' [197°]

An isomeride acid [161°] may be obtained from its methyl ether [179°] which accompanies the preceding ether $\text{C}_4\text{H}_4(\text{NO}_2)_2\text{NH}(\text{CO}_2\text{Me})$ [115°] is also formed in the nitration

Nitro-pyrrole carboxylic acid [146°] Formed by boiling di nitro pyrrole with potash solution (Ciamcian a Danesi, *G* 12, 40) Minute needles (containing aq)— $\text{NH}_4\text{A}'$ prisms or scales

NITRO PYRROLE-DI-METHYL DIKE

$\text{C}_4\text{H}_2(\text{NO}_2)_4$ *see* $\text{C}_4\text{H}_2(\text{NO}_2)_2\text{N}(\text{CO}_2\text{CH}_3)_2$ [149°] Formed by nitrating pyrrole di methyl di ketone (Ciamcian a Silber, *G* 16, 347, *B* 18, 1467, 19, 1078) Needles (from water)

NITRO PYRROLE METHYL KETONE By

nitrating $\text{NH}-\begin{array}{c} \text{CH CH} \\ \text{C=C} \end{array}$ two compounds are formed [197°] and [156°] Both yield pps of $\text{C}_4\text{H}_2\text{N}_2\text{O}_4\text{Ag}$ (Ciamcian a Silber, *B* 18, 1413, 1457) A compound $\text{C}_4\text{H}_2(\text{NO}_2)_2\text{N CO CH}_3$ [114°] crystallising in yellow needles (containing aq) may also be obtained

(*B* 1) **NITRO QUINOLINE** $\text{C}_8\text{H}_6\text{N}_2\text{O}_2$ *see* $\text{CH C}(\text{NO}_2)\text{C CH CH}$ [72°] Formed, together with the (*B* 4) isomeride, by the nitration of quinoline, especially in presence of fuming H_2SO_4 , in the cold (Claus a Kramer, *B* 18, 1243, Noeltling a Trautmann, *B* 23, 8654) Colourless needles (containing aq)

(*B* 2) **Nitro-quinoline** $\text{NO}_2\text{C}-\begin{array}{c} \text{CH C} \\ \text{CH C} \end{array}\text{CH CH}$ [150°] (La Coste, *B* 16, 669), [164°] (*C* a *K*) Formed by boiling *p* nitro aniline (25 pts), glycerin (60 pts), nitro benzene (15 pts), and H_2SO_4 (50 pts) for 4 hours Needles (containing aq)— $\text{B}'\text{H}_2\text{PtCl}_6$, small yellow needles

Methyl iodide $\text{B}'\text{MeI}$ Needles

(*B* 3) **Nitro-quinoline** $\text{CH CH C}(\text{NO}_2)\text{C CH CH}$ [181 5°] Formed from *m* nitro aniline, picric acid, glycerin, and H_2SO_4 (Claus a Stiebel, *B* 20, 3095) Needles— $\text{B}'\text{HCl}$ [225°]— $\text{B}'\text{HNO}_3$,— $\text{B}'\text{H}_2\text{PtCl}_6$, prisms

(*B* 4) **Nitro-quinoline** $\text{CH CH}-\begin{array}{c} \text{CH C} \\ \text{CH C} \end{array}\text{CH CH}$ [89°] Formed by nitrating quinoline, and also by boiling *o* nitro-aniline with glycerin, nitrobenzene, and H_2SO_4 (Kömgas, *B* 12, 449, La Coste, *B* 16, 678, Claus, *B* 18, 1243, Noeltling, *B* 23, 8654) Formed also by heating quinoline (*B* 4)-sulphonic acid with HNO_3 (Claus a

Küttner, *B* 19, 2886), and by warming $\text{C}_8\text{H}_6(\text{NO}_2)(\text{OMe})\text{CH CH CHO}$ with alcoholic ammonia (Miller a Kinkelin, *B* 22, 1716) Prisms— $\text{B}'\text{H}_2\text{PtCl}_6$, orange needles

(*B* 2, 4) **Di-nitro quinoline**

$\text{C}(\text{NO}_2)_2\text{CH C CH CH}$ [150°] Formed by $\text{CH C}(\text{NO}_2)\text{C N CH}$ heating (4, 2, 1) di nitro aniline with glycerin, nitro benzene, and H_2SO_4 (La Coste, *B* 15, 562) Long slender needles

By the nitration of quinoline two isomeric di nitro-quinolines [183°] and [134°] may be got (Claus a Kramer, *B* 18, 1243) Their platino chlorides $\text{B}'\text{H}_2\text{PtCl}_6$ form yellow crystals

NITRO-QUINOLINE-(Py 3) CARBOXYLIC ACID $\text{C}_8\text{H}_5(\text{NO}_2)(\text{CO}_2\text{H})\text{N}$ [220°] Formed by boiling (*Py* 3) methyl quinoline (quinaldine) with nitric acid (*S* *G* 1 4) (Doebner a Miller, *B* 15, 3076) Crystals, sl sol cold water— AgA'

NITRO RESORCIN $\text{C}_6\text{H}_3(\text{NO}_2)(\text{OH})_2$ [4 3 1]

[115°] Formed together with a volatile (2, 3, 1) isomeride [85°], in the preparation of diazoresorcin by the action of nitrous acid on an ethereal solution of resorcin (Weselsky, *A* 164, 1, *M* 1, 887) Lemon yellow needles. When heated with concentrated sulphuric acid it yields $\text{O}(\text{C}_6\text{H}_4(\text{NO}_2)\text{OH})_2$ (Hazura, *M* 4, 610, 5, 188) which forms $\text{Ba}(\text{C}_6\text{H}_4\text{N}_2\text{O}_4)_2$ 2aq and $\text{BaC}_6\text{H}_4\text{N}_2\text{O}_4 \cdot 5\frac{1}{2}\text{aq}$ The isomeride [85°] is converted by the action of nitrous acid into $\text{C}_6\text{H}_4(\text{OH})\text{O}(\text{NOH})(\text{NO}_2)$ [1 3 4 2] (De la Harpe a Reverdin, *Bl* [2] 49, 760)— $\text{BaH}_2\text{A}'_2 \cdot 5\text{aq}$ — $\text{BaH}_2\text{A}'_2 \cdot \text{aq}$ — $\text{BaH}_2\text{A}'_2 \cdot 2\text{aq}$ golden needles

Methyl ethers

$\text{C}_6\text{H}_3(\text{NO}_2)(\text{OH})(\text{OMe})$ [4 3 1] [95°] Volatile with steam— $\text{C}_6\text{H}_3(\text{NO})_2(\text{OMe})(\text{OH})$ [4 3 1]

[144°] Not volatile with steam
Ethyl ethers $\text{C}_6\text{H}_3(\text{NO}_2)(\text{OEt})(\text{OH})$ [79°]

Volatile with steam [131°] Non volatile
Di acetyl derivative $\text{C}_6\text{H}_3(\text{NO})_2(\text{OAc})_2$

[91°] Tables (from alcohol) (Errera, *G* 15, 273)
Di benzoyl derivative
 $\text{C}_6\text{H}_3(\text{NO}_2)(\text{OBz})_2$ [111°] Got by nitration (Schiaparelli a Abelli, *G* 13, 257, Errera, *G*, 15, 271)

Di m nitro-di benzoyl derivative [123°]

Di nitro resorcin $\text{C}_6\text{H}_2(\text{NO}_2)_4(\text{OH})$ [4 2 3 1] [142°] Formed by the action of nitrous fumes, or of cold HNO_3 , upon di nitroso resorcin (Benedikt a Hubl, *M* 2, 323, Barr, *B* 21, 1514, Von Kostanecki, *B* 21, 3122) Formed also by boiling di nitro *m* amido phenol with dilute KOH aq (Lippmann a Fleissner, *M* 6, 814, 7, 98) Golden leaflets— $\text{K}_2\text{A}'_2 \cdot \frac{1}{2}\text{aq}$ — BaA'_2 — $\text{Ag}_2\text{A}'_2$ red pp

Methyl ether $\text{C}_6\text{H}_3(\text{NO}_2)_2(\text{OMe})(\text{OH})$

[75°] Needles (Aronheim, *B* 12, 30)
Di nitro resorcin $\text{C}_6\text{H}_2(\text{NO}_2)_4(\text{OH})_2$ [213°] Formed by nitration of the di acetyl derivative of resorcin (Typke, *B* 16, 552) Yellow prisms or needles— $(\text{NH}_4)_2\text{A}'_2$ — $\text{BaH}_2\text{A}'_2$ — BaA'_2 red scales with violet lustre, v sl sol water

Ethers $\text{C}_6\text{H}_3(\text{NO}_2)_2(\text{OMe})_2$ [67°],

$\text{C}_6\text{H}_3(\text{NO}_2)_2(\text{OEt})_2$ [75°] Obtained by nitration of the ethers of resorcin (Homig, *B* 11, 1039, Aronheim, *B* 12, 82)

Tri nitro resorcin

$\text{C}_6\text{H}(\text{NO}_2)_3(\text{OH})_2$ [6 4 2 3 1] *Styphnic acid*
Oxyphnic acid *Mol w* 245 [175°]. *S.* 64 at 14° (Stenhouse, *Pr.* 19, 410).

Formation—1 By the action of boiling nitric acid on extract of Brazil wood, or sapan wood, euxanthone, gum ammoniac, asafetida, galbanum, pucedanin, ostruthin, saganenium, or the aqueous extract of fustic or sandal wood (Chevreul, *A Ch* 66, 116, 73, 43, Erdmann, *J pr* 37, 409, 38, 355, Botiger a Will, *A* 58, 273; Rothe, *J pr* 46, 376, Gorup Besanex, *A* 183, 336, Stenhouse, *C J* 19, 236) Graebe, *B* 22, 1405) —2 By the action of nitric acid on *m*-nitro phenol, on γ , δ , or ϵ di nitro phenols, and on β or γ tri nitrophenol (Bantlin, *B* 10, 524, 11, 210), Henriques, *A* 215, 340) —3 By the nitration of resorcin or of either di nitroresorcin (Merr a Zetter, *B* 12, 681, Benedikt a Hubl, *M* 2, 326, Von Kostanecki, *B* 21, 3122) —4 By boiling tri nitro phenylene di methyl diamine with potash (Romburg, *R T C* 7, 6) —5 Together with three di nitro benzoic acids, by adding *o* nitro benzoic acid to a mixture of fuming HNO_3 and H_2SO_4 (Griess, *B* 7, 1224, Salkowski, *B* 8, 637)

Properties—Yellow laminae Ppd from its aqueous solution by HCl V sol alcohol and ether

Salts— $(\text{NH}_4)\text{HA}'$ — $(\text{NH}_4)_2\text{A}''$ monoclinic needles, a b c = 1.166 2 09, β = 76° 52' — $\text{Na.A}'$ 2aq — KHA' aq — $\text{K.A}''$ — BaA' 3aq — BaA'' aq — SrA' 2aq — CaA'' 3aq — $\text{Pb}(\text{OH})\text{A}''$ — $\text{MnH.A}'$ 12aq — CoA'' 3aq — $\text{CaK.A}'$ 4aq — $\text{NiK.A}'$ 3aq — CuA'' 4aq — $\text{Cu}(\text{NH}_4)_2\text{A}''$ 7aq — $\text{Cuk.A}'$ 4aq — $\text{Ag.A}''$ aq slender needles

Methyl ether Me A' [124°] Formed by nitrating $\text{C}_6\text{H}_5(\text{OMe})_2$ (Hong, *B* 11, 1039)

Di ethyl ether Et A' [121°] Yields tri nitro *m* phenylene diamine on heating with ammonia (Nolting a Collin, *B* 17, 239)

NITRO RESORCIN SULPHONIC ACID

$\text{C}_6\text{H}_3(\text{NO})_2(\text{OH})_2(\text{SO}_3\text{H})$ [124°] Formed by sulphonating nitro resorcin [115°] (Huzara, *M* 4, 610) Minute crystals (containing 1' aq) — BaA''' 4aq yellow needles — BaA''' 2aq yellow scales — $\text{Ba.A}'''$ 10aq blood red needles

A nitro resorcin disulphonic acid is formed by oxidising the corresponding nitroso compound with H_2O (Ulzer, *M* 9, 1130)

NITRO SALICYLIC ACID v Nitro-oxy BENZOIC ACID

NITROSAMINES Compounds containing nitrosyl (NO) united to nitrogen They are described under the amines from which they are derived by displacement of hydrogen by nitrosyl Nitrosamines are formed by the action of nitrous acid upon secondary bases They are neutral substances and may be reconverted into the parent base by boiling with tin and HCl aq, with zinc and H_2SO_4 , with aniline, or with alcoholic potash (Geuther, *A* 128, 151, Griess, *B* 7, 218, Witt, *C J* 33, 203) Many aromatic nitrosamines are converted into *p* nitroso compounds by alcoholic HCl , the nitrosyl entering the benzene nucleus (Fischer a Hepp, *B* 20, 1247, 2471)

NITROSATES This name is given by Wallach (*A* 241, 288) to compounds formed by the union of nitrogen peroxide with unsaturated hydrocarbons Thus Guthrie's 'amylene nitrite' (vol 1 p 210) would be called amylene nitrosate and may be represented as nitroso-amyl nitrate of formula $\text{C}_5\text{H}_9(\text{NO})(\text{O NO}_2)$ or $\text{C}_5\text{H}_9(\text{NOH})(\text{ONO}_2)$ This body reacts with aromatic

bases forming $\text{C}_5\text{H}_9(\text{NOH})\text{NPh}$ [141°], $\text{C}_5\text{H}_9(\text{NOH})\text{NHCH}_3$ [141°], and the isomeric $\text{C}_5\text{H}_9(\text{NOH})\text{NHCH}_2\text{H}_5$ [115°], which yield nitrosamines melting at 128°, 148°, and 150° respectively Amylene 'nitrosate' reacts in like manner with *o* anisidine, piperidine, and diethylamine, forming bases melting at 139°, 98°, and 72° respectively These bases are termed 'nitrol amines' by Wallach

The term nitrosite is given by Wallach to compounds resulting from the union of N_2O with unsaturated hydrocarbons Thus terpinene nitrosite $\text{C}_{10}\text{H}_{16}\text{N}_2\text{O}$ may be viewed as a nitroso nitrite with formula $\text{C}_{10}\text{H}_{16}(\text{NO})(\text{ONO})$ or $\text{C}_{10}\text{H}_{16}(\text{NOH})(\text{ONO})$ They readily exchange ONO for NHR' or $\text{NR}'\text{R}''$ when acted upon by bases, forming nitrolamines Thus terpinene nitrosite acted upon by ethylamine yields 'terpinene nitrol-ethylamine' $\text{C}_{10}\text{H}_{18}(\text{NOH})\text{NHET}$ [131°]

NITROSO ACETIC ETHER v OXIMIDO ACETIC ETHER

NITROSO ACETOACETIC ETHER $\text{C}_5\text{H}_7\text{NO}$, $\text{CH}_3\text{COCH}(\text{NOH})\text{COEt}$ [54°] Formed by the action of nitrous acid on aceto acetic ether, and on acetyl malonic ether (V Meyer a Zublin, *B* 11, 320, Wiegand, *B* 15, 1050, Ceresole, *B* 15, 1326, Lang, *B* 20, 1327) Prisms, sl sol water, sol alkalis Forms $\text{CH}_3\text{C}(\text{NOH})\text{C}(\text{NOH})\text{COEt}$

Anilide $\text{C}_6\text{H}_5\text{N}_2\text{O}$ [100°] Formed by the action of nitrous acid on the anilide of acetoacetic acid (Knorr, *A* 236, 80) Prisms

NITROSO TRIACETONAMINE v ACETRONAMINE

NITROSO ACETONE $\text{C}_4\text{H}_7\text{NO}$, $\text{CH}_3\text{COCH}(\text{NOH})\text{CH}_3$ or $\text{CH}_3\text{COCH}_2\text{NO}$ *Oxim of pyruvic aldehyde Oxim of methyl glyoxal* [65°] Formed by the action of nitrous acid on acetoacetic ether (V Meyer a Zublin, *B* 11, 695, Ceresole, *L* 15, 1326), and by warming acetone with amyl nitrite and HCl (Claisen, *B* 20, 252) Silvery leaflets or prisms, boiling with decomposition at about 200° V sol water and alcohol, volatile with steam May be sublimed

Reactions—1 Dilute HCl aq at 140° forms acetic and formic acids and ammonia (Treadwell and Steiger, *B* 15, 1059) —2 Tin and HCl give di methyl pyrazine.—3

Hydroxylamines hydrochloride forms methyl glyoxim or acetoximic acid (vol 1 p 38) and a compound, $\text{C}_4\text{H}_9\text{N}_2\text{O}_2$, which detonates at 238°–247°, and forms an explosive hydrochloride $\text{C}_4\text{H}_9\text{N}_2\text{O}_2\text{HCl}$ [113°] (Scholl, *B* 23, 3578) —4 **Phenyl hydrazine** yields $\text{CH}_3\text{C}(\text{NHPH})\text{CH}(\text{NOH})$ [134°] (Pechmann, *B* 21, 2994) —5 **Phenyl methyl-hydrazine** yields $\text{CH}_3\text{C}(\text{NMPH})\text{CH}(\text{NOH})$ [118°]

Methyl ether $\text{C}_4\text{H}_9\text{O}(\text{NOME})$ (115° uncor) Formed by heating nitroso acetone with NaOME (Meyer a Ceresole, *B* 15, 8067, 16, 833) Colourless oil

Ethyl ether $\text{C}_4\text{H}_9\text{O}(\text{NOET})$ (130°).

Benzyl ether $\text{C}_4\text{H}_9\text{O}(\text{NOCH}_2\text{C}_6\text{H}_5)$ [46°].

Di-nitroso-acetone $\text{CH}(\text{NOH})\text{COCH}(\text{NOH})$ [144°] Formed by the action of nitrous acid on acetone dicarboxylic acid (Pechmann a Wehsarg, *B* 19, 2465, 21, 2992) Prisms, sl. sol cold water, decomposed by boiling water into HCy , CO_2 , and water Explodes when heated

Phenyl hydrazide $\text{NHPH OCH}(\text{NOH})$ [145°] Needles. Yields a mono-acetyl derivative [133°].

Phenyl-methyl-hydrazide [187°]**Crystals**

Oxim $\text{CH}(\text{NOH})\text{C}(\text{NOH})\text{CH}(\text{NOH})$ *Tri nitroso propane* [171°] Crystalline powder

NITROSO-ACETOPHENONE $\text{C}_6\text{H}_5\text{NO}_2$ *re* $\text{C}_6\text{H}_5\text{COCHNOH}$ *Oxim of phenyl glyoxylaldehyde* [128°] Prepared by the action of amyl nitrite and NaOEt on acetophenone (Claisen a Manasse, *B* 20, 2194, Braun, *B* 22, 556) Thin, monoclinic plates, $a, b, c = 2.762, 1.2146, b = 66^\circ 54'$ Sol cold water, soluble in aqueous Na_2CO_3 On heating with AcO it yields $\text{C}_6\text{H}_5\text{COCN}$ SnCl_4 in HCl aq, reduces it to ω amido acetophenone and di phenyl pyrazine When its compound with NaHSO_3 is boiled with H_2SO_4 it yields $\text{C}_6\text{H}_5\text{COCHO}$ Hy droxylamine hydrochloride yields $\text{C}_6\text{H}_5\text{N}_2\text{O}_2$ [207°-211°] (Scholl, *B* 23, 3580)

Oxim v Oxim of PHENYL GLYOXAL

NITROSO-ANILINE $\text{C}_6\text{H}_5(\text{NO})(\text{NH})$ [14°] [174°] Formed by heating nitroso phenol with NH_4Cl , ammonium acetate, and ammonium carbonate (Fischer a Hepp, *B* 20, 2175, 21, 684) Steel blue needles (from benzene) Decomposed by NaOHAq into NH_3 and nitroso phenol Tin and HCl reduce it to phenylene *p* diamine Phenyl hydrazine hydrochloride forms $\text{C}_6\text{H}_5\text{N}_2\text{O}$ [125°] Phenyl methyl hydrazine yields $\text{C}_6\text{H}_5\text{N}_2\text{O}$ [151°] — $\text{C}_6\text{H}_5\text{NaN}_2\text{O}$ 2aq yellow crystals

NITROSO-ANTHRONE v ANTHRACENF

NITROSO-BENZENE $\text{C}_6\text{H}_5\text{NO}$ On distilling with steam, the product of the action of nitrosyl chloride on HgPh_2 , there is obtained a pungent green liquid, which yields aniline on reduction (Baeyer, *B* 7, 1638) By oxidising the di-oxim of quinone with alkaline K_2FeCy_6 , there is obtained a golden yellow pp, probably di nitroso benzene $\text{C}_6\text{H}_4(\text{NO})_2$ [14°] It yields *p* phenylene diamine on reduction, and *p* di nitro benzene on warming with HNO_3 Hydroxylamine hydrochloride converts it into quinone dioxim

NITROSO-BENZYL ALLYL-THIO UREA $\text{PhC}(\text{NOH})\text{NHCSNHC}_6\text{H}_5$ Formed by melting together allyl thiocarbimide and benzamidoxim (Koch, *B* 24, 899) Fine needles

ISO-NITROSO-BENZYL-AMINE v BENZ ENYL AMIDOXIM*p* **NITROSO-DI-BENZYL-ANILINE**

$\text{C}_6\text{H}_5(\text{NO})\text{N}(\text{CH}_2\text{C}_6\text{H}_5)_2$ [92°] Formed by adding amyl nitrite to a solution of di benzyl aniline in alcoholic HCl Thin steel blue plates, or small green crystals V sol ether and CS_2 , m. sol. alcohol On reduction it gives *u* di benzyl-*p*-phenylene diamine (Matzudaira, *B* 20, 1616)

NITROSO-BENZYL-MALONIC ACID

$\text{C}_6\text{H}_5\text{C}(\text{NO})(\text{CO}_2\text{H})_2$ [120°] Obtained from its ether, which is got from nitroso malonic ether, NaOEt , and benzyl chloride (Conrad a Bischoff, *A* 204, 121, 209, 215) Plates On fusion, or on boiling with water, it gives benzyl alcohol, CO_2 , and $\text{HCy} - \text{K}_2\text{A}'$ aq (dried at 100°)

NITROSO-BENZYL TOLYL-THIO UREA

$\text{PhC}(\text{NOH})\text{NHCSNH}_2\text{C}_6\text{H}_4\text{Me}$ [67°] Formed by melting *p* tolyl thio carbimide with benzamidoxim (Koch, *B* 24, 897)

NITROSO-BETORCIN $\text{C}_6\text{H}_5\text{NO}_2$ *re*

$\text{C}_6\text{H}(\text{NO})\text{Me}_2(\text{OH})_{1.435}$ [143°] Obtained by the action of nitrosyl sulphate $\text{SO}_3\text{H}(\text{NO})$ on a solu-

tion of betorein (Stenhouse a Groves, *O J* 87, 404, Von Kostanecki, *B* 19, 2823) Red prisms (from HOAc)

p **NITROSO-ISOBUTYL-ANILINE**

$\text{C}_6\text{H}_5(\text{NO})\text{NHCH}_2\text{Pr}$ [94°] Formed from iso butyl aniline, NaNO_2 , and HCl (Wacker, *A* 243, 297) Steel blue crystals, v sol alcohol Yields $\text{C}_6\text{H}_5(\text{NH}_2)\text{NHCH}_2\text{Pr}$ on reduction Boiling alkalis split it up into *p* nitroso phenol and iso butylamine Further treatment with NaNO_2 and HCl yields $\text{C}_6\text{H}_5(\text{NO})\text{N}(\text{NO})\text{CH}_2\text{Pr}$

a **NITROSO-n BUTYRIC ACID** $\text{C}_4\text{H}_7\text{NO}_2$ *re*

$\text{C}_4\text{H}_7\text{C}(\text{NOH})\text{CO}_2\text{H}$ [151°] Formed by the action of nitrous acid and NaOH upon ethyl acetoacetic ether (Wienig, *B* 15, 1057) Flat prisms — AgA' insoluble powder

B Nitroso butyric acid

$\text{CH}_3\text{C}(\text{NOH})\text{CH}_2\text{CO}_2\text{H}$ [140°] Obtained from its ether, which is got by the action of hydroxylamine on acetoacetic ether (Westenberger, *B* 16, 2996) — AgA' white pp

(aB) Di isonitroso butyric acid $\text{C}_4\text{H}_7\text{N}_2\text{O}_4$ *re* $\text{CH}_3\text{C}(\text{NOH})\text{C}(\text{NOH})\text{CO}_2\text{H}$ Obtained from its ethyl ether [140°] which is formed, together with the anhydride $\text{C}_4\text{H}_7\text{N}_2\text{O}_4$ [133°], by the action of hydroxylamine on nitroso acetoacetic ether (Ceresole a Kockert, *B* 17, 821) The acid is crystalline and gives the salts BaA' , $2\frac{1}{2}\text{aq}$ and AgA' The anhydride gives BaA' , $2\frac{1}{2}\text{aq}$ and $\text{Ag}_2\text{A}''$

NITROSO CARVACROL

$\text{C}_6\text{H}_4(\text{NO})(\text{CH}_3)(\text{CH}_2)(\text{OH})$ [153°] Yellow prisms (Paterno a Canzoneri, *B* 12, 388) Reduced by SnCl_2 to amido carvacrol [304°]

NITROSO-CARVENE v CARVONIM

NITROSO COMPOUNDS Compounds containing nitrosyl NO They are usually obtained by the action of nitrous acid They may be divided into nitrosamines (*g v*) in which nitrosyl is attached to nitrogen, and nitroso compounds proper in which nitrosyl is attached to carbon Compounds in which nitrosyl is attached to oxygen are called nitrites Compounds containing the divalent radicle oximodogen NOH are often called isonitroso compounds, the group CNOH being isomeric, and often interchangeable, with the group CHNO The isonitroso compounds may be obtained by the action of hydroxylamine on aldehydes, ketones, and ketonic compounds, and may thus be termed oximides, oximes, or oxims of these ketonic bodies Nitrous acid acting upon the group COCH_2 frequently converts it into $\text{COC}(\text{NOH})$ the new body may be termed either a nitroso derivative of the original ketone, or a monooxim of the diketone COCO , the latter terminology is that usually employed in this dictionary Nitrous acid acting upon a secondary amine forms a nitrosamine, with tertiary aromatic amines and with phenols it yields nitroso compounds, the nitrosyl taking up the *para* position When the *p* position is occupied the nitrosyl can in some cases still enter the benzene nucleus in the *o* position The nitroso phenols $\text{HO C} \begin{smallmatrix} \text{CH CH} \\ \text{CH CH} \end{smallmatrix} \text{CNO}$ and the oxims of the

mono quinones $\text{CO} \begin{smallmatrix} \text{CH CH} \\ \text{CH CH} \end{smallmatrix} \text{CNOH}$ are identical, and will be described under the quinones Some aromatic nitrosamines are converted into

the isomeric *p* nitroso compounds by the action of alcoholic HCl

NITROSO-CREATININE *v* CREATININE

NITROSO-CRESOL *v* Oxim of TOLUQUINONE

DI-NITROSO CRESOREICIN $C_6H_4Me(NO)_2(OH)_2$ or $C_6H_4Me(NO)_2O$ [1 3 5 2 4] Formed from cresorein and HNO_2 (Von Kostanecki, *B* 20, 3135) Pale green plates (containing aq) Explodes above 160° Sl sol water With HNO_3 it yields di nitro cresorein [90°]

DI-NITROSO-CYMENE $C_{10}H_{12}N_2O_2$ [72°] Formed by oxidising the di oxim of thymoquinone with alkaline K_2FeC_4 (Kehrmann a Messinger, *B* 23, 3560) Greenish yellow pp, smelling like iodine After one fusion it melts at 130°

NITROSO-ETHYL-ACETONE is the (β) oxim of METHYL ETHYL DIKETONE (*q v*)

NITROSO-ETHYL-ANILINE $C_8H_9N_2O$ *see* [4 1] $C_8H_9(NO)(NH_2Et)$ [78°] Obtained by adding alcoholic HCl to an ethereal solution of the nitrosamine of ethyl aniline (Fischer a Hepp, *B* 19, 2993) Green plates, v sol alcohol Yields $C_8H_9(NH_2)(NH_2Et)$ (270°) on reduction On heating with aqueous NaOH it is split up into nitroso phenol and ethylamine — $B'HCl$ needles, v sol water

Nitroso di ethyl aniline [1 4] $C_8H_9(NO)NEt_2$ [84°] Formed from di ethyl aniline and nitrous acid (Kopp, *B* 8, 621) Green prisms (from ether) Decomposed by boiling dilute NaOH into nitroso phenol and diethylamine Salts — $B'H_2P_2Cl_4$ — $B'I_3$ [118 5°] (Dafert, *M* 4, 506) — $B'I_3$ [127°] — $B'H_2SO_4$ — $B'2C_2H_5(NO)_2(OH)$ — $B'HCy$ [171°] (Lippmann, *M* 6, 544) Orange crystals (from alcohol)

NITROSO DI ETHYL KETONE *v* (a) Oxim of METHYL ETHYL DIKETONE

a NITROSO ETHYL-PHENYL AMINE

$C_8H_9(NH_2)CH(NO)CH_3$ Formed by heating $C_8H_9(NH_2)CHNO$ with KOH, methyl alcohol, and MeI at 100° (Gabriel a Meyer, *B* 14, 2339) Yellowish oil — $B'HCl$ prisms

Acetyl derivative

$C_8H_9(NHAc)CH(NO)CH_3$ [109°]

NITROSO ETHYL o TOLUIDINE C_8H_9NO *see* $C_8H_9Me(NO)NH_2Et$ [140°] Green plates (Fischer, *B* 19, 2994)

NITROSO ETHYL-o-XYLIDINE $C_{10}H_{11}NO$ *see* $C_8H_9(NO)Me(NH_2Et)$ [1 2 3 4] [124°] Green crystals (Menton, *A* 263, 327) — $B'HCl$ needles

NITROSO FORMANILIDE *v* FORMIC ACID

a NITROSO GLUTARIC ACID

$CO_2H C(NO)CH_2CH_2CO_2H$ [152°] Produced by boiling furazyl propionic acid

$O < N C CH_2CH_2CO_2H$ [86°] with potash, the semi nitrile $CN C(NO)CH_2CH_2CO_2H$ [87°] being formed at the same time (Wolff, *A* 260, 112) Prisms Yields amido-glutaric acid on reduction — $BaA''1\frac{1}{2}aq$ needles, v sl sol water

Amidoxim

$CO_2H CH_2CH_2C(NO)C(NO)NH_2$ [158°] Formed by the action of hydroxylamine on the semi nitrile [87°] Needles, sl sol water

NITROSO GUANIDINE *v* GUANIDINE

NITROSO HEXOIC ETHER

$CH_3C(NO)CH_2CH_2CO_2Et$ Formed from ethyl-acetoacetic ether and hydroxylamine (Westenberger, *B* 16, 2997) Oil.

Nitroso hexoic acid $CH_3C(NO)CMe_2CO_2H$ [97°] By the action of amyl nitrite and nitric acid on tri methyl ethylene there is formed a compound $CMe_2(ONO)C(NO)CH_3$ which when heated with alcoholic KCy yields the nitrile $CH_3C(NO)CMe_2CN$ [100°] (230°) which forms on saponification the corresponding amide [164°] and acid [97°] (Wallach, *A* 248, 166) The acid is split up by heat into CO_2 and the oxim of methyl isopropyl ketone

NITROSO-INDOXYL *v* ψ -ISATIN OXIM

NITROSO MALONIC ACID $CH(NO)(CO_2H)_2$ or $C(NO)(CO_2H)_2$ Oxim of mesoxalic acid

Formation — 1 By treating barbituric acid with nitrous acid and warming the resulting violonic acid with potash (Baeyer, *A* 131, 292) 2 From its ether, which is got by passing nitrous fumes into sodium malonic ether (Conrad a Bischoff, *B* 13, 599, *A* 209, 211) — 3 By the action of hydroxylamine on mesoxalic acid (Meyer a Muller, *B* 16, 608)

Properties — Needles Decomposes at 126° with violence At 40° its aqueous solution gives off CO_2 and HCy Sodium amalgam reduces it to amido malonic acid

Salts — $KA''\frac{1}{2}aq$ — $PbA''aq$ — $AgA''\frac{1}{2}aq$

Ethyl ether Et_2A'' SG $\frac{1}{2}$ 1 149 Oil

NITROSO MESITYL OXIDE

$CMe_2CHCOCHNOH$ [102°] Prisms (Claisen a Manasse, *B* 22, 526)

NITROSO METHYL ACETONE *v* Oxim of DI METHYL DIKETONE

NITROSO DI METHYL p AMIDO BENZOIC ACID $C_8H_9(NH_2)(NMe_2)CO_2H$ [224°] Formed by the action of nitrous acid on $C_8H_9(NMe_2)CO_2H$ (Bischoff, *B* 22, 342) Plates (from alcohol) Yields $C_8H_9(NMe_2)CO_2H$ on reduction with $SnCl_2$ and HCl Salts — $B'H_2C_2O_4$ [178°-181°] — $B'C_2H_5(NO)_2OH$ [168°] — $B'HCl$ slender needles

Methyl ether $MeA''6\frac{1}{2}aq$ [101°] — $B'HCl$ — $B'C_2H_5(NO)_2OH$ Golden yellow needles

NITROSO DI METHYL-AMIDO BENZO PHENONE $C_8H_9CO C_6H_4(NH_2)NMe_2$ Oil (Bischoff, *B* 22, 340)

Nitroso-tetra-methyl-diamido-benzophenone *v p* 263

p NITROSO-METHYL ANILINE $C_8H_9N_2O$ *see* $C_8H_9(NO)NMe_2H$ [118°] Formed by adding alcoholic HCl to an ethereal solution of the nitrosamine $C_8H_9NMe_2(NO)$ (Fischer a Hepp, *B* 19, 2991) Steel blue prisms (from water) Decomposed by $NaOHAq$ into *p* nitroso phenol and methylamine Yields $C_8H_9(NH_2)(NMe_2H)$ on reduction Nitrous acid forms the nitrosamine $C_8H_9(NO)NMe_2NO$ [101°] Nitric acid (SG 1 13) yields the nitro-compound $C_8H_9(NO_2)NMe_2NO$ [104°]

Nitroso-di-methyl aniline $C_8H_9(NO)NMe_2$ or $C_8H_9 < NMe_2 > O$ Mol w 160 [85°] Formed by the action of nitrous acid on di-methyl aniline hydrochloride (Baeyer a Caro, *B* 7, 963, Schraube, *B* 8, 616, Wunster, *B* 12, 523, 1825, Meldola, *C J* 39, 37) Green plates (from ether)

Reactions — 1 Reduced by *tin* and $HClAq$ to $C_8H_9(NH_2)(NMe_2)$ — 2 Resolved by boiling alkali into nitroso phenol and dimethylamine — 3 K_2FeC_4 oxidises it to $C_8H_9(NO_2)(NMe_2)$ — 4 Alcoholic potash forms $NO(C_8H_9NMe_2)$ — 5 $HClAq$ at 105° yields $C_8H_9(NH_2)(NMe_2)$.

$C_6H_4Cl_2(NH_2)(NMe_2)$ and $C_6H_4Cl_2(NH_2)_2$ (Möhlan, *B* 19, 2010) — 6 *Phenyl hydrazine acetate* yields $C_{12}H_{11}N_3O$ [108°] This base is also formed by the action of diazobenzene on nitroso dimethylaniline (O Fischer, *B*, 21, 2610, 22, 623) Phenyl methyl hydrazine yields the compound $C_{12}H_{15}N_3O$ [141°]

Salts — $B'HCl$ yellow needles — $B'H_2SO_4$ — $B'H_2C_2O_4$ 2aq — $B'H_2C_2O_4$ — $B'_2H_2FeCy_2$ aq — $B'_2H_2FeCy_2$ 2aq — B'_2AgNO_3 — B'_2H_2Cy [222°] (Lippmann a Fleissner, *M* 6, 537)

Combinations — B'_2I_2 [116°] (Dafert, *M* 4, 506) — B'_2I_2 [124°] — B'_2PhNH_2 steel blue crystals — $B'_2NH_2C_6H_4Me$ [14] — B'_2PhOH — $B'_2C_6H_5$ dark green crystals — $B'_2HCyC_6H_5$ — $B'_2HCyC_6H_4NO_2$ — $B'_2HCyC_6H_5$ — $(B'HCy)(PhN_2)_2$

NITROSO-METHYL OXINDOLE v. Oxim of

METHYL-ISATIN

NITROSO-METHYL-o-TOLUIDINE

$C_8H_9Me(NO)(NHMe)$ [152] [151°] Formed by the action of HCl on the isomeric nitrosamine $C_8H_9Me(NMeNO)$ (Kock, *A* 243, 308) Green plates Split up by boiling $NaOHAq$ into nitroso-*o* cresol and methylamine $KMnO_4$ oxidises it to nitro methyl *o* toluidine — $B'H_2Cl_2$ aq [110°] Yellow cubes

Nitroso di-methyl *m*-toluidine

$C_8H_9Me(NO)NMe_2$ [125] [92°] Formed by the action of nitrous acid on di methyl *m* toluidine (Wurster a Riedel, *B* 12, 1796, 13, 126) Light green needles Decomposed by boiling $NaOHAq$ into NMe_2H and the oxim of *m* tolouquinone — $B'HCl$ — $B'_2H_2FeCy_2$ aq — $B'_2H_2FeCy_2$ 2aq yellow needles

NITROSO-METHYL-o-XYLIDINE

$C_8H_9Me(NO)(NHMe)$ [1263] [161°] Green needles, sl sol water (Menton, *A* 263, 323) — $B'HCl$ crystalline meal

NITROSO-NAPHTHALENE $C_{10}H_7(NO)$ [89°]

Formed from $Hg(C_{10}H_7)_2$ and $NOBr$ in CS_2 (Bayer, *B* 7, 1639, 8, 615) Yellow crystals

Di-nitroso-naphthalene $C_{10}H_6(NO)_2$ [14]

Formed by the action of alkaline K_2FeCy_4 on the dioxim of (α)-naphthoquinone (Nietzki a Guitermann, *B* 21, 433). Pale yellow powder, exploding at 120° Insol water and alcohol

Di-nitroso naphthalene $C_{10}H_6(NO)_2$ [12]

[126°] Formed from (β) naphthoquinone dioxim and K_2FeCy_4 (Leuckart, *B* 19, 174) Needles, m sol alcohol

NITROSO-NAPHTHOL v Oxim of NAPHTHOQUINONE

NITROSO (β) NAPHTHOLSULPHONICACID

$C_{10}H_6(NO)(OH)SO_3H$ or $C_{10}H_6(NO)(OH)O(SO_3H)$ [123] or [132] Formed from ammonium (β) naphthol sulphate, $NaNO_2$, and HCl (Meldola, *C* J 39, 41) Very soluble orange crystals Its solution gives with phenol in $HOAc$ a blue colour, becoming red on dilution, and with diphenyl amine a blue colour, remaining blue on dilution Tin and HCl reduce it to the amido acid — BaA' aq orange needles — $BaC_2H_3NSO_3$ 2aq green needles — $Ag(NH_4)A'$ — $(NH_4)A'$ aq green — MgA'' 3aq orange — ZnA'' 3aq — PbA'' aq

An isomeric acid, obtained by reducing benzene azo (β) naphthol sulphonic acid with aqueous ammonium sulphide, crystallises in sparingly soluble grey needles (Griess, *B* 14, 2042)

NITROSO-DI-(α) NAPHTHYL-AMINE

$C_{10}H_7(NO)NHO_2H$. [169°] Formed from

$(C_{10}H_7)NNO$ and alcoholic HCl (Fischer a Hepp, *B* 20, 1248, Wacker, *A* 243, 301) Dark red needles With boiling dilute H_2SO_4 it yields the mono oxim of (α) naphthoquinone and (α) naphthylamine — $B'HCl$ green needles -

Nitroso (β) naphthylamine

$C_{10}H_7(NO)NH_2$ [12] [162°] Formed by heating the (α) oxim of (β) naphthoquinone with NH_4Cl and NH_4OAc (Ilinski, *B* 17, 391, Harden, *A* 255, 150) Dark green needles (from dilute alcohol Yields naphthylene *o* diamine on reduction and the di oxim of (β) naphthoquinone on treatment with hydroxylamine — $B'HCl$ — $B'_2H_2PtCl_6$ — $B'_2H_2SO_4$ aq

(α) NITROSO - (β) - NAPHTHYL-ETHYL-

AMINE $C_{11}H_9(NO)NHEt$ [121°] Formed by the action of alcoholic HCl at 6° on the nitrosamine $C_{11}H_9N(Et)(NO)$, and also by the action of ethyl amine on the (α) oxim of (β) naphthoquinone (Fischer a Hepp, *B* 20, 2471, 21, 686) Green crystals Yields a crystalline nitrosamine $C_{12}H_{11}N_2O_2$, decomposing at 105°

(α) Nitroso-(α) naphthyl ethyl amine

$C_{11}H_9N_2O$ v $C_{10}H_7(NO)NHEt$ [14] [133°] Formed in like manner (Kock, *A* 243, 310) Brown pp Reduced by stannous chloride to $C_{10}H_7(NH)(NHEt)$ Yields ethylamine and $C_{10}H_7(NO)H$ *o* when boiled with $NaOHAq$ — $B'HCl$ — Picrate $C_{18}H_{15}N_3O_8$ [174°] — $C_{12}H_{13}N_2O_2Na$ white spangles, v sol water

Tetrahydride $C_8H_7 < C(NHEt)CH$

[119°] Formed from the isomeric nitrosamine and alcoholic HCl (Bamberger a Helwig *B* 22, 1814) Needles — $B'HCl$ golden crystals Nitroso naphthyl di ethyl amine $C_{10}H_7(NO)NEt_2$ [165°] Formed by adding $NaNO_2$ to a well cooled solution of the base (B E Smith, *C* J 41, 182) Reddish golden scales Gives a blue colour with H_2SO_4

NITROSO-NITRATES v NITROSATES

NITROSO-NITRO ANTHRONE $C_{14}H_9N_2O_4$

[263°] Formed by the action of alkalis on 'hydro anthracene nitrite (Liebermann, *B* 14, 467, cf vol 1 p 277)

NITROSO-NITRO-BUTANE $C_4H_9N_2O_4$ v

$CH_3C(NO)(NO_2)C_2H_5$ *Pseudobutylnitrole* [58] Formed by the action of KOH , KNO_3 , and dilute H_2SO_4 on β nitro butane (Meyer a Locher, *A* 180, 186) and of NO on $CH_3C(NO)C_2H_5$ (Scholl, *B* 21, 508) White prisms (from chloroform), yields a blue liquid on fusion Insol water and alkalis

The isomeric $PrCH(NO)(NO_2)$ and $PrCH(NO)(NO_2)$ are oils (Demole, *B* 7, 790, Züblin, *B* 10, 2084)

NITROSO NITRO PENTANE $Et_2C(NO)(NO_2)$

[68°] Formed from Et_2CNOH and N_2O_4 (Scholl, *B* 21, 509)

NITROSO-NITRO-PROPANE $C_3H_7N_2O_4$ v

$CH_3C(NO)(NO_2)CH_3$ [76°] Formed from acetoxim and N_2O_4 (Scholl, *B* 21, 508) The isomeric compound $CH_3CH_2CH(NO)NO_2$ or $CH_3CH_2C(NO)NO_2$ melts at 60° (Meyer, *A* 175, 114)

NITROSO-NITRO RESORCIN $C_6H_4N_2O_4$ v

$C_6H_4(OH)(NO_2)O(NOH)$ [1234] Formed from nitro resorcin [85°] and nitrous acid (De la Harpe a Reverdin, *B* 21, 1405) Brown needles, not melted at 200°, explodes at a higher temperature.

NITROSO ORCIN $C_8H_7Me(NO)(OH)$. Formed from orcin, NaOHAq, and amyl nitrite (Kramer, B 17, 1888). Dark red prisms. When heated with orcin and H_2SO_4 , it gives the dyestuff $C_{11}H_{11}NO_3$.

Di nitroso orcin $C_8H_7Me(NO)_2(OH)_2$ [1 2 4 3 5]. Formed by adding H_2SO_4 , containing N_2O , to a solution of orcin (Stenhouse a Groves, C J 31, 544). Yellow prisms. Blackens about 140° without melting. Alcoholic hydroxylamine hydrochloride at 100° yields $C_8H_7Me(NO)_2$, whence AcO forms $C_8H_7Me(N_2O)_2$ [47°] (Goldschmidt, B 20, 1607). The compound $C_8H_7Me(NO)_2$ yields, on oxidation by potassium ferriocyanide, the product $C_8H_7Me(NO)_2$ [103°], crystallising in pale-yellow needles.

NITROSO OXANTHRANOL $C_{11}H_9NO$. Formed by boiling 'hydro anthracene nitrite' with alkalis (Liebemann, B 14, 471). Orange flakes, sol alkalis.

NITROSO OXINDOLE v **ISATIN OXIM**

NITROSO OXY METHYL QUINOLINES

C_8H_7N $C_8H_7Me(NO)(OH)$. The following crystalline compounds have been obtained by the action of nitrous acid on the oxy methyl quinolines (Noelting a Trautmann, B 23, 3665) — (B 3) nitroso (B 4) oxy (B 1) methyl quinoline, (L 1) nitroso (B 4) oxy (B 2) methyl quinoline, (B 4) nitroso (B 1) oxy (B 2) methyl quinoline, (B 1) nitroso (B 4) oxy (L 3) methyl quinoline, and (B 2) nitroso (B 1) oxy (B 4) methyl quinoline. Friedlander and Muller (Py 20, 2011) have obtained in like manner (Py 1,3) nitroso-oxy (Py 4) methyl quinoline crystallising in red needles.

NITROSO OXY DI PHENYL AMINE

$C_6H_5(NO)(OH)NPh$ [4 3 1]. Formed from $C_6H_5(OH)NPh(NO)$ and alcoholic HCl (Kohler, B 21, 909). Red needles, m sol alcohol.

NITROSO (B 4) OXY QUINOLINE

$C_8H_7(NO)(OH)N$. Formed from o oxy quinoline and nitrous acid (Luppmann a Fleisner, M 10, 794). Yellow needles. The isomeric nitroso (B 2) oxy quinoline crystallises from $HOAc$ in golden needles (Mathews, B 21, 1886).

(Py 2) Nitroso (Py 1,3) di oxy quinoline

$C_6H_5 \begin{smallmatrix} CO C(NO)H \\ N C(OH) \end{smallmatrix}$ [208°]. Formed by the action of nitrous acid on (Py 1,3) di oxy quinoline (oxycarbostyryl (Baeyer a Homolka, B 16, 2216). Orange prisms. Decomposed by cone $HClAq$ into isatin and hydroxylamine. $SnCl_4$ yields tri oxy quinoline.

NITROSO PHENOL v **MONO-OXIM OF QUINONE**

NITROSO PHENYL ACETIC ACID v **Oxim**

of **PHENYL GLYOXYLIC ACID**

p NITROSO DI PHENYL-AMINE

$C_6H_5(NO)NPhH$ [143°]. Formed from diphenyl nitrosamine and alcoholic HCl (O Fischer a Hepp, B 19, 2991, 21, 677, 2614). Green plates (from benzene). With phenyl hydrazine hydrochloride it yields a compound $C_{18}H_{16}N_2O$ [112°]. Free phenyl hydrazine in ether yields amido diphenylamine [75°] and $C_6H_5N_2O$ [173°] v Bromo aniline yields $C_6H_5Br.N$ [243°]. Tin and HCl reduce it to $C_6H_5(NH_2)NPhH$ [65°] (Ikuta, A 243, 274). Aqueous NaOH yields aniline and quinone oxim. — $B'HCl$ bronze tablets or needles.

Acetyl derivative [97°]. Red prisms.

Nitrosamine $C_6H_5(NO)NPh(NO)$ [98°].

NITROSO-PHENYL BENZYL-THIO-UREA

$PhC(NO)NHCSNPh$ [172°]. Crystals (from alcohol) (Koch, B 24, 394).

NITROSO-PHENYL (a)-NAPHTHYL AMINE

$C_{10}H_7(NO)NPh$ [150°]. Formed from phenyl-naphthyl nitrosamine and alcoholic HCl (Fischer a Hepp, B 20, 1247). Brownish yellow crystals. Yields $C_{10}H_7(NH_2)NPh$ on reduction. Boiling dilute H_2SO_4 splits it up into aniline and quinone oxim — $B'HCl$ green plates.

TRI NITROSO PHLOBOGLUCIN

$C_6(NO)_3(OH)$. Prepared by the action of KNO_3 and $HOAc$ on phloroglucin (Benedikt, B 11, 1374). Needles, sol water and alcohol — K_4A''' needles, exploding above 130° .

TRI-NITROSO-PROPANE v. *Oxim* of Di-

NITROSO ACETONE

NITROSO PROPIONIC ACID v. *Oxim* of **Pyruvic acid**

NITROSO PROPYL-ANILINE $C_8H_7N_2O$ v $C_8H_7(NO)NPrH$ [59°]. Formed from the isomeric $C_8H_7NPr(NO)$ and alcoholic HCl (Wacker, A 243, 291). Steel blue needles, v sol alcohol. Yields $C_8H_7(NH_2)NPrH$ on reduction, and quinone oxim and $NPrH_2$ on boiling with alkalis. $NaNO_2$ and HCl yield $C_8H_7(NO)NPr(NO)$ [69°].

Nitroso di propyl-aniline $C_8H_7(NO)NPr_2$ [42°]. Formed from di propyl aniline (241°) and HNO_2 (Mandl, M 7, 99). Green trimetric crystals, $a b c = 576 \cdot 1 \cdot 277$. Yields quinone oxim and dipropylamine on warming with potash. HCl forms $C_8H_7N_2O$ [140°].

NITROSO PROPYL CRESOL $C_8H_7NO_2$ v $C_8H_7MePr(NO)(OH)$. *Cymoquinone oxim* [140°] is formed from propyl cresol, KNO_3 , and $HOAc$ (Mazzara, G 12, 167). The isomeric $C_8H_7MePr(NO)(OH)$ melts at 167° .

NITROSO RESORCIN $C_6H_4(NO)(OH)_2$ [4 3 1]. *Oxy quinone oxim*. Formed from $C_6H_4(OH)_2(ONa)$ and amyl nitrite (Fèvre, Bl [2] 39, 585, C R 96, 790). Golden crystals (containing aq), turning brown at 112° . Its solution is turned deep green by a ferrous salt. $SnCl_4$ reduces it to amido resorcin. H_2O yields nitroresorcin (Ulzer, M 9, 1128) — NH_4A 2aq — $KA'Aq$ — $NaA'Aq$ — AgA' brown needles.

Methyl ether MeA' (Aronheim, B 12, 80).

Ethyl ether EtA' . Yellow flakes.

Di nitroso resorcin $C_6H_4(NO)_2(OH)_2$ [4 2 3 1].

Formed by the action of nitrous acid on resorcin (Fitz, B 8, 631, Kostanecki, B 22, 1345). Yellowish plates (containing aq). Explodes at 115° . Forms coloured lakes. Yields di amido-resorcin on reduction — NH_4A' — NaA' dark-green powder.

NITROSO RESORCIN DISULPHONIC ACID

The salt $C_6H_4(NO)(OH)(OK)SO_3K$ is formed by the action of KNO_2 and $HOAc$ on potassium resorcin disulphonic acid (Ulzer, M 9, 1127). It forms violet crystals.

NITROSO SUCCINIC ACID $C_4H_5NO_4$ v $CO_2HCH_2C(NO)CO_2H$. Obtained from its mono ether, which is got by allowing di nitroso succino succinic ether to stand with water (Ebert, A 229, 65). Crystals, decomposing below 126° — CaA'' 4aq.

Mono-ethyl ether $EtHA''$ [111°] — NH_4EtA'' — $CaEtA''$ 2aq — $CaC_4H_5NO_4$ 2aq — $BaC_4H_5NO_4$ aq — $Zn(EtA'')_2$ — $AgEtA''$.

An isomeric ether $EtHA''$ [54 7°] is obtained by the action of $NaOEt$ on the oxim of oxalacetic

ether (Prutti, *C O* 1888, 1460, 1890, 938, Hantzsch, *B* 23, 11)

Di-ethyl ether Et.A'. Oil Identical with the oxim of oxalacetic ether

Di nitroso succinic acid
 $\text{CO}_2\text{H C}(\text{NOH})\text{C}(\text{NOH})\text{CO}_2\text{H}$ [130°] Formed from carboxy tartaric acid and hydroxylamine (Muller, *B* 16, 2985) Prisms— $\text{Ag}_2\text{A}'$ explosive pp

NITROSOSULPHATES and **NITROSO SULPHURIC ACIDS** *v* **SULPHATES** and **SULPHURIC ACID** in vol iv

NITROSO THIOGLYCOLLIC ACID
 $\text{HS C}(\text{NOH})\text{CO}_2\text{H}$ Formed by boiling nitrosothiohydantoin with baryta water (Maly a. Andreasch, *M* 1, 168, *B* 13, 601) Crystals, *v* sol ether Gives a blue colour with FeCl_3 Decomposed by boiling water or alcohol into CO_2 , hydrogen sulphocyanide, and H_2O — BaA' aq

NITROSO-THIOHYDANTOIN $\text{C}_2\text{H}_3\text{N}_2\text{OS}$ Formed from thiohydantoin and HNO_2 (Maly, *B* 12, 967) Crystalline powder, *sl* sol water

NITROSO-THYMOL *v* *Oxim* of **THYMOQUINONE**

DI NITROSO-TOLUENE $\text{C}_6\text{H}_4\text{Me}(\text{NO})_2$ [6 3 2or1] [α 144°] Formed by oxidising toluquinone dioxim with K_2FeCy_4 (Nietzki, *B* 21, 492, Mehne, *B* 21, 734) Amorphous insoluble powder, volatile with steam Gasified on fusion Reconverted by hydroxylamine into toluquinone dioxim

NITROSO o-TOLUIDINE $\text{C}_6\text{H}_4\text{NO}$ *z* $\text{C}_6\text{H}_3\text{Me}(\text{NO})(\text{NH}_2)$ [1 5 2] [116°] Formed by heating toluquinone mono oxim (nitroso *o* cresol, with acetate and chloride of ammonium (Mehne, *B* 21, 731) Small green needles with blue reflex Yields NH_3 , and nitroso *o* cresol on heating with aqueous NaOH Hydroxylamine yields toluquinone dioxim

Nitroso-m-toluidine
 $\text{C}_6\text{H}_3\text{Me}(\text{NO})(\text{NH}_2)$ [1 2 5] [178°] Formed in like manner from nitroso *m* cresol Resembles its isomeride and yields the same dioxim on treatment with hydroxylamine

o-NITROSO-VALERIC ACID is the *Oxim* of **PROPYL GLYOXYLIC ACID**

γ-Nitroso-valeric acid
 $\text{CH}_3\text{C}(\text{NOH})\text{CH}_2\text{CH}_2\text{CO}_2\text{H}$ *Oxim* of *acetylpropionic acid* [96°] Formed from *β* acetylpropionic acid (levulic acid) and hydroxylamine (Muller, *B* 16, 1617) Prisms Yields levulic acid on treatment with tin and HCl H_2SO_4 at 100° forms methyl succinimide (Bredt, *A* 251, 816, cf Rischbicht, *B* 20, 2671)— $\text{BaA}_2\cdot 2\text{aq}$ — AgA' white pp

Ethyl ether Et.A' Oil
γ-Di nitroso valeric acid
 $\text{CH}(\text{NOH})\text{C}(\text{NOH})\text{CH}_2\text{CH}_2\text{CO}_2\text{H}$ [136°] Formed from glyoxyl propionic acid and hydroxylamine (Wolf, *A* 260, 93) Prisms Converted by conc H_2SO_4 at 70° into furazyl propionic acid $\text{O} \begin{smallmatrix} \text{N} \\ \text{C} \end{smallmatrix} \text{CH}_2\text{CH}_2\text{CO}_2\text{H}$ [86°]. — $\text{BaA}'\cdot 3\text{aq}$ thin needles

NITROSO-XYLENOL *v* *Oxim* of **XILOQUINONE**

180-NITRO STEARIC ACID $\text{C}_{18}\text{H}_{33}(\text{NO})\text{O}_2$ Formed by boiling stearic acid (100 g) with HOAc (1500 cc) and HNO_3 (250 g of SG 1.48) for four days (Claus, *J pr* [2] 43, 161) Yellowish

buttery mass, *v* sol ether and alcohol, *insol* water and ligroin Yields stearic acid on reduction It is therefore not a true nitro compound. — $\text{K}_2\text{A}'$ — $\text{K}_2\text{A}'\cdot\text{KHCO}_3$ — $\text{Na}_2\text{A}'$ granular mass, *insol* ether— SrA' — CaA' — CuA' light green

NITRO-STYCHNINE *v* **STYCHNINE**

NITRO-STYRENE The *o*, *m*, and *p*-isomerides $\text{C}_6\text{H}_4(\text{NO})\text{CH CH}_3$, melting at 14°, −5°, and 29° respectively, are formed by boiling the acids $\text{C}_6\text{H}_4(\text{NO})\text{CHBrCH CO H}$ with aqueous Na CO_3 (Einhorn, *B* 16, 2213, Prausnitz, *B* 17, 597, Basler, *B* 16, 3005) The corresponding dibromides $\text{C}_6\text{H}_4(\text{NO})_2\text{CHBrCHBr}$ melt at 52°, 79°, and 73° respectively

ω-Nitro styrene $\text{C}_6\text{H}_5\text{CH CH NO}_2$ [58°] (250° 260°) Formed by heating benzoic aldehyde with nitro methane and ZnCl_2 for 8 hours at 160° (Priests, *A* 225, 319) Formed also by boiling styrene with HNO_3 (Simon, *A* 31, 269, Blyth a. Hofmann, *A* 53, 297) Yellow crystals, yielding benzoic acid on oxidation When freshly prepared it is soluble in NaOHAq , but the solution slowly decomposes, yielding benzoic aldehyde and resin Diluted H_2SO_4 yields benzoic aldehyde, hydroxylamine, and CO Conc HClAq yields hydroxylamine and PhCHCl CO H [78°]

ω-Di-nitro-styrene $\text{C}_6\text{H}_3(\text{NO})_2\text{CH CH}(\text{NO})_2$ [107°] Formed, together with the *ω* isomeride, by nitration of *ω* nitro styrene (Priests) Yellow needles (from alcohol)

ωp Di-nitro styrene [199°] Formed as above, and also by the action of HNO_3 and H SO_4 on *p* nitro cinnamic acid at 0° (Friedlander a. Mahly, *A* 229, 221, *B* 16, 851) Yellow crystals, *sol* KOHAq Dilute H_2SO_4 decomposes it into *p* nitro benzoic aldehyde, hydroxylamine, and CO

ωm-Di-nitro-styrene [122°] Formed from *m* nitro cinnamic acid, HNO_3 , and H SO_4 at 0° (Friedlander a. Lazarus, *A* 229, 233) Yellowish plates (from water) Conc H_2SO_4 at 100° gives CO and *m* nitro benzaldoxim

o-NITRO STYRYL-ACROLEIN $\text{C}_6\text{H}_4\text{NO}_2$ *z* $\text{C}_6\text{H}_3(\text{NO}_2)_2\text{CH CH CO CH O}$ [153°] Formed from *o* nitro cinnamic aldehyde, aldehyde, and very dilute NaOH (Einhorn, *B* 17, 2026) Pale yellow crystals (from dilute alcohol)

o-NITRO-STYRYL ACRYLIC ACID
 $\text{C}_6\text{H}_4(\text{NO}_2)\text{CH CH CH CO}_2\text{H}$ [218°] Formed from *o* nitro cinnamic aldehyde, Ac_2O , and sodium acetate, and obtained also by oxidising $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CH}_2\text{CO CH}_3$ with aqueous NaOCl (Diehl a. Einhorn, *B* 18, 2331) Slender felted needles, *v* sol hot alcohol

o-NITRO STYRYL GLYOXYLIC ACID
 $\text{C}_6\text{H}_4(\text{NO}_2)\text{CH CH CO CO}_2\text{H}$ [136°] Formed from *o* nitro benzoic aldehyde, pyruvic acid, and HClAq (Baeyer a. Drewson, *B* 15, 2862) Crystals Converted by alkalis into indigo

DI-m-NITRO-DI STYRYL KETONE [232°] Sparingly soluble flakes (Von Miller a. Rohde, *B* 22, 1839)

o NITRO STYRYL METHYL KETONE
 $\text{C}_6\text{H}_4(\text{NO}_2)\text{CH CH CO CH}_3$ [60°] Formed, together with the *p* isomeride [110°], by nitration of styryl methyl ketone (Baeyer a. Drewson, *B* 15, 2858, 16, 1958) Formed also by boiling $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CH}_2\text{CO CH}_3$ with Ac_2O , or $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{CH}_2\text{CH CO CH}_3\cdot\text{CO}_2\text{CH}_3$ with dilute

H_2SO_4 (Fischer & Kuzel, *B* 16, 35) Long needles, sol alcohol

***m* NITRO STYRYL-QUINOLINE** $\text{C}_{11}\text{H}_{12}\text{N}_2\text{O}_2$
 $\text{C}_6\text{H}_5\text{C}_6\text{H}_4\text{NCHCHC}_6\text{H}_4\text{NO}_2$ [132°]
 Formed by heating methyl quinoline (lepidine) with *m* nitro benzoic aldehyde and KHSO_4 at 160° (Heymann & Königs, *B* 21, 1429) Needles (from alcohol)

An isomeride [139°] is formed from (*Py* 3)-methyl quinoline and *m* nitro benzoic aldehyde (Wallach, *B* 16, 2009, Wartman, *B* 23, 3646). It gives the crystalline salts B'HCl — B'HN_2 — $\text{B'C}_6\text{H}_5(\text{NO}_2)_2\text{OEt}$ — B'H.PtCl_4 1½ aq

***p* Nitro styryl quinoline** [165°] Formed from (*Py* 3) methyl quinoline and *p* nitro-benzoic aldehyde (Bulach, *B* 20, 2047, 22, 285) Needles Yields a dibromide $\text{C}_{11}\text{H}_{12}\text{N}_2\text{O}_2\text{Br}_2$ [276°] which turns brown at 230°

NITRO DI-STYRYL DI-VINYL KETONE *v*
 NITRO PHENYL BUTYL PHENYL BUTYL KETONE

NITRO α SULPHO-BENZOIC ACID $\text{C}_6\text{H}_4\text{NSO}_3$
 $\text{C}_6\text{H}_4(\text{NO}_2)(\text{SO}_3\text{H})(\text{COH})$ [4 2 1] [110°]
 Formed by oxidising nitro toluene sulphonic acid (Hart, *Am* 1, 350, Hedrick, *Am* 9, 411, Kastle, *Am* 11, 177) White needles (from hot water) — KHA 1 aq long needles — KA — BaA

Chloride $\text{C}_6\text{H}_4(\text{NO}_2)(\text{SO}_2\text{Cl})(\text{COCl})$ [60°]
Amide $\text{C}_6\text{H}_4(\text{NO}_2)(\text{SO}_2\text{NH}_2)(\text{CO}_2\text{H})$ [172°]

Formed by oxidising *p* nitro toluene sulphonic amide with K_2FeO_4 or alkaline KMnO_4 (Nojes, *Am* 8, 171, 11, 161) Small granules (from hot water), melting at 177° when slowly heated, but at 172° when quickly heated Changes slowly when kept at 180° into the anhydride

$\text{C}_6\text{H}_4(\text{NO}_2) \left\langle \begin{smallmatrix} \text{SO} \\ \text{CO} \end{smallmatrix} \right\rangle \text{NH}$ [209°] which yields the salts $\text{KC}_6\text{H}_4\text{N}_2\text{SO}_3$, S 96 at 48.5° , $\text{Ba}(\text{C}_6\text{H}_4\text{N}_2\text{SO}_3)_2$ 3 aq, and $\text{AgC}_6\text{H}_4\text{N}_2\text{SO}_3$. The free amic acid yields the salts $\text{Ba}(\text{C}_6\text{H}_4\text{N}_2\text{SO}_3)_2$ aq, $\text{AgC}_6\text{H}_4\text{N}_2\text{SO}_3$ 1 aq, and $\text{Ag}_2\text{C}_6\text{H}_4\text{N}_2\text{SO}_3$ aq

Nitro *m* sulphobenzoic acid
 $\text{C}_6\text{H}_4(\text{NO}_2)(\text{SO}_3\text{H})(\text{COH})$ [τ 3 1] Formed by nitrating *m* nitrobenzoic acid (Lumprecht & Usler, *A* 106, 27) Crystalline — BaA 3 aq — BaA 1½ aq — BaH_2A 2 aq radiating prisms

Nitro *p* sulphobenzoic acid
 $\text{C}_6\text{H}_4(\text{NO}_2)(\text{SO}_3\text{H})(\text{CO}_2\text{H})$ [2 4 1] Formed by oxidising (2, 1, 1) nitro toluene sulphonic acid (Hart *Am* 1, 352) — KHA — BaA 2 aq granules

Nitro *p* sulphobenzoic acid
 $\text{C}_6\text{H}_4(\text{NO}_2)(\text{SO}_3\text{H})(\text{CO}_2\text{H})$ [3 4 1] [181°] Formed by nitrating *p* sulphobenzoic acid (Hart, *Am* 1, 342, Remsen, *A* 178, 288) Prisms — KHA 1½ aq — BaA 4 aq — BaH_2A 5 aq — CaA 5 aq — CuA 5 aq bluish green crystals

NITRO SULPHO TOLUIC ACID $\text{C}_6\text{H}_4\text{NSO}_3$
 $\text{C}_6\text{H}_4\text{Me}(\text{NO}_2)(\text{SO}_3\text{H})(\text{CO}_2\text{H})$ [1 6 4 3] Formed by oxidation of nitro *m* xylene sulphonic acid $\text{C}_6\text{H}_4\text{Me}(\text{NO}_2)(\text{SO}_3\text{H})$ [1 3 6 4] with KMnO_4 (Lumprecht, *B* 18, 2191) — AK 1 aq fine white silky needles

NITRO SULPHYDRO CINNAMIC ACID
 $\text{C}_6\text{H}_4(\text{NO}_2)\text{CHC}(\text{SH})\text{CO}_2\text{H}$ [240°] Formed from $\text{C}_6\text{H}_4(\text{NO}_2)\text{CHC}(\text{SH})\text{CO}_2\text{SH}$, nitric acid and H_2SO_4 (Bondzynski, *M* 8, 355) Crystals (from alcohol) — BaA long needles

NITRO α -SULPHYDRO CINNAMOYL SULPHOCYANIDE $\text{C}_6\text{H}_4(\text{NO}_2)\text{CHC}(\text{SH})\text{COSCN}$
 The α -compound [189°] is formed, together with the *p*-derivative [251°] by nitrating sulphydro cinnamoyl sulphocyanide These bodies may

also be got from thioglycoyl sulphocyanide and the corresponding nitro- benzoic aldehyde (Bondzynski, *M* 8, 355) Both are crystalline

NITRO-SULPHYDRO METHYL IMIDAZOLE
Methyl derivative $\text{C}_6\text{H}_4\text{Me}(\text{NO}_2)\text{SMe}$ [85°]

Formed from $\text{NM} \left\langle \begin{smallmatrix} \text{CH} \\ \text{C(SMe)N} \end{smallmatrix} \right\rangle \text{CH}$ and dilute

HNO_3 (Wohl & Marekwald, *B* 22, 1358) Yellow needles, sl sol cold water — B'H.PtCl_4 [197°]

NITRO SULPHYDRO PHENYL IMIDAZOLE
Methyl derivative $\text{C}_6\text{H}_5\text{Ph}(\text{NO}_2)\text{SMe}$

[116°] Formed from $\text{NPh} \left\langle \begin{smallmatrix} \text{CH} \\ \text{C(SMe)N} \end{smallmatrix} \right\rangle \text{CH}$ and dilute HNO_3 (Wohl & Marekwald, *B* 22, 1357)

NITROSYL BROMIDE, CHLORIDE, and SULPHATE *v* Nitrogen oxybromides and oxychlorides, *p* 569, and Nitro sulphamic acid, vol iv, *p* 601

NITRO TEREPHTHALIC ACID $\text{C}_6\text{H}_4\text{NO}_2$
 $\text{C}_6\text{H}_4(\text{NO}_2)(\text{CO}_2\text{H})_2$ [2 4 1] [270°] Formed by nitrating terephthalic acid (De la Rue & Muller, *A* 121, 90, Burkhardt, *B* 10, 145) Crystalline — A , A' powder (Skraup, *M* 7, 148)

Methyl ether MeA [70°] Prisms (Ahrens, *B* 19, 1636)

Amide $\text{C}_6\text{H}_4(\text{NO}_2)(\text{CONH}_2)_2$ Prisms

NITRO TEREPHTHALIC ALDEHYDE
 $\text{C}_6\text{H}_4(\text{NO}_2)(\text{CHO})_2$ [86°] Formed from terephthalic aldehyde, KNO_3 , and H_2SO_4 at 110° (Löw, *A* 231, 364) Large needles (by sublimation) With acetone and NaOH at 60° it gives the indigo reaction

NITRO THIENOL $\text{C}_6\text{H}_3\text{S}(\text{NO}_2)(\text{OH})$ [116°]
 Formed by treating a dilute H_2SO_4 solution of amido thiophene with nitrous acid and boiling for some time Colourless needles Sol water and ether Dissolves in alkalis with a yellow colour (Stadler, *B* 18, 2319)

NITRO-(α)-THIENYL-GLYOXYLIC ACID
 $\text{C}_6\text{H}_4(\text{NO}_2)\text{S CO CO}_2\text{H}$ [92°] Formed by oxidation of nitro thienyl methyl ketone [123°] (Peter, *B* 18, 541) Crystals

NITRO (a) THIENYL METHYL KETONE
 $\text{C}_6\text{H}_4(\text{NO}_2)\text{S CO CH}_3$ Two isomerides [86°] and [123°] are formed by nitrating (a) thienyl methyl ketone with fuming HNO_3 at -8° (Peter, *B* 17, 2646, 18, 541) They both yield the same di nitro thienyl methyl ketone [167°]

***m* NITRO THIOBENZOIC ALDEHYDE**
 $(\text{C}_6\text{H}_4(\text{NO}_2)\text{CHS})_2$ A grey powder formed by passing H_2S into an alcoholic solution of *m* nitro benzoic aldehyde (Bertagnini, *A* 79, 269) Insol. ordinary solvents

NITRO THIONYL ANILINE
 $\text{O}_2\text{H}_4(\text{NO})\text{N}(\text{SO})$ Formed from nitro-aniline and thionyl chloride (Michaelis & Humme, *B* 24 755) The *m* compound melts at 63.5° , the *p* isomeride at 70° Decomposed by hot water

NITRO THIOPHENE $\text{C}_6\text{H}_4(\text{NO}_2)\text{S}$ [44°] (225° cor) Prepared by passing air charged with thiophene vapour through fuming HNO_3 (Meyer & Stadler, *B* 17, 2648, 18, 533) Pale-yellow monoclinic prisms

Di nitro thiophene $\text{C}_6\text{H}_2(\text{NO}_2)_2\text{S}$ [52°] (290°) Formed, together with the isomeride [78°], by further nitration of nitro thiophene (Meyer & Stadler, *B* 17, 2648, 2779, 18, 530, 1778) Yellow monoclinic plates, *m* sol water Converted by repeated steam distillation into the isomeride [78°] A drop of KOH aq added to the alcoholic solution gives a splendid red colour, destroyed by excess of KOH . Forms the

double compounds $C_6H_4(NO_2)_2SC_6H_5$ [50°] and $C_6H_4(NO_2)_2SC_6H_4$ [162°]

Di-nitro-thiophene [78°] Formed as above
Yellow needles, volatile with steam

NITRO THIOPHENE SULPHONIC ACID
 $C_6H_4(NO_2)(SO_3H)_2$ Formed from nitro thiophene and fuming H_2SO_4 (Stadler, B 18, 534)
White hygroscopic crystals—AgA'

Chloride Oil

Amide $C_6H_4(NO_2)_2S(SO_2NH_2)$ [178°]

NITRO (a) THIOPHENIC ACID $C_6H_4NSO_3$, α $C_6H_4(NO_2)(CO_2H)$ [146°] Formed from (a) thiophenic acid and conc HNO_3 at 50° (Römer, B 20, 116) Needles, slowly changed by water into a variety melting at about 125° A little NaOH colours its alcoholic solution magenta CuA' —AgA' needles

Ethyl ether EtA' [71°]

NITRO THYMOL $C_6H_4Me(C_6H_4)(NO_2)(OH)$ [140°] Formed by oxidation of nitroso thymol (R Schiff, B 8, 1501, Liebermann, B 10, 612)

Di nitro-thymol [55°] Formed by nitration of thymol or its sulphonic acid (Lallemand, A Ch [3] 49, 152) Crystalline—KA'—BaA', 3aq—CaA', 6aq—AgA' lemon yellow pp

Ethyl ether EtA [53°] Tables (from alcohol) (Ladenburg & Engelbrecht, B 10, 1218)

Tri-nitro-thymol [111°] Got by nitrating di nitro thymol Yields a methyl ether [92°] (Atochley, Z 1871, 415)

NITRO-TOLUAMIDOXIM $C_6H_4N_2O_2$, α $C_6H_4Me(NO_2)C(NO_2)(NH_2)$ [4 2 1] [161°]

Formed by heating nitro toluic nitrile with alcoholic hydroxylamine (Weise, B 22, 2430)
Needles—B'HCl white crystalline mass

o-NITRO-TOLUENE $C_6H_4Me(NO_2)$ [1 2] Mol w 137 (218°) SG 1.168 (Streng, B 24, 1987) SV 1423 (Lossen, A 254, 73) Formed, together with the *p* isomeride, by nitration of toluene (Glénard & Boudault, C R 19, 505, Hofmann & Muspratt, A 53, 221, Kekulé, Z [2] 3, 225, Rosenstiehl, A Ch [4] 27, 433) Formed also by elimination of NH_3 from (2, 1, 4) nitro *p* toluidine (Beilstein & Kuhlberg, A 155, 1, 158, 348) Liquid, solidifies at -10.5° After administration to dogs it appears in the urine as *o*-nitro benzoic acid and crystalline $C_6H_4N_2O_6 \cdot 2\frac{1}{2}aq$ (Jaffé, Russ Zeit Pharm 1878, 513, Noyes, Am 5, 99) Long boiling with alkaline K_2FeO_4 yields *o*-nitro benzoic acid Zinc dust and alcoholic NaOH reduce it to *o*-azoxy toluene [59°] (Gutermann, B 20, 2016) Its product of sulphonation differs from that of *p*-nitro toluene in giving no red colour when boiled with alkalis (Reverdin & Harpe, BI [2] 50, 44)

m-Nitro-toluene $C_6H_4Me(NO_2)$ [1 3] [16°] (230°) SG 2.1168 SV 1440 Occurs in small quantity in crude nitro toluene (Monnet, Reverdin, A Nöling, B 12, 445, 18, 1337) Prepared from *o*- or *p*-toluidine by successive acetylation, nitration, saponification, diazotisation, and boiling with alcohol (Beilstein & Kuhlberg, A 155, 24, 158, 346, Buchka, B 22, 829) Yields *m*-nitro-benzoic acid on oxidation $SnCl_4$ in $HClAq$ reduces it to pure *m* toluidine, while zinc dust and $HClAq$ yield chloro *m*-toluidine also Boiling with KOH in MeOH forms $(C_6H_4Me)_2N_2O$ [39°]

p-Nitro-toluene $C_6H_4Me(NO_2)$ [1 4] [54°] (234°) (Streng, S.V.S. 121-7 (Schiff, A 223, 261).

Formed, together with the *o*-isomeride, by nitration of toluene Trimetric crystals Much less volatile with steam than *o*-nitro-toluene Oxidised by boiling alkaline K_2FeO_4 to *p* nitro-benzoic acid (Noyes, B 16, 52) Reduced by iron and $HClAq$ to pure *p*-toluidine, while zinc and $HClAq$ yield chlorinated toluidine Zinc dust and NaOH reduce it to $(C_6H_4Me)_2N_2$ [144°], two azoxy compounds $(C_6H_4Me)_2N_2O$ [75°] and [70°] and $(C_6H_4Me)_2N_2H_2$ [126°] (Janovsky & Reimann, B 22, 40) CrO_3Cl_2 followed by water yields nitro toluquinone (Etdard, C R 87, 989) NaOMe yields a brownish red product reduced by $SnCl_4$ to $C_6H_4(C_6H_4NH_2)_2$ (Bender & Schultz, B 19, 3237)

ω-Nitro-toluene $C_6H_4CH_3(NO_2)$ *Phenyl-nitro-methane* Formed by the action of acids upon the di sodio-derivative of nitro benzyldiene

phthalide $C_6H_4 \begin{matrix} \diagup C(ONa) \\ \diagdown C(Na)NO_2 \end{matrix} \begin{matrix} \diagup \\ \diagdown \end{matrix} C_6H_4$ (Ga-

briel & Koppe, B 18, 1254, 19, 1145) Liquid, boiling with decomposition at 226° Reduced by tin and HCl to benzylamine By heating with fuming HCl at 150° it yields benzoic acid and hydroxylamine Gives a white crystalline sodium derivative

Di-nitro-toluene $C_6H_4Me(NO_2)_2$ [1 2 4] Mol w 182 [70°] (300°) SVS 1375 (Schiff) S (CS₂) 219 at 17° Obtained by nitration of toluene (Deville, A 44, 307, Cahours, C R 24, 555, Nöling & Witt, B 18, 1336) and *o* or *p* nitro toluene Obtained also by eliminating NH_3 from di-nitro toluidine [195°] (Staedel, A 239, 220) Monoclinic needles, al sol cold alcohol Yields tolylene *m* diamine on reduction (Baeyer, B 7, 1638) By partial reduction with ammonium sulphide the *p*-nitro-group is reduced, giving $C_6H_4Me(NO_2)(NH_2)$ [1 2 4] of melting point [78°] On the other hand, by alcoholic $SnCl_4$ (3 mols) the *o* nitro-group is first reduced, giving $C_6H_4Me(NH_2)(NO_2)$ [1 2 4] of melting point [107°] (Anschütz & Heusler, B 19, 2161)

s-Di-nitro-toluene $C_6H_4Me(NO_2)_2$ [1 3 5] [93°] Formed by eliminating NH_3 from di nitro-*o* toluidine [208°] or di nitro *p* toluidine [166°] (Staedel, B 14, 901, A 217, 189, Neville & Winther, C J 41, 416, Hubner, A 222, 75) Yellow needles (from HOAc mixed with benzene), v sol benzene and alcohol Volatile with steam Yields di nitro benzoic acid [204°] on oxidation

Di-nitro-toluene $C_6H_4Me(NO_2)_2$ [1 2 3] [63°] Formed by heating (2, 3, 1, 4) di nitro toluic acid with dilute $HClAq$ (Rozanski, B 22, 2681)
Hair-like needles (from dilute HOAc)

Di-nitro-toluene $C_6H_4Me(NO_2)_2$ [1 2 5] [48°] (N & L), [52°5] (R) Formed by heating (1, 4, 3, 6)-di nitro toluic acid with dilute $HClAq$ at 250° (Rozanski, B 22, 2879) Formed also from toluquinone dioxim and HNO_3 (Nietzki & Gutermann, B 21, 428) Yellow crystals (from ligroin) Appears also to occur among the products of the nitration of toluene (Lumprecht, B 18, 1402)

Di-nitro-toluene $C_6H_4Me(NO_2)_2$ [1 2 6] [61°] Formed by eliminating NH_3 from di nitro *p* toluidine [168°] (Staedel, A 217, 206, 225, 884) Formed also by nitration of toluene Needles (from alcohol).

Di-nitro toluene $C_6H_4Me(NO_2)_2$ [1 3 4]? [60°]
S (CS₂) 2 188 at 17° A product of nitration of *m*-nitro toluene (Beilstein a Kuhlberg) Long needles (from CS₂)

(a) **Tri-nitro-toluene** $C_6H_3Me(NO_2)_3$ [1 2 4 6]
 Mol w 227 [82°] **S** (CS₂) 386 at 17° Formed by nitration of toluene (Wilbrand, A 128, 178) Flat needles, v sol hot alcohol Forms with aniline a compound $C_6H_4Me(NO_2)_2 \cdot PhNH_2$ [84°] (Hepp, A 215, 365)

(β) **Tri nitro toluene** $C_6H_3Me(NO_2)_3$ [112°]
 Formed, together with the (γ) isomeride, by nitration of *m* nitro toluene (Hepp) Triclinic prisms (from acetone)

(γ) **Tri nitro toluene** $C_6H_3Me(NO_2)_3$ [104°]
 Formed as above (Hepp) Trimetric plates, $a b c = 937.1 \ 672 \ V$ sol cold alcohol

NITRO-TOLUENE ω PHOSPHONIC ACID

$C_6H_4(NO_2)_2CH_2PO(OH)_2$ Formed by dissolving toluene ω phosphonic acid in fuming HNO₃ (Lutthauer, B 22, 2144) Yellow needles, decomposing at 217° without melting The acid $(C_6H_4(NO_2)_2CH_2)PO(OH)_2$ melts at 212°

NITRO-TOLUENE SULPHONIC ACID

$C_6H_4Me(NO_2)_2SO_3H$ Formed by reduction of $C_6H_4Me(NO_2)_2SO_3Cl$ by sodium amalgam (Otto a Gruber, A 145, 24) Crystalline —NaA' $\frac{1}{2}$ aq

Di nitro toluene sulphonic acid

$C_6H_3(CH_3)(NO_2)_2SO_3H$ Formed by reduction of di nitro toluene sulphonic chloride with zinc dust (Perl, B 18, 71) V sol water and alcohol

Salts —A'K.—A'Ba.—A'Pb 3aq minute prisms

o NITRO-TOLUENE SULPHONIC ACID

$C_6H_4Me(NO_2)(SO_3H)$ [125] Formed from $C_6H_4Me(NO_2)(NH_2)(SO_3H)$ [1 2 4 5] by heating its diazo derivative with alcohol at 100° (Foth, A 230, 305)

Chloride $C_6H_4Me(NO_2)(SO_3Cl)$ [50°].

Amide [133 5°] Long needles

o Nitro toluene sulphonic acid

$C_6H_4Me(NO_2)(SO_3H)$ [1 2 4] Formed by sulphonating *o* nitro toluene or by nitrating toluene *p* sulphonic acid (Beilstein a Kuhlberg, A 155, 18, Engelhardt a Bek, Z [2] 5, 209, Kornatzki, A 221, 180)

Salts —BaA', 2aq **S** (of BaA') 58 at 19 5° —PbA', 2aq **S** (of PbA') 77 at 18°

Chloride Oil (Otto a Gruber, A 145, 23)

Amide [128°] (O a G), [139°] (K), [144°] (Neale, A 203, 73) Yields a benzoyl derivative $C_6H_4Me(NO_2)_2SO_2NHBz$ [180°] whence the salts $C_6H_4Me(NO_2)_2SO_2NKBz$, $Ca(C_6H_4N_2SO_3)_2$ 2aq and $Ba(C_6H_4N_2SO_3)_2$ may be prepared, and whence PCl₅ produces $C_6H_4Me(NO_2)_2SO_2NCClPh$ [125°] from which ammonium carbonate forms $C_6H_4N_2SO_3$ [128°] (Anna Wolkoff, Z 1871, 422, B 5, 141)

p Toluide [181°] Crystals

o-Nitro-toluene sulphonic acid

$C_6H_4Me(NO_2)(SO_3H)$ [1 2 3, 5 or 6] Formed from (2, 1, 4) nitro toluidine by sulphonating and eliminating NH₂ (Foth) Its salts are v e sol water

Chloride [50°] Thick prisms

Amide [135°] Needles

o-Nitro toluene sulphonic acid

$C_6H_4Me(NO_2)(SO_3H)$ [1 2 or 6 8] Formed from *p*-toluidine sulphonic acid by nitration and elimination of NH₂ (Peschmann, A 178, 314, Foth,

A 280, 308) —BaA', 2aq plates, al sol cold water

Chloride $C_6H_4Me(NO_2)SO_3Cl$ [58 5°].

Amide $C_6H_4Me(NO_2)SO_3NH_2$ [163 5°]

m Nitro-toluene sulphonic acid Formed by sulphonating *m* nitro toluene (B a K) —BaA', 2aq **S** (of BaA') 1 145 at 17 5° —PbA', 2 $\frac{1}{2}$ aq **S** (of PbA') 3 62 at 18°

p-Nitro toluene sulphonic acid
 $C_6H_4Me(NO_2)(SO_3H)$ [1 4 2] [134°] **S** 210 at 23°, 250 at 28° Formed by sulphonating *p* nitro toluene (Javorsky, Z 1865, 222, B a K, Jenssen, A 172, 230, Hart a Remsen, B 10, 1046, Am 1, 349, Schwanert, A 186, 351, Noyes, Am 8, 168, Hauser, Bl [3] 3, 797) Trimetric crystals (containing 2 $\frac{1}{2}$ aq) NaOH aq yields an azoxy compound reduced by zinc dust to di amido stilbene disulphonic acid

Salts —NH₄' long prisms, not decomposed by H₂SO₄ at 100° —NaA' 2aq —KA' **S** 2 62 at 16° —BaA', 3aq **S** 3 34 at 18 5° —CaA', 4aq —CaA', aq —CaA', 6aq —PbA', 3aq **S** (of PbA') 15 3 at 19° —PbA', 2aq

Chloride [44°] Tablets (from ether)

Amide [187°] Needles

Nitro-toluene exo-sulphonic acid

[1 4] $C_6H_4(NO_2)CH_2SO_3H$ Formed by nitration of $C_6H_5CH_2SO_3H$ (Mohr, A 221, 217)

Chloride Oil When gently distilled it gives SO₃ and $C_6H_4(NO_2)CH_2Cl$ [1 4] [71 5°]

Amide [204°] Prisms Accompanied by an isomeride [140°–160°]

Nitro-toluene disulphonic acid

$C_6H_4Me(NO_2)(SO_3H)_2$ Formed by boiling *p* bromo toluene disulphonic acid with fuming HNO₃ (Kornatzki, A 221, 198) —K₂A' The same, or an isomeric acid, is obtained by displacing Br by H and nitrating the product, it gives the salts K₂A' and BaA' 3aq

Di-nitro-toluene exo-sulphonic acid

$C_6H_3(NO_2)_2CH_2SO_3H$ Formed from the acid $C_6H_4(NO_2)CH_2SO_3H$ and a mixture of H₂SO₄ (2 pts) and fuming HNO₃ (1 pt) (Mohr, A 221, 225) —KA' —BaA', 4aq —PbA', 4aq

Di-nitro-toluene sulphonic acid

$C_6H_4Me(NO_2)_2SO_3H$ [1 2 6 4] [165°] Formed by nitrating toluene *p* sulphonic acid (Schwanert, B 10, 28, A 186, 342) Trimetric prisms (containing 2aq) —KA' **S** 52 at 14 5° **S** (94 p o alcohol) 09 at 22° —NH₄' **S** 4 2 at 18° —BaA', 4aq **S** 3 at 17° —CaA' 2aq —PbA', 2aq —PbA' 3aq **S** (of PbA') 2 64 at 14 5° (B a K, Z [2] 6, 796)

Chloride [125°] Crystals (from ether)

Amide [203°] Laminæ

(a) NITRO-*o*-TOLUIC ACID $C_6H_4NO_2$ etc

$C_6H_4Me(NO_2)(CO_2H)$ [2 5 1] [179°] Formed, together with the (β) acid, by nitration of *o* toluic acid, and, together with the (γ) acid by oxidation of nitro-*o*-xylene with dilute HNO₃ (Jacob sen, B 16, 1957, 17, 162) Small crystals, v. al sol water Yields amido toluic acid [195°] and oxy toluic acid [179°] —KA' aq —CaA', 2aq —BaA', 2aq slender needles

(β) Nitro-*o*-toluic acid

$C_6H_4Me(NO_2)CO_2H$ [2 5 1] [145°] Formed as above. Long needles Yields oxy toluic acid [183°] —BaA', 2aq —CaA', 2aq

(γ)-Nitro-*o*-toluic acid

$C_6H_4Me(NO_2)CO_2H$ [2 4 1]. [152°]. Formed as

above Long needles Yields oxy toluic acid [179°] —BaA', 5aq easily soluble prisms

(a) Nitro-*m*-toluic acid

$C_6H_4Me(NO_2)CO_2H$ [3 6 1] [219°] Formed, together with a small quantity of its (3, 2, 1)-isomeride [182°], by nitration of *m* toluic acid (Jacobsen, *B* 14, 2353, Ahrens, *Z* 1869, 183, Krausler, *Z* 1866, 370, Panatovic, *J pr* [2] 33, 64) Monoclinic prisms —BaA', 2aq —CaA', 4aq m sol water

(β) Nitro-*m*-toluic acid $C_6H_4Me(NO_2)CO_2H$ [3 2 1] [182°] Formed as above (Jacobsen)

s Nitro-*m*-toluic acid

$C_6H_4Me(NO_2)CO_2H$ [3 5 1] [167°] Formed by oxidation of *s*-nitro *m*-xylene with $KMnO_4$ and acetic acid (Thöl, *B* 18, 360) Silky needles, v sol water —BaA', 4aq *S* 308 at 15° Needles

Nitro *m* toluic acid

$C_6H_4Me(NO_2)CO_2H$ [3 4 1] [214°] Formed by oxidation of crude nitro-xylene (Beilstein a Kreusler, *A* 144, 168, Remsen a Kuhara, *Am* 3, 426) and of nitro isocymene (Kelbe, *A* 221, 161) —NH₄A' 2aq —MgA' 7aq —CaA', 2aq —BaA', 4aq very soluble needles

Ethyl ether EtA' [55°] (B a K)

Amide [151°] (B a K)

Nitrile $C_6H_4Me(NO_2)CN$ [80°].

Nitro-*p*-toluic acid

$C_6H_4Me(NO_2)(CO_2H)$ [4 3 1] [190°] Formed by boiling cymene or *p*-toluic acid with fuming HNO_3 (Noad, *A* 63, 297, Fittica, *A* 172, 309, Fittig, *A* 168, 251, Ahrens, *Z* [2] 5, 102) Monoclinic prisms, sl sol cold water —BaA', 4aq —CaA', 3aq —CuA', 4aq —Cu₂A', (OH) —Cu₂A', (OH) aq (Noyes, *Am* 10, 472) —Pb(OH)A' —AgA' The ethers MeA' and EtA' are crystalline

Nitro-*p*-toluic acid

$C_6H_4Me(NO_2)(CO_2H)$ [4 2 1] [161°] Obtained by heating its nitrile with HCl aq at 195° Long needles —BaA', 4aq —BaA', 5aq (Noyes, *Am* 10, 472) —CaA', 2aq —CuA', aq —AgA' needles

Amide [153°] Slender needles

Nitrile $C_6H_4Me(NO_2)CN$ [99°] (G, W),

[101°] (N) Formed from (3, 1, 4) nitro-*p*-toluidine by Sandmeyer's reaction (Glock, *B* 21, 2662, Weiss, *B* 22, 2429, Von Niementowski, *J pr* [2] 40, 4, 21, 1535, 1992) Needles (from alcohol) Yields on reduction with tin and HCl the compounds (C_6H_4MeCy)₂N₂O [182°] and $C_6H_4Me(NH_2)CN$ [94°] Does not form an imido-ether when treated with alcoholic HCl (Pinner, *B* 23, 2919)

A nitro toluic acid [218°] was obtained by Ahrens together with the acids [219°] and [190°] by the action of HNO_3 on crude xylene

Di-nitro-*o* toluic acid

$C_6H_4Me(NO_2)_2CO_2H$ [2 5 3 1] [206°] Formed by nitrating *o* toluic acid (Jacobsen a Wierss, *B* 16, 1957, Racine, *A* 239, 77) Needles Yields di nitro-phthalic acid [226°] on oxidation —BaA', 2aq v e sol water

Methyl ether MeA' [74°] Needles

Di-nitro-*p* toluic acid

$C_6H_4Me(NO_2)_2CO_2H$ [4 3 5 1] [158°] Formed by nitrating *p* toluic acid (Brückner, *B* 8, 1678) Plates (from hot water) —KA' 2aq —CaA', 2aq —BaA', 2aq —AgA'

Di-nitro *p* toluic acid

$C_6H_4Me(NO_2)_2CO_2H$ [4 2 5 1] [188°] Formed, together with the isomeride [249°], by nitrating

(2, 4, 1)-nitro toluic acid (Bozansky, *B* 22, 2676) Radiating needles Yields (1, 4, 3) di-nitro toluene on heating with dilute HCl at 250° —NaA' 3aq —BaA', 2aq —CaA', 2aq

Di nitro-*p*-toluic acid

$C_6H_4Me(NO_2)_2CO_2H$ [4 2 3 1] [249°] Formed as above (R) Trimeric prisms —CaA', aq —BaA', 4aq long needles

NITRO-*m*-TOLUIC ALDEHYDE $C_6H_4Me(NO_2)CHO$ An oil, formed as well as di nitro *m* toluic aldehyde [112°], by nitrating *m* toluic aldehyde (Bornemann, *B* 17, 1473)

NITRO-*o*-TOLUIDINE

$C_6H_4Me(NH_2)(NO_2)$ [1 2 8] [97°] Obtained from acetyl-*o*-toluidine by nitration and saponification (Lellmann a Wurthner, *A* 228, 240), and also by heating its sulphonic acid with dilute H_2SO_4 at 180° (Nietzki a Polln, *B* 28, 138) Prisms (from dilute alcohol)

Acetyl derivative [158°] Plates

Nitro-*o*-toluidine $C_6H_4Me(NH_2)(NO_2)$ [1 2 4] [107°] Obtained by nitrating *o* toluidine (1 pt) dissolved in H_2SO_4 (10 pts) (Nölting a Collin, *B* 17, 268) and by reduction of (4, 2, 1) di nitro toluene [70 5°] (Graeff, *A* 229, 343, Lumprecht, *B* 18, 1400, Anschütz, *B* 19, 2161) Orange monoclinic prisms Sweet taste —B', H_2SO_4 , plates, decomposed by water

Acetyl derivative [151°] Needles

Nitro-*o* toluidine $C_6H_4Me(NH_2)(NO_2)$ [1 2 5] [128°] Obtained from its acetyl derivative which is got by nitrating acetyl-*o* toluidine (Beilstein a Kuhlberg, *A* 158, 345) Small lemon yellow needles (from water)

Acetyl derivative [197°] Needles

Nitro *o* toluidine $C_6H_4Me(NH_2)(NO_2)$ [1 2 6] [92°] Formed by reduction of (6, 2, 1) di nitro toluene (Cunrath, *A* 172, 223, Ullmann, *B* 17, 1957) Formed also, together with the (1, 2, 4) isomeride, by nitrating *o* toluidine in presence of a large excess of H_2SO_4 (Green a Lawson, *priv com*) Bright yellow slender needles —B'HCl

Acetyl derivative [158°] Needles

Benzoyl derivative [167°]

Nitro *m* toluidine $C_6H_4Me(NH_2)(NO_2)$ [1 3 6] [184°] Formed from its acetyl derivative, which is got by nitrating acetyl *m* toluidine (Beilstein a Kuhlberg, *A* 158, 348) Obtained also by heating the ethyl ether [54°] of nitro cresol [129°] with NH_3 aq at 150° (Staedel, *A* 209, 214) Needles, v sol alcohol Yields tolylene *p* diamine [64°] on reduction (Fileti a Crosa, *G* 18, 298)

Acetyl derivative [102°] Cubes

Nitro *m*-toluidine $C_6H_4Me(NH_2)(NO_2)$ [1 3 4] [109°] Formed by heating the ethyl ether [51°] of nitro-cresol [56°] with NH_3 aq for 8 hours at 150° (Staedel, *A* 259, 225) Golden plates, m sol alcohol

s Nitro *m* toluidine

$C_6H_4Me(NH_2)(NO_2)$ [1 3 5] [98°] Formed from *s* di nitro toluene (Becker, *B* 15, 1138, Neville a Winther, *C J* 41, 416, Staedel, *A* 217, 199) Needles —B'HCl [56°] Prisms

Benzoyl derivative [177°]

Nitro *m* toluidine $C_6H_4Me(NH_2)(NO_2)$ [1 3 2] [58°] Formed by reducing (2, 3, 1) di nitro-toluene (Lumprecht, *B* 18, 1401) —B'HCl —B', H_2SO_4 , tables, v sol hot water

Acetyl derivative [186°] Needles

Nitro-*p*-toluidine $C_6H_4Me(NH_2)(NO_2)$ [1 4 3].

[116°] Formed from acetyl-*p* toluidine by nitration and hydrolysis (Beilstein, A Kuhlberg, A 155, 23, Lorenz, A 172, 177, Hubner, A 208, 813, Cosack, B 13, 1088, Ehrlich, B 15, 2009, Gattermann, B 18, 1483, Lellmann, A 221, 7, Nöling, B 17, 263) Formed also by heating nitro-*p* cresol with NH_4Aq at 180° (Barr, B 21, 1543) Red monoclinic prisms, $a b c = 1.358 : 1.755$, $\beta = 54^\circ 51'$ (Panebianco, G 9, 358) — B'HCl — B'HNO_3 crystals, decomposed by water

Acetyl derivative [95°] Needles Reduced by means of ammonium sulphide at 0° to $\text{C}_6\text{H}_4\text{Me}(\text{NHAq})_2 \cdot \text{N}_2\text{O}$ [196°] and the compound $\text{C}_6\text{H}_4\text{Me} \begin{smallmatrix} \text{N} \\ \text{NH CMe} \end{smallmatrix} \text{O}$ [234°], whence B'HCl , B'H PtCl_4 , and B'HNO_3 (Bankievitch, B 22, 1396)

Trichloroacetyl derivative [55°].

Valeryl derivative [88°] Yields on reduction $\text{C}_6\text{H}_4\text{Me} \begin{smallmatrix} \text{N} \\ \text{NH} \end{smallmatrix} \text{C}_6\text{H}_5$ [145°]

Benzoyl derivative [143°] Needles

Nitro *p* toluidine $\text{C}_6\text{H}_4\text{Me}(\text{NH}_2)(\text{NO}_2)$ [1 4 2] [77 5°] Formed by reduction of (4, 2, 1) di-nitro toluene (B a K) and, together with a small quantity of the preceding isomeride, by the action of nitric acid on a solution of *p* toluidine in H_2SO_4 (Hubner, B 10, 1716, Nolting a Collin, B 17, 263, Foth, A 230, 299) Monoclinic needles — B'HCl [220°] — B'HNO_3 — B'H SO_4 2aq stellate needles — B'AgNO_3 [132°] Greenish yellow crystals (Mixer, Am 1, 241)

Acetyl derivative [144 5°] Needles (Wallach, A 234, 353)

Benzoyl derivative [172°] Pale yellow prisms (Bell, C N 30, 202)

Di-nitro-*o* toluidine

$\text{C}_6\text{H}_4\text{Me}(\text{NH}_2)(\text{NO}_2)_2$ [1 2 3 5] [208°] Formed from di nitro *o* cresol and its ethers by the action of NH_3 (Staedel, B 14, 900, A 217, 185, 203, Van Romburgh, R T C 3, 398, Barr, B 21, 1543) Yellow prisms (from xylene)

Di-nitro-*m* toluidine

$\text{C}_6\text{H}_4\text{Me}(\text{NH}_2)(\text{NO}_2)_2$ [1 3 4 6] [193°] Formed from the ethyl ether of di nitro *m* cresol and NH_4Aq at 100° (Staedel, A 259, 220), and from $\text{C}_6\text{H}_4\text{MeBr}(\text{NO}_2)_2$ and NH_3 (Jackson, B 22, 1232) Formed also by reducing (γ) tri nitro toluene (Hepp, A 215, 368) Yellow crystals, yielding di nitro toluene [71°] on elimination of NH_3 .

Di-nitro-*p* - toluidine $\text{C}_6\text{H}_4\text{Me}(\text{NH}_2)(\text{NO}_2)_2$ [1 4 3 5] [168°] S (CS₂) 32 at 18° Formed by nitration of acetyl or benzoyl *p* toluidine and hydrolysis of the product (Beilstein, B 13, 242, Hubner, A 222, 73) Formed also by the action of ammonia on the ethers of di nitro *p* cresol (Staedel, A 217, 183) Needles, sl sol alcohol Yields chrysamic acid on oxidation

Acetyl derivative [195°] Needles Yields on reduction an azoxy-compound [236°], an azo-compound [244°], and $\text{C}_6\text{H}_4\text{N}_2\text{O}_3$ [256°] The compounds $\text{C}_6\text{H}_4\text{Me}(\text{NO}_2) \begin{smallmatrix} \text{N} \\ \text{NH} \end{smallmatrix} \text{CMe}$ [246°] and

$\text{C}_6\text{H}_4\text{Me}(\text{NH}_2) \begin{smallmatrix} \text{NH CMe} \\ \text{N} \end{smallmatrix} \text{O}$ [266°] may also be got by reduction (Bankievitch, B 21, 2404)

Trichloroacetyl derivative [142°]. Prisms or needles (Friederici, B 11, 1975)

Benzoyl derivative [186°] Needles
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An isomeric body [203°] is got by nitrating benzoyl-(2,1,4) nitro toluidine (Cunert, A 172, 229).

Di-nitro-*p* - toluidine $\text{C}_6\text{H}_4\text{Me}(\text{NH}_2)(\text{NO}_2)_2$ [1 4 2 6] [168°] Formed from tri nitro-toluene [82°] and ammonium sulphide (Tiemann, B 3, 218, Beilstein, B 13, 242, Staedel, A 226, 854). Needles, v sol alcohol

Di-nitro-*p* - toluidine $\text{C}_6\text{H}_4\text{Me}(\text{NH}_2)(\text{NO}_2)_2$ [94°] Formed by heating (β) tri nitro toluene with alcoholic NH_3 at 100° (Hepp) Golden needles (from HOAc)

Tri-nitro-*m* - toluidine $\text{C}_6\text{H}_4\text{Me}(\text{NH}_2)(\text{NO}_2)_3$ [1 3 2 4 6] [136°] Formed by the action of NH_3 on the ethyl ether of tri nitro *m* cresol (Nolting a Salis, B 15, 1864, A Ch [6] 4, 128, Staedel, A 259, 222) Small cubes, sol alkalis, forming a red solution

NITRO-*p*-TOLUIDINE SULPHONIC ACID $\text{C}_6\text{H}_4\text{Me}(\text{NH}_2)(\text{NO}_2)(\text{SO}_3\text{H})$ [1 4 2 5] S 1603 at 15° Formed by sulphonating nitro-*p* toluidine (Limpicht, B 18, 2183, Foth, A 230, 298) — KA' aq — BaA' 4aq — PbA' 3 1/2 aq

DI-NITRO TOLYL-ACETIC ACID

$\text{C}_6\text{H}_4\text{Me}(\text{NO}_2)_2 \text{CH}_2 \text{CO}_2\text{H}$ [173°] Formed by nitration (Senkowsky, M 9, 856) Yields the ethers MeA' [41°] and EtA' [68°] crystallising in needles

NITRO *p*-TOLYL-AMIDO-ACETIC ACID

$[\text{1 3 4}] \text{C}_6\text{H}_4\text{Me}(\text{NO}_2) \text{NH CH CO}_2\text{H}$ [190°] Formed from nitro *p* toluidine and bromo acetic acid (Plochl, B 19, 9, Leuckhart B 20, 24) Prisms — $\text{NH}_4\text{A'}$ — BaA' 1 aq — PbA' 2 purple red needles

Ethyl ether EtA' [65°] Needles

NITRO *o* TOLYLAMIDO BENZOIC ACID

$[\text{3 4 1}] \text{C}_6\text{H}_3(\text{NO}_2)(\text{NHCH}_2\text{CO}_2\text{H}) \text{CO}_2\text{H}$ [211°] Formed by heating *o* toluidine with bromo nitrobenzoic acid (Heidensleben, B 23, 3451) Brown needles — NaA' 2 aq red needles

Ethyl ether EtA' [106°] Leaflets

***m* Nitro *p* tolyl amido benzoic acid**

$\text{C}_6\text{H}_4\text{Me NH C}_6\text{H}_4(\text{NO}_2) \text{CO}_2\text{H}$ [257°] Formed from *p* toluidine and bromo nitro benzoic acid (Schöpf, B 22, 3288, H) — NaA' dark red needles

Ethyl ether EtA' [115°] Leaflets

NITRO DI *p* TOLYL AMINE

$\text{NH}(\text{C}_6\text{H}_4)(\text{C}_6\text{H}_4\text{NO}_2)_2$ [85°] The *benzoyl derivative* [167°] is formed, together with that of di nitro di tolyl amine [191°], by nitrating benzoyl di *p* tolyl amine (Lellmann, B 15, 831)

Hexa nitro di *p* tolyl amine

$\text{NH}(\text{C}_6\text{H}_4\text{Me}(\text{NO}_2)_3)_2$ [258°] Formed from di-tolyl-nitrosamine and fuming HNO_3 (Lehne, B 13, 1545) Trimetric crystals

NITRO TOLYL-ISOBUTYRIC ACID

$[\text{1 3 6}] \text{C}_6\text{H}_4\text{Me}(\text{NO}_2) \text{CH}_2 \text{CHMe CO}_2\text{H}$ [139°] Formed from iso tolyl nitro toluene and dilute HNO_3 (S G 1 12) at 200° (Effront, B 17, 2326) — AgA' colourless plates

NITRO TOLYLENE DIAMINE $\text{C}_6\text{H}_4\text{N}_2 \text{O}_2$ *ie* $\text{C}_6\text{H}_4\text{Me}(\text{NO}_2)(\text{NH}_2)_2$ [1 2 2 6] [154°] Obtained from its *acetyl derivative* [253°], which is got by nitrating di acetyl tolylene diamine (Tiemann, B 3, 9, Ladenburg, B 8, 1211) Needles, with violet reflex Yields a *di-benzoyl derivative* [245°] (Ruhemann, B 14, 2656)

Nitro-tolylene diamine *Benzoyl derivative* $\text{C}_6\text{H}_4\text{Me}(\text{NO}_2)(\text{NH}_2)(\text{NHBz})$ [1 2 5 4] [139°]. S S

Formed by reducing benzoyl-di-nitro-toluidine (Hübner, *A* 208, 317) Red needles (from water).

Nitro tolylene diamine $C_6H_4Me(NO_2)(NH_2)_2$ [1 6 4 2] ? [132°] Formed by reducing (6,4,2,1)-tri-nitro-toluene (Tiemann, *B* 3, 218) Red prisms (from water)

Tri-nitro-tolylene-diamine

$C_6Me(NO_2)_3(NH_2)_2$ [1 2 4 6 3 5] [222°] Formed from $C_6Me(NO_2)_3Br$, and alcoholic NH_3 (Palmer, *B* 21, 3501) Small yellow prisms

NITRO-TOLYLENE-TETRA-METHYL-DIAMINE $C_6H_4Me(NO_2)(NMe)_2$ [63°] Formed by nitration (Niementowski, *B* 20, 1888) Prisms

DI-NITRO-DI-TOLYL-ETHYLENE-DIAMINE

$C_6H_4(NH C_6H_4Me NO_2)_2$ [195°] Formed from (8,1,4) nitro toluidine and ethylene bromide (Gattermann & Hager, *B* 17, 779) Red plates

NITRO-TOLYL-HYDRAZINE SULPHONIC ACID

$C_6H_4Me(NO_2)(N_2H_4)(SO_3H)$ [1 2 4 5] Formed from nitro-*p*-diazo-toluene sulphonic acid and a cooled solution of $SnCl_2$ (Lamprecht, *B* 18, 2194) Tables— BaA' , 4sq yellow prisms

NITRO-TOLYL-METHYLENE PHTHALIDE

$C_6H_4 \begin{array}{c} \diagup O=C(NO_2) C_6H_4 \\ \diagdown CO \end{array}$ [144°] Formed from tolyl-methylene-phthalide by the action of nitrous acid, the resulting $C_6H_4 \begin{array}{c} \diagup C(NO_2) CH(NO_2) C_6H_4 \\ \diagdown CO \end{array}$

[138°] being boiled with dilute alcohol (Heilmann, *B* 23, 3163) Needles

NITRO TOLYL METHYLENE-PHTHALIMIDINE

$C_6H_4 \begin{array}{c} \diagup C=C(NO_2) C_6H_4 \\ \diagdown CO NH \end{array}$ [159°] Formed from tolyl-methylene-phthalimidine and nitrous acid (Heilmann, *B* 23, 3161) Needles (from alcohol)

NITRO *m*-TOLYL-PROPIONIC ACID

$C_6H_4Me(NO_2)C_6H_4CO_2H$ [130°-136°] Formed from (2,5,1) iodo isobutyl toluene and HNO_3 (S G 1 25) at 200° Efront, *B* 17, 2327) Needles (from water)

DI NITRO DI-TOLYL-PROPIONIC ACID

$(C_6H_4MeNO_2)_2CMe CO_2H$ [129°] Formed, together with the tetra-nitro-acid [225°] by nitrating di-tolyl propionic acid (Haiss, *B* 15, 1476) Yellowish crystals

DI NITRO-DI-*o*-TOLYL-SULPHAZIDE

$C_6H_4N_2O_2S$ [2 4 1] $C_6H_4Me(NO_2) NH NH SO_2 C_6H_4(NO_2)Me$ [1 4 2] ? [142°] Obtained by the action of alcoholic SO_2 upon nitro-*o*-diazo toluene (Lamprecht, *B* 20, 1241) Small yellow prisms

NITRO-TOLYL-THIOCARBAMIC ETHER

[1 2 4] $C_6H_4Me(NO_2) NH CS OEt$ [96°] Formed by boiling with alcohol nitro-tolyl-thiocarbamide, which is produced by the action of Ac_2O on phenyl-nitro tolyl-thio-urea (Steuemann, *B* 16, 2337) Needles, v sol alcohol

NITRO-TOLYL-THIO-UREAS

The following compounds have been prepared by Steuemann (*B* 16, 2337) from $C_6H_4Me(NO_2)(NH_2)$ [1 2 4]

$NH_2CS NH C_6H_4Me(NO_2)$ [176°]

[4 1] $C_6H_4Me NH CS NH C_6H_4Me(NO_2)$ [169°]

$CS(NH C_6H_4Me(NO_2))_2$ [207°]

DI-NITRO-DI-*p*-TOLYL-UREA

$CO(NH C_6H_4NO_2)_2$ [c. 233°] Formed from di-*p* tolyl-guanidine, alcohol, and HNO_3 (S G

1 4) (A G Perkin, *C* 7, 87, 698) Needles (from xylene)

NITRO-URACIL v Nitro-DI OXY PY IMIDINE

NITRO-URAMIDO-BENZOIC ACIDS

The three following acids are obtained by boiling the three di-nitro-uramido-benzoic acids with aqueous NH_3 (Griess, *B* 5, 193) —
 $C_6H_4(NO_2)(NH CO NH_2) CO_2H$ [6 3 1] Crystals.
 $C_6H_4(NO_2)(NH CO NH_2) CO_2H$ [4 3 1] Needles.
 $C_6H_4(NO_2)(NH CO NH_2) CO_2H$ [2 3 1] Plates
 The isomeric $C_6H_4(NO_2)(NH CO NH_2) CO_2H$ [5 3 1] is formed, together with nitro di-uramido benzoic acid, by the action of potassium cyanate on nitro amido benzoic acid (Griess, *B* 17, 2184) It yields the salt BaA' , 5sq

Nitro-diuramido-benzoic acid

$(NH_2 CO)_2N C_6H_4(NO_2) CO_2H$ forms crystals (containing 2aq), and yields the salt BaA' , 7½aq
 Three crystalline di-nitro-*m*-uramido benzoic acids are got by nitrating *m* uramido benzoic acid A di nitro-*p*-uramido benzoic acid is formed by nitrating *p*-uramido-benzoic acid

NITROUS ACID v Nitrogen, p 567.

NITROUS ETHER v ETHYL NITRITE

NITRO-UVITIC ACID

$C_6H_4Me(NO_2)(CO_2H)_2$ [1 x 3 5] [227°] Formed, together with an isomeride $C_6H_4NO_2$, ½aq [250°] by nitrating uvitic acid (Böttinger, *B* 9, 804, *A* 189, 171) Prisms (containing 2aq) — K_2A'' aq — BaA'' aq — CaA'' 8aq needles, m sol hot water

NITRO VALERIC ACID $C_6H_4(NO_2) CO_2H$

Formed by the action of nitric acid on isovaleric acid and on di isomyl ketone (Dessaignes, *A* 79, 374, Bredt, *B* 15, 2319, Brazier, a Gossleth, *A* 75, 262, Schmidt, *B* 6, 602) Monoclinic tables AgA' prisms (from hot water)

DI NITRO VINYL FURFURANE

$C_6H_4(NO_2)O CH CH NO_2$ [144°] Formed by nitrating vinyl furfuran (Priebe, *B* 18, 1362) Yellow needles Yields a dibromide [111°]

NITRO VINYL PHENOL *Methyl derivative* $C_6H_4(NO_2)(CH CH NO_2)OMe$ [3 1 4]

[163°] Formed from the methyl derivative of *p* coumaric acid and HNO_3 (Einhorn a Grabfield, *A* 243, 369) Yellow needles (from alcohol)

o **NITRO-*o* XYLENE** $C_6H_4Me(NO_2)$ [1 2 4]

Mol w 151 [29°] (258°) S G ¾ 1 139 Formed by nitration of *o* xylene (Jacobsen, *B* 17, 160) Long yellow prisms

o-Nitro-*o*-xylene $C_6H_4Me(NO_2)$ [1 2 3] (250°

1 V at 739 mm) S G ½ 1 147 Formed, together with the preceding isomeride, by the action of H_2SO_4 and HNO_3 on *o* xylene (Nöling a Forel, *B* 18, 2669) Liquid

c Nitro-*m*-xylene $C_6H_4Me_2(NO_2)$ [1 3 2],

(225°) at 745 mm S G ½ 1 112 Formed from nitro xylidine [78°] by eliminating NH_3 (Grevink, *B* 17, 2430), and, together with the (1,3,4) isomeride, by nitration of *m* xylene with HNO_3 and H_2SO_4 at 0° (N a F) Liquid

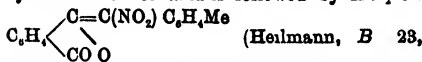
p-Nitro-*m*-xylene $C_6H_4Me_2(NO_2)$ [1 3 4] (244° cor) S V 164 5 Formed by nitrating *m*-xylene (Harmsen, *B* 13, 1558) or its dihydride (Wallach, *A* 258, 380), and by eliminating NH_3 from nitro-xylidine [128°] (G) Liquid

s-Nitro-*m*-xylene $C_6H_4Me_2(NO_2)$ [1 3 5], [75°], (263° 1 V) at 739 mm. Formed by eliminating NH_3 from nitro-xylidine [70°] (Wroblewsky, *A* 207, 94, *BI* [2] 84, 332, Thol,

B. 18, 360, Nörling & Forel, B 18, 2678)
Needles, volatile with steam

Nitro *p*-xylene $C_6H_4Me_2(NO_2)$ [1 4 2] (239°
1 V) at 739 mm SG 1⁶ 1132. Formed by
nitration of *p*-xylene (Jannasch, A. 176, 55,
N a F) Liquid.

***m*-Nitro *m*-xylene** $C_6H_4Me_2(CH_3, NO_2)$. Formed
by the action of alkalis followed by HCl₂ on



3164) Oil, with irritating odour

Di-nitro *m* xylene $C_6H_4Me_2(NO_2)_2$ [1 3 4 2]
Mol w 196 [82°] Formed, together with the
isomeride [93°], by nitrating *m* xylene with HNO₃
and H₂SO₄ at 5° (Grevingk, B 17, 2422) Plates,
v sol alcohol

Di nitro *m*-xylene $C_6H_4Me_2(NO_2)_2$ [1 3 4 6]
[93°] Formed by nitrating *m* xylene or its di-
hydride (Luhmann, A 144, 274, Fittig, A 148,
5, Wallach, A 258, 332) Crystals (from alcohol)

Di-nitro-*p*-xylene $C_6H_4Me_2(NO_2)_2$ [1 4 2 3]
[93°] Formed, together with the isomeride
[124°], by nitrating *p*-xylene (Rammer, B [2]
9, 434, Fittig, A 136, 307, 147, 17, Jannasch,
A 171, 79, Nörling, B 19, 144, Lellmann, A
228, 252) Monoclinic crystals Yields xylene-
diamine [75°]

Di nitro *p*-xylene $C_6H_4Me_2(NO_2)_2$ [1 4 2 6]
[124°] Needles

Di-nitro *p*-xylene $C_6H_4Me_2(NO_2)_2$ [1 4 2 5].
Formed in small quantity by nitrating *p* xylene
(L) Long yellow needles (from alcohol)

Tri nitro-*m* xylene $C_6H_3Me_3(NO_2)_3$ [1 3 2 4 6].
Mol w 241 [c 182°] Formed by nitration of
m xylene or its dihydride (Luhmann, Grevingk,
Tilden, C J 45, 416, Wallach, A 258, 333)
Crystals, insol hot alcohol

Tri nitro *p* xylene $C_6H_3Me_3(NO_2)_3$ [1 4 2 3 5]
[140°] Formed by nitration of *p* xylene (Fittig,
Nörling, B 19, 145). Crystals (from benzene)

NITRO *m*-XYLENE PHOSPHONIC ACID

$C_6H_4Me_2(NO_2)PO(OH)_2$. Two acids of this formula,
[100°] and [182°], are formed by nitrating
m-xylene (α)-phosphonic acid (Weller, B 20,
1722, 21, 1492) From *m* xylene (β) phosphonic
acid an isomeric acid [107°] is obtained, and
another isomeride [224°] may be obtained from
p xylene phosphonic acid

NITRO *m*-XYLENE SULPHONIC ACID

$C_6H_4Me_2(NO_2)(SO_3H)$ [1 3 6 4] [132°] Formed
by sulphonating nitro *m* xylene and by nitrating
m-xylene (α) sulphonic acid (Harmsen, B 13,
1558, Lumprecht, B 18, 2191, Claus & Schmidt,
B 19, 1418) Crystals NaA' aq—NaA' 2aq—
KA'—NH₄A'—BaA', 8½aq—CaA', 6aq S 6 35
at 18°—MgA', 9aq—CuA', 6aq—PbA', 4aq—
AgA' aq

Amide [179°] (L); [187°] (O a S)

Chloride [98°] (L)

Nitro-*m*-xylene sulphonic acid

$C_6H_4Me_2(NO_2)(SO_3H)$ [1 3 5 4] [100°] Formed,
with the preceding and succeeding acid, by
nitrating (1,3,4) xylene sulphonic acid (O a S)
Plates—KA'—NaA' aq—BaA', 1½aq—CaA', 6aq
—PbA', 4aq—CuA', 6aq—AgA' aq. needles, v.
sol. water

Amide [108°] Needles.

Chloride [97°]

Nitro-*m*-xylene sulphonic acid

$C_6H_4Me_2(NO_2)SO_3H$ [1 3 2 4] [144°] Formed
as above (O a S) Plates (containing aq)—
KA' ½aq—NaA' aq—BaA',—CaA',—PbA',—
CuA', 2aq—AgA' ½aq

Chloride [96°] Needles.

Amide [172°] Needles

Di-nitro-xylene sulphonic acid

$C_6H_4Me_2(NO_2)_2SO_3H$ [1 3 6 5 4] [70°] Formed,
as well as the following isomeride, by nitrating
m-xylene sulphonic acid (Claus & Schmidt, B.
19, 1425) Plates.—KA'—NaA' aq—BaA', ½aq.
—BaA', 2½aq—CaA', 5aq—PbA', 4½aq—
CuA', 4aq pale green plates

Chloride [118°] Crystals.

Amide [158°] Needles

Di-nitro-xylene sulphonic acid

$C_6H_4Me_2(NO_2)_2SO_3H$ [1 3 6 2 4] Needles (con-
taining 2aq) (Lumprecht, B 18, 2192, O. a. S).
—NaA' aq—KA'—BaA', 3aq—CaA', 8½aq—
CuA', 2½aq—PbA', 3½aq prisms.

Chloride [123°] Prisms

Amide [193°] Prisms

NITRO-*m*-XYLENOL $C_6H_4NO_2$ s.s.

$C_6H_4Me_2(NO_2)(OH)$ [1 3 x 4] [68 5°] Formed
by nitrating *m*-xynolol (Lako, A 182, 32)
Needles—KA' 3aq dark red plates

Nitro *m* xynolol $C_6H_4Me_2(NO_2)(OH)$ [95°]
Formed by the action of nitrous acid on the
nitro xylidine obtained by partial reduction of
di nitro *m* xylene (Pfaff, B 16, 616, 1136)
Needles—KA' 2aq red crystals

Methyl ether MeA' [57°] Needles

(α) Nitro *p*-xynolol

$C_6H_4Me_2(NO_2)(OH)$ [1 4 6 3] [115°] (O), [122°]
(G a S) Formed by oxidising nitroso-*p*-xylene-
ol (the oxum of phlorone) with alkaline K₂FeCy₄.
(Olver, G 12, 162, Goldschmidt & Schmid, B.
18, 569) Needles, sol. hot water

(β)-Nitro *p*-xynolol $C_6H_4Me_2(NO_2)(OH)$.
[236°] Formed by nitrating *p* xynolol (O).
Oil—BaA', purple scales

(γ)-Nitro *p*-xynolol [89°] Formed by heat-
ing *p* xynolol sulphonic acid with fuming HNO₃.
(O) Light-yellow scales—KA' aq—BaA', aq.
light yellow scales

Nitro-*p*-xynolol $C_6H_4Me_2(NO_2)(OH)$ [1 4 3 5].
[91°] Formed by the action of nitrous acid on
the corresponding nitro xylidine (Von Kosta-
necki, B 19, 2320) Buff-coloured plates;
perhaps identical with the preceding isomeride.

Nitro-*p*-xynolol *Ethyl ether*

$C_6H_4Me_2(NO_2)(OEt)$ [85°] Formed from *p*-
xylidine by nitration and treatment with nitrous
acid (Nörling, Witt, a. Forel, B 18, 2667)

Di-nitro-*o*-xynolol

$C_6H_4Me_2(NO_2)_2(OH)$ [1 2 3 5 4] [128°] Formed
as a by-product by nitrating *o*-xylene (Nörling
& Pick, B 21, 3158) Obtained also from (1, 2, 4)-
xylidine. Needles, al. sol. cold water

Di-nitro-*o*-xynolol

$C_6H_4Me_2(NO_2)_2(OH)$ [1 2 3 4 6] [82°] Obtained
by nitrating and diazotising (1, 2, 3)-xylidine (N.
a. P.) Orange yellow needles, m. sol water

Di-nitro-*p*-xynolol [121°] Formed from *p*-
xynolol, H₂SO₄, and HNO₃ (Kostanecki, B 19,
2321) Yellow plates (from water)

NITRO-*m*-XYLENOL SULPHONIC ACID

$C_6H_4Me_2(NO_2)(OH)(SO_3H)$ [1 3 x 6 4] Formed
from nitro-xylidine sulphonic acid by the diazo
reaction (Lumprecht & Sartig, B. 18, 2190, A.

130, 340) —BaA', 3aq —PbA', 3aq The *ethyl derivative* forms $C_6HMe_2(NO_2)(OEt)(SO_3K)$ aq, crystallising in plates

NITRO-*m*-XYLIDINE $C_6H_4N_2O_2$ *ie* $C_6H_2Me_2(NO_2)(NH_2)$ [1 3 5 4] [76°] Formed from acetyl *m*-xylidine by nitration and saponification (Wroblewski, A 207, 91, Nörling a Forel, B 18, 2677) Orange needles

Acetyl derivative [178°] Needles

Nitro-*m*-xylidine

$C_6H_4Me_2(NO_2)(NH_2)$ [1 3 6 4] [123°] Formed by reduction of di nitro *m* xylene (Fitting, A 147, 18, Wallach, A 258, 332) and by nitration of 1,3,4) *m* xylidine (1 pt) dissolved in H_2SO_4 (10 pts) (Nörling a Collin, B 17, 265) Orange needles —B'HCl —B'H₂SO₄ —B'H₂C₂O₄

Acetyl derivative [160°]

Diacetyl derivative [116°].

Nitro-*s*-xylidine

$C_6H_4Me_2(NO_2)(NH_2)$ [1 3 4 5] [54°] Formed by nitration of *s*-*m*-xylidine dissolved in 10 pts of conc H_2SO_4 (Nörling a Forel, B 18, 2679) Yellow needles Volatile with steam

Nitro-*m* xylidine

$C_6H_4(CH_3)_2(NO_2)(NH_2)$ [1 3 2 4] or [1 3 4 2] [78°] Formed by reduction of di-nitro *m* xylene [82°] with alcoholic ammonium sulphide (Grevink, B 17, 2425) Yellow needles, sol hot water

Acetyl derivative [149°] White needles

Nitro-*p*-xylidine $C_6H_4Me_2(NO_2)(NH_2)$ [1 4 5 2] [142°] Formed by nitration of acetyl *p* xylidine and saponification, or by nitration of *p* xylidine dissolved in conc H_2SO_4 (Nörling, Witt, a Forel, B 18, 2666) Formed also by reducing di nitro *p*-xylene (Kostanecki, B 19, 2318) Brownish yellow crystals On reduction it gives the *para* diamine [147°]

Acetyl derivative [166°] Needles

Nitro-*p* xylidine $C_6H_4Me_2(NO_2)(NH_2)$ [1 4 3 5] [96°] Formed by reduction of di nitro *p* xylene [123°] (Fitting, A 147, 22, Kostanecki, B 19, 2320) Needles (from alcohol)

Acetyl derivative [180°] Needles

Di-nitro-*p* xylidine

$C_6H_4Me_2(NO_2)_2(NH_2)$ [1 4 3 5 2] [203°] Formed from tri nitro-*p* xylene and alcoholic NH_3 (Nörling, B 19, 145) Needles (from HOAc)

Di nitro-xylidine [192°] Formed by reduction of tri nitro-xylene (Bussenius a Eisenstuck, A 118, 165, Beilstein, A 133, 45) Yellow needles

NITRO XYLIDINE SULPHONIC ACID

$C_6H_4Me_2(NO_2)(NH_2)SO_3H$ [1 3 x 6 4] S 0818 at 8° Formed by nitrating xylidine sulphonic acid (Lumprecht a Sartig, B 18, 2189, A 230, 338) Needles —KA' 1½aq —BaA' 1½aq —PbA' aq yellow silky needles

NITRO-XYLYLENE DIAMINE $C_6H_4N_2O_2$ *ie* $C_6H_4(NO_2)(NH_2)_2$ [215°] Formed by reducing tri-nitro-*m* xylene (Bussenius a Eisenstuck, A 118, 159, Fitting, A 148, 6, Wallach, A 258, 338) Red prisms With EtI at 105° it yields crystalline $C_6H_4Me_2(NO_2)(NH_2)(NH_2Et)$ —B'HCl —B'H₂Cl₂ —B'H₂PtCl₂ 3aq —B'H₂SO₄ 2aq —B''(H₂SO₄)₂ 2aq —B''H₂SO₄

NITEYL CHLORIDE *v* Nitrogen oxychlorides, under Nitrogen, p 570

NOBLE METALS The elements Gold, Ruthenium, Rhodium, Palladium, Iridium, Osmium, and Platinum are sometimes classed

together as the *noble metals*. The application of the term *noble* to metals is a survival of the alchemical notion of a close connexion between physical and moral properties. Gold was regarded by the alchemists as the purest metal, as that which most nearly approached their ideal element. As gold did not change when heated in air, and did not dissolve in any single acid, these two properties came to be looked on as characteristic, and when these properties were found to belong also to certain other metals these other metals were placed in the same class as gold. Silver dissolves in nitric acid, but it does not change by heating in air, hence Ag was often classed with Au as a noble metal. The term *noble* is applied in this article to the seven metals named above, as a convenient form of expressing the fact that these metals have many properties in common. The seven metals are all found uncombined in nature, they are all very lustrous, heavy, generally ductile and malleable, hard (except Au), some are not oxidised by heating in air, others combine with O at high temperatures.

Au has been known from prehistoric times. Pt was discovered about 1750, and the other Pt metals from 1803 to 1845. The name platinum is said to be derived from *platina del Pinto* (*platina*=diminutive of *plata*=silver), a name by which the natural alloy of the metal was known in New Granada, from whence it was first obtained. The names palladium and rhodium were given by Wollaston who separated these metals from Pt ore, in 1803, the first suggested by the planet Pallas discovered about that time, and the second from *ῥόδον*=a rose, in allusion to the colour of solutions of the salts of the metal. Tennant discovered two other metals in Pt ore, in 1803, he called one iridium, because of its many coloured salts (*iris*=the rainbow), and the other osmium, because of the peculiar smell of its volatile oxide (*οσμή*=a smell). In 1845, Claus separated a sixth metal from Pt ore from the Ural, and called it ruthenium, in allusion to Russia.

Ru, Rh, Pd, and Ir, Os, Pt, form the second and third sections or families of Group VIII in the periodic classification of the elements, the first family of this group is composed of Fe, Ni, and Co. Au belongs to Group I, it is a member of the family Cu, Ag, Au. As Cu follows immediately after Fe, Ni, Co in the *long period* beginning with K and ending with Br, and as Ag follows Ru, Rh, Pd, in the *long period* Rb to I, so Au is placed in immediate succession to Os, Ir, Pt in the *long period* which, when it is complete, will begin with an alkali metal resembling Os and end with a halogen more or less like I. Looking at the position of Au in the periodic scheme of classification (*v* Table, p 204, vol II) one may say that the relations of this element to Os, Ir, and Pt will be found very similar to the relations of Ag to Ru, Rh, and Pd, and also to the relations of Cu to Fe, Ni, and Co.

The table on page 629 presents some of the important properties of the metals Ru, Rh, and Pd.

These three metals form *oxides* MO and MO₂. Ru and Rh also form M₂O₃. Ru forms a volatile oxide RuO₄, Rh forms RhO₄, and Pd₂O is known. The oxides MO are basic, forming salts

	RUTHENIUM	RHODIUM	PALLADIUM
Atomic weights	101.4	102.7	106.3
	The only compound of these elements which has been gasified is RuO_4 . The S.H. of each has been determined directly. Molecular weights unknown.		
Spec gravities (approx)	12.8	12.1	11.4
Atomic weight / spec gravity	8.5	8.6	9.8
Melting points (approx)	above 2000°	2000°	1500°-1600°
Spec heats	0.61	0.58	0.6
Occurrence and preparation	The metals Rh, Ru, and Pd, occur in small quantities in many Pt ores, they are usually separated in the form $2\text{NH}_4\text{Cl}$ MCl_2 , on strongly heating these double chlorides, the metals remain.		
Physical properties	White, lustrous, hard, brittle, crystalline, less ductile and malleable than Pd, fuses in O H flame.	Greyish white, very hard, malleable, not very ductile, may be fused in O H flame, and is thus obtained in crystals.	White, hard, lustrous, ductile and malleable, most fusible of the Pt metals. Crystallises in octahedral and also in hexagonal forms.
Chemical properties	Oxidised by heating powdered metal in air, also by heating with KOH or K_2CO_3 , forming K_2RuO_4 , which is sol water. Combines with Cl at red heat, dissolves very slowly in <i>aqua regia</i> .	Oxidised at red heat, when in powder, combines with Cl at red heat, unacted on by any acid, but when alloyed with Pb, Cu, &c, dissolves in <i>aqua regia</i> . Fused with KHSO_4 , forms a soluble Rh K sulphate. Absorbs H rapidly. When in very fine powder, decomposes H_2CO_3 to H and CO_2 , and $\text{C}_2\text{H}_6\text{O}$ to H and $\text{C}_2\text{H}_4\text{O}$.	Oxidises superficially at moderate temp, at higher temps the oxide is reduced. Absorbs H rapidly, probably forming a definite compound. Dissolves in hot conc HClAq or H_2SO_4 , also sol HNO_3 and <i>aqua regia</i> .

$\text{MX}(\text{X} = \text{SO}_4, \text{etc})$ Ru_2O_3 and Rh_2O_3 are also basic, the corresponding salts are M_3X . A few salts corresponding with RuO_4 have been isolated, but none derivable from Rh_2O_3 or PdO_2 has been prepared. RhO_2 has not yet been proved to be basic. RuO_4 is not known, but a few salts (*ruthenates*) have been isolated in which RuO_4 forms the negative radicle, e.g. K_2RuO_4 . An acid HRuO_4 is known, the K salt (KRuO_4) is said to be isomorphous with KClO_4 and KMnO_4 . RuO_4 is a solid melting at c 26°, it is very volatile, easily reduced to lower oxides, explodes at c 108° giving RuO_2 and O, with KOHAq it forms K_2RuO_4 . The chlorides generally correspond with the oxides MO , M_2O_3 , and MO_2 , the chlorides form double chlorides with more positive chlorides, the salts MCl , 2XCl are best regarded as salts of the acids H_2MCl_2 (e.g. K_2PdCl_4 and $(\text{NH}_4)_2\text{RuCl}_6$). The cyanides MCy_2 form double salts, an acid H_2RuCy_2 is known. The sulphides generally correspond with the oxides MO , M_2O_3 , and MO_2 , PdS forms thio-salts, Na_2PdS_2 , &c; Rh_2S_3 dissolves in alkali sulphides the sulphides M_2S_3 and MS_2 are therefore more or less acidic. The three metals form many ammoniacal compounds, which are best regarded as salts of various radicles obtained by replacing H in two or more NH_3 molecules by Ru, Rh, or Pd.

In the table on page 680 are presented some of the more important properties of the metals Os, Ir, Pt, and Au.

Au differs considerably in its chemical properties from Os, Ir, and Pt. The oxides of Os, Ir, and Pt are MO and MO_2 , M_2O_3 , represented by Os_2O_3 and Ir_2O_3 , OsO_4 . Very few salts of Os have been prepared as yet, OsO and Os_2O_3 are probably basic. OsO_2 is not known, but osmates, e.g. K_2OsO_4 , have been isolated. OsO_4 is solid, with low melting and boiling points, it is slowly soluble in water, seems to form very unstable salts with alkalis, but does not decompose carbonates. Os forms a peculiar acid $\text{H}_2\text{N}_2\text{OsO}_4$, perhaps HO OsO_4 , $\text{N N OsO}_4\text{OH}$, it also forms ammoniacal bases, in which OsO seems to replace H_2 in N_2H_4 , and OsO_2 to replace H_2 in N_2H_6 . The acid H_2OsCy_2 is known. Very few salts of Ir are known, Ir_2O_3 is basic. The chloride IrCl_3 , corresponding to IrO_2 , is known, and also such salts as IrO SO_4 , in which an acid radicle seems to replace O in IrO_2 . Some uridates have been obtained, e.g. $\text{K}_2\text{O } 2\text{IrO}_4$. The acid H_2IrCy_2 , and salts of this acid, exist. Some ammoniacal bases are known, in which Ir seems to replace H_2 of N_2H_4 . A few salts corresponding with PtO are known, but they have been but little studied. PtO forms salts such as $\text{Pt}(\text{SO}_4)_2$. PtO_2 also combines with some positive oxides to form salts $x\text{PtO}_2 \cdot y\text{RO}$, e.g. $3\text{PtO}_2 \cdot \text{Na}_2\text{O}$. PtO_2 also combines with acidic and basic oxides to form complex salts, e.g. $\text{PtO}_2 \cdot 10\text{MoO}_3$ (or 10WO_3) $\cdot 4\text{Na}_2\text{O} \cdot x\text{H}_2\text{O}$. A very large number of salts of complex ammoniacal

	OSMIUM	IRIDIUM	PLATINUM	GOLD
<i>Atomic weights</i>	190.3	192.5	194.3	197
	The only compound of these elements which has been gasified is OsO ₄ . The S.H. of each has been determined directly Molecular weights are unknown.			
<i>Spec. gravities (approx)</i>	22.5	22.4	21.4	19.5
<i>Atomic weight</i>	85	86	9	101
<i>Spec. gravity</i>				
<i>Melting points (approx)</i>	Melts in the electric arc	1900°	1800°	1200°
<i>Spec. heats</i>	0.311	0.326	0.324	0.324
<i>Occurrence and preparation</i>	The three metals Os, Ir, and Pt, occur in small quantities, associated (? alloyed) with each other, and frequently with Ru, Rh, and Pd. They are usually separated as 2NH ₄ Cl MOCl ₄ , and are obtained by strongly heating these compounds			Occurs native, generally alloyed with Ag. Prepared by removing earthy impurities, or by crushing auriferous quartz, forming an amalgam of Au and Hg, and removing Hg by heat, sometimes by smelting, or by treatment with Cl &c
<i>Physical properties</i>	White, with tinge of blue, very hard, crystalline, brittle, also a black, amorphous, powder. Os does not melt at the highest temp of the O-H flame, it is the heaviest substance known.	White, lustrous, hard, crystalline, brittle, but fairly malleable at red heat, also a hard black powder. Melts in O-H flame.	Silver white, very lustrous, fairly hard, very malleable and ductile, expands by heat less than any other metal. Melts in O-H flame.	Very lustrous, yellow metal, crystallises in octahedra, good conductor of heat and electricity, most malleable of all metals, very ductile. Also obtained as a lustreless, brown yellow, powder.
<i>Chemical properties</i>	Oxidised readily, to OsO ₄ , by heating finely powdered metal in air, also oxidised by HNO ₃ , when in fine powder. Oxidised by fusion with KOH or KNO ₃ . Combines directly with Cl.	When finely divided, oxidises slowly when heated in air, and dissolves in <i>aqua regia</i> , in compact form is insol in all acids. Oxidised by fusion with KOH and KNO ₃ . Combines directly with Cl.	Not oxidised by heating in air or O, but by fusion with KOH. Dissolves in <i>aqua regia</i> . Combines directly with Cl. Absorbs H rapidly, and gives it off again at red heat.	Not oxidised by heating in air. Dissolves in <i>aqua regia</i> . Combines directly with Cl, Br, and I. Compounds are easily decomposed, yielding Au.

bases, containing Pt, are known. Pt is rather remarkable for the number of compounds which it forms with H and acidic radicals, these compounds are acids, and corresponding with each is a series of salts; the acids in question are H₂PtCl₆, H₂PtCl₅, H₂PtBr₆, H₂PtI₆, H₂Pt(NO₃)₆Cl₂, H₂Pt(NO₃)₅Cl₃, H₂PtO(NO₃)₄, H₂PtCy₄, H₂PtCy₃Cl₃, H₂Pt(SCN)₄, H₂Pt₂S₄ (salts of H₂Pt₂S₄ exist).

Au forms three oxides, Au₂O, AuO, and Au₂O₃. Au₂O is slightly sol. cold water, a few salts corresponding with Au₂O₃ are known, e.g. Au(NO₃)₃, AuSO₄, corresponds with AuO. Au₂O₃ forms aurates, e.g. KAuO₄, by reacting with alkalis. The sulphides, Au₂S and AuS, form thio-salts with the alkali sulphides, e.g. NaAuS₂,

but these thio salts have been examined very slightly. The chlorides are AuCl and AuCl₃, the acids HAuCl₄, HAuBr₄, and HAuCy₄, and salts of the form M'AuCl₄, are known. The compounds of Au are easily decomposed with separation of Au.

The Pt metals fall into two families. (1) Ru, Rh, Pd, and (2) Os, Ir, Pt. The elements, as a group, possess the physical characters of metals very distinctly marked, but the existence of acidic oxides and sulphides points to the non-metallic nature of these elements. Au is physically more metallic, and chemically more non-metallic, than any of the other noble metals. For the relations of Au to Cu and Ag, and

the relations of this family to the members of Group VIII (Fe, Ni, Co, Ru, Rh, Pd, Os, Ir, Pt), and also to the other members of Group I (Li, Na, K, Rb, Cs), *v* COPPER GROUP OF ELEMENTS, vol. ii p 250, *v* ALSO IRON GROUP OF ELEMENTS, this vol p 65. For details about the individual noble metals, *v* GOLD, vol ii p 647, Iridium, this vol p 46, Osmium, Palladium, in this vol, PLATINUM, RHODIUM, and RUTHENIUM in vol 17.

M M P M

NOMENCLATURE The nomenclature of chemistry is based on the system introduced by Lavoisier, De Morveau, Berthollet, and De Fourcroy in 1787. The leading principles laid down by the French chemists were (1) that every substance is to be regarded as an element until it is proved to be otherwise, (2) that the name of a compound is to exhibit the elements, and as far as possible the relative proportions of the elements, of which it is composed. The names given to elements are not based on any uniform principle, some are known by the names given them for centuries, more recently discovered elements are named, sometimes from the names of compounds of them well known before the elements were discovered, sometimes from the localities where the material was found from which the element was prepared for the first time, sometimes from a characteristic property of the element, sometimes from fanciful considerations, and sometimes to express the pride of the discoverer in his own nationality. To all more recently discovered metals have been given names ending in *um*. Binary compounds are designated by names ending in *ide*, this termination being applied to the name of the more negative element, thus all binary compounds of O are called oxides. As Cl is more negative than S, it is better to call S_2Cl_2 sulphur chloride than chlorine sulphide. When two oxides, chlorides, &c., of an element exist, they are generally distinguished by throwing the name of the more positive element into adjectival form, and using the termination *ic* to indicate more of the negative element, and *ous* to denote less of the negative element, relatively to a fixed quantity of the more positive element, thus the compounds $FeCl_2$ and $FeCl_3$ are known as ferrous chloride and ferric chloride respectively. When more than two oxides, chlorides, &c., of an element are known, it is customary either to use prefixes *di-* (or *bi*), *tri-* (or *ter*), &c., or to indicate the relative proportions of the elements by such prefixes as *hypo* and *per*. It is also customary to give names to certain oxides for the purpose of indicating their acidic character. Thus the five oxides of N have been named as follows at different times

N_2O Nitrous oxide, nitrogen monoxide (might also be called hyponitrous anhydride)

NO Nitric oxide, nitrogen dioxide

N_2O_3 Nitrogen trioxide, nitrogen sesquioxide, nitrous anhydride

N_2O_4 Nitrogen dioxide, nitrogen tetroxide, nitroso-nitric anhydride, nitrogen peroxide

N_2O_5 Nitrogen pentoxide, nitric anhydride, nitrogen peroxide

The same name—nitrogen dioxide—has been given to two different compounds, NO and NO_2 ;

and the name peroxide has been used for NO_2 and N_2O_5 . This illustrates a difficulty. The prefixes *mono*, *di*-, &c. are sometimes employed to designate the first, second, third, &c. members of a series of oxides, chlorides, &c. of the same element, without implying anything as to the number of O, Cl, &c. atoms in the molecules of the various compounds, but the same prefixes are employed sometimes to imply one, two, &c. atoms of O, Cl, &c. On both systems of naming N_2O is called monoxide, on the first system NO is called dioxide, but on the second system it must be called monoxide, hence the second system of naming gives the same name to two different compounds. To get over this difficulty N_2O may be called dinitrogen monoxide, and NO mononitrogen monoxide, but such names are cumbersome. The prefix *per-* is generally employed to designate the highest compound of a series, & the one with relatively most negative element, but a higher compound may be discovered, in such a case the prefix, if used at all, must be moved from the older to the more recently discovered substance. Salts are named from the acids of which they are metallic derivatives. If there are two acids containing the same elements, to that with relatively more of the negative radicle is given a name ending in *-ic*, and its salts are called *-ates*, to the other acid is given a name ending in *-ous*, and its salts are called *-ites*. By the use of prefixes *per*, *hypo*, &c., four or five acids and their salts may be named, e.g.

$HClO$ Hypochlorous acid.

$HClO_2$ Chlorous acid

$HClO_3$ Chloric acid

$HClO_4$ Perchloric acid

Compounds which probably contain the OH group are generally called hydroxides, and those containing the SH group are called hydro sulphides or sulphhydrates. These names more or less imply a special view of the structure of the compounds, as the same view is not always held by all chemists, it seems preferable to call a commonly occurring compound, such as KOH, potash, rather than potassium hydroxide. The nomenclature of organic chemistry must be based on certain conceptions regarding the structure of carbon compounds. These compounds are so numerous, and many of them show such small differences in empirical composition, while not a few are identical in composition, that it would be impossible to frame a systematic nomenclature without the help of the conceptions of molecular structure which lie at the root of organic chemistry. In other words names cannot be found for the vast variety of carbon compounds without considering the properties and functions of these compounds as well as their composition, but the only way we have of expressing, at present, the chemical properties of carbon compounds is in terms of the molecular and atomic theory. For complicated examples *v* AZO-COMPOUNDS, vol i p 369, to some simpler cases *v* HYDROCARBONS, vol. ii p 715.

No attempt is made in this article to trace the historical development of chemical nomenclature, nor to discuss fully the present system of naming used in the science. Chemical nomenclature is a subject the details of which must be

learned gradually by studying chemical substances and their reactions. The article NOMENCLATURE in the first edition of this Dictionary contains references to the more important older memoirs on the subject. In addition to these should be added a report on chemical nomenclature by a committee of the British Association (*B A* 1884 89). Reference should also be made to the suggestions of the Council of the Chemical Society (*C J* 35, 277). M M P M

NONADECANE *v* ENNDECANE

NONANE *v* ENNANE

NONAPHTHENE $C_{14}H_{10}$ (136° cor.) SG $\frac{20}{20}$ 763 H C γ 1,380,900 H C β 1,383,400 Occurs in Russian petroleum (Markownikoff, *J R* 15, 331, Ossipoff, *J R* 20, 645). It is ψ cumene hexahydride as it yields ψ cumene sulphonic acid on treatment with H_2SO_4 (Konovaloff, *C C* 1887, 1133, *J R* 22, 4, 118). Br and AlBr₃ yield tri bromo ψ cumene. Nitric acid (SG 14) yields $C_8H_7NO_2$ (219°), which on reduction yields $C_8H_7NH_2$ (172°–177°), SG $\frac{2}{2}$ 873, smelling like conine. Chlorine yields C_8H_7Cl (186°), whence C_8H_7I (110° at 200 mm), C_8H_7OAc (209°), and C_8H_7OH (191°), SG $\frac{20}{20}$ 8972 may be successively derived. Nonaphthyl iodide is converted by Ag_2O into $(C_8H_7)_2O$ (801°), SG $\frac{20}{20}$ 866. Nonaphthylene C_8H_8 (136°), SG $\frac{2}{2}$ 807 may be obtained from nonaphthyl chloride.

Isononaphthene $C_{14}H_{18}$ (151°) H C γ 1,381,700 H C β 1,384,200 Occurs also in Russian petroleum.

NONIC ACID $C_{10}H_{18}O_2$ [187°] Formed by the action of bromine and alcoholic potash on isopropyl isovaleric acid (Wohlbruck, *B* 20, 2336). Plates.

NONOIC ACID *v* ENNOIC ACID

NONYL The radicle C_9H_{19} , called ENNYL in this Dictionary.

Di-nonyl is OCTODECANE

NONYL ALCOHOL *v* ENNYL ALCOHOL

NONYLENE *v* ENNYLENE

NONYLENIC ACID *v* ENNENOIC ACID

NONYLIC ACID *v* ENNOIC ACID

NORMETHYLEMIPIC ACID *v* Methyl derivative of DI OXY-PHTHALIC ACID

NORMETHYLNITROHEMIPIC ACID *v* Methyl derivative of NITRO DI OXY PHTHALIC ACID

NORNARCOTINE *v* NARCOTINE

NOROPIANIC ACID *v* OPIANIC ACID

NORWEGIUM Ng (?) At w c 219 This name was given by Dahll to a substance separated by him from *nickel glance* from the Norwegian island of Osterö and ranked by him among the elements (*B* 12, 1731, 13, 250). According to Prochazka (*A C J* 2, 213) the element exists in some specimens of unrefined Pb (along with Bi, Cu, and Ni). The claim of Ng to rank as an element cannot be regarded as yet satisfactorily established. Ng is said to show great resemblances to Bi, to melt at c 254°, to form a fusible oxide resembling Bi_2O_3 . The hydroxide is soluble in KOHAq, also in a large excess of NH_3 or Na carbonate solution. If the oxide is Ng_2O_3 , the at w is approximately 219, if the oxide is NgO , the at w is approximately 146. M M P M

NOTATION The expression of the composition, and, as far as possible, the properties, of compounds by the use of symbols and formulæ. The subject is discussed sufficiently in the articles EQUATIONS, CHEMICAL (*q v* vol II p 433), FORMULÆ (*q v* vol II p 572), and ISOMERISM (*q v* this vol p 79). M M P M

NUCIN *v* JUGLONE

NUCITANNIN Occurs in walnuts (Phipson, *C N* 20, 116). Decomposed by dilute acid into sugar and red amorphous rufic acid $C_{14}H_{12}O_4$, which yields the salts $CaC_8H_4O_4$, and $PbC_8H_4O_4$.

NUCLEIN *v* PROTEIDS, Appendix C

NUCLEO ALBUMIN *v* PROTEIDS, Appendix C

NUCLEO PROTEIDS *v* PROTIDS, Appendix C

NUMBERS, LAW OF EVEN Laurent (*A Ch* [3] 18, 266) said that the sum of the mono valent, trivalent, and pentavalent elements contained in any well defined and stable compound is always an even number. This *law of even numbers* was long an article of belief among orthodox chemists. If by an *n* valent element is meant one the atom of which combines directly with *n* other atoms to form a molecule, then there are several exceptions to the so called law, e.g. themolecules $InCl_2$, $InCl_3$, and probably $InCl$ exist as gases, so do the molecules $FeCl_2$ and $FeCl_3$, the molecules WCl_2 and WCl_6 , &c. M M P M

NUPHARIN $C_{14}H_{12}N_2O_2$. An amorphous substance in the rhizome of *Nuphar luteum* (Gruning, *J* 1882, 1156).

NUX VOMICA *v* STRYCHNINE

O

n-OCTADECANE $C_{18}H_{38}$ [28°] (317°). SG $\frac{2}{2}$ 775, $\frac{2}{2}$ 768 Occurs in paraffin from brown coal. Formed by reduction of stearic acid with HI and P and by the action of Na on ennyl iodide (Krafft, *B* 15, 1703, 16, 1723, 19, 2221, 21, 2261). Hexagonal tables.

OCTADECINENE $C_{18}H_{34}$ [30°] (164° at 15 mm) SG $\frac{2}{2}$ 8016 Formed by heating $C_{18}H_{34}Br_2$ with alcoholic potash (Krafft, *B* 17, 1874). Plates.

OCTADECIC ACID $(C_{17}H_{33})_2CH CO_2H$ [39°] (above 300°) Prepared by heating di octylmalonic acid (Conrad & Bischoff, *B* 13, 597). White crystals.

OCTADECYL ALCOHOL $C_{18}H_{37}OH$ [59°] (210° at 15 mm) SG $\frac{2}{2}$ 8048, $\frac{2}{2}$ 7849 Occurs in crude cetyl alcohol, and is prepared by reducing stearic aldehyde with zinc dust and acetic acid (Krafft, *B* 16, 1722, 17, 1627).

Acetyl derivative [α 31°] (223° at 15 mm)

OCTADECYL-BENZENE $C_{18}H_{38}$, C_8H_8 [86°] (249° at 15 mm) Formed from octadecyl iodide, iodobenzene, and sodium (Krafft, *B* 19, 2984) Yields a solid sulphonic acid

OCTADECYLENE $C_{18}H_{34}$ [18°] (179° at 15 mm) $SG \frac{1}{2}$ 7910, $\frac{2}{3}$ 7881 Formed by distilling octadecyl palmitate (Krafft, *B* 16, 8024)

Octadecylene $C_{18}H_{34}$ *Anthemene* [64°] $SG \frac{1}{2}$ 942 VD 127 ($H=1$) Obtained from *Anthemus nobilis* by extracting the blossoms with ligroin (Naudin, *Bl* [2] 41, 488) Minute needles

OCTADECYLENE BROMIDE *v* **DI-BROMO-OCTADECANE**

OCTADECYL IODIDE $C_{18}H_{37}I$ [33 5°] (K), [43°] (S) From the alcohol, I , and P (Krafft, *B* 19, 2984, Schweizer, *Ar Ph* [3] 22, 753)

OCTADECYL-PHENOL $C_{18}H_{37}O$, C_8H_8 , OH [84°] (277° at 15 mm) Formed by fusing octadecylbenzene sulphonic acid with potash (Krafft, *B* 19, 2985) Plates (from alcohol)

n **OCTANE** C_8H_{18} , Mol w 114 (125°) $SG \frac{1}{2}$ 7188 VD 4 03 (Lemoine, *Bl* [2] 41, 163) CE (0°-10°) 001186, (0°-100°) 001331 (Thorpe, *C J* 37, 217) SV 186 3 Occurs in American petroleum (Pelouze & Cahours, *A* 127, 197, Schorlemmer, *C J* 15, 419) Formed by the action of zinc and HCl_{aq} on sec octyl iodide (Schorlemmer, *C J* 27, 1029), by the action of sodium amalgam on *n* octyl iodide (Zincke, *A* 152, 15), and by the action of sodium on *n* butyl iodide (Schorlemmer, *A* 161, 280) Obtained also by distilling whale oil under pressure (Engler, *B* 22, 595) Oil

Iso-octane $Pr CH_3 CH_2$, Pr *Di isobutyl* (108°) $SG \frac{1}{2}$ 709, $\frac{2}{3}$ 693 (W), $\frac{1}{2}$ 711 (Thorpe) CE (0°-10°) 001205, (0°-100°) 001401 $\mu_p = 1.3943$ at 16° (W) $R_p = 64.47$ SV 184 5 (Schiff, *A* 220, 88) VD 3 94 Formed by electrolysis of potassium isovalerate (Holbe, *A* 69, 261, *C S Mem*, 3, 378, *C J* 2, 157) Formed also by the action of sodium on isobutyl iodide and on a mixture of isoamyl and isopropyl iodides (Wurtz, *A* 93, 112, 96 364, Schorlemmer, *Pr* 16, 37, *A* 144, 188, W C Williams, *C J* 31, 541, 35, 125) Occurs also among the products of the distillation of whale oil under pressure (Engler, *B* 22, 595) Oil

Octane C_8Me_8 , *Hexa methyl ethane* [97°] (106°) Formed from tert butyl iodide and sodium (Lwow, *Bl* [2] 35, 169)

Octane C_8H_{18} (119°) VD 3 97 (obs) $SG \frac{1}{2}$ 712 Formed by reduction of conine, comceine, or conhydrine by prolonged heating with HI and P (Hofmann, *B* 18, 12)

OCTANE DICARBOXYLIC ACID $C_{18}H_{34}O_4$, [184°-194°] A product of the action of sodium on bromo methyl ethyl acetic ether (Pagenstecher, *A* 195, 121) Crystals (from water)

Octane dicarboxylic acid
 $CO_2H CH_2 CHMe (CH_2)_6 CO_2H$ [44°] *Methyl-asetic acid* Formed by heating the tetracarboxylic acid (Perkin, jun, *C J* 51, 218) Crystals— AgA'' , *Ethyl ether Et_2A''*. Oil

Octane tetracarboxylic ether
 $(CO_2Et)_4 CH CHMe (CH_2)_6 CH (CO_2Et)_2$ (275° at 60 mm.). A product of the action of di bromo-

methyl pentamethylene on malonic ether (Perkin, *C J* 53, 217) Syrup

Octane tetradeca-carboxylic ether
 $CO_2Et CH_2 [C(CO_2Et)_2]_6 CH_2 CO_2Et$ Formed from chloro butane heptacarboxylic ether and sodium butane heptacarboxylic ether (Bischoff, *B* 21, 2116) Viscid oil

OCTENE *v* **OCTYLENE**

OCTENOIC ACID $C_8H_{14}O_2$, *is*
 $Pr CH_2 CH CH CH_2 CO_2H$ (231°) Formed by distilling isobutyl paraconic acid (Fittig, *B* 21, 920, *A* 255, 103) Oil

Octenoic acids are also formed by oxidation of octenoic aldehyde (Fossek, *M* 2, 622), and by reduction of suberene carboxylic acid $C_8H_{14}O_2$ (Spiegel, *A* 211, 119) They are volatile with steam

OCTENOIC ALDEHYDE $C_8H_{14}O$ (150° at 18 mm) Formed by heating isobutyric aldehyde with conc $NaOAc_{aq}$ at 150° (Fossek, *M* 2, 614) Liquid, volatile with steam Forms a mirror with ammoniacal $AgNO_3$, and a crystal line compound with $NaHSO_4$ Yields acetic and isobutyric acids on oxidation

Octenoic aldehyde $PrCH CH_2 CHO$ (173°). Formed from butyric aldehyde and aqueous $NaOH$ or $NaOAc$ (Raupenstrauch, *M* 8, 108) Oil Reacts with phenyl hydrazine Yields $C_8H_{14}O$ (161°) on reduction with iron and $HOAc$

Octenoic aldehyde $C_8H_{14}O$ (230° \pm) $SG \frac{1}{2}$ 958 Formed by passing dry HCl into iso butyric aldehyde (Oeconomides, *Bl* [2] 36, 209) Oil, resimified by potash Reduces ammoniacal $AgNO_3$, forming a mirror

OCTENYL ALCOHOL $C_8H_{16}O$, *is*
 $CH CH CH CH_2 CH_2 OH$ *Di ethyl allyl carbinol* (156°) $SG \frac{1}{2}$ 889 CE (0°-33°) 00104. Formed from di ethyl ketone, allyl iodide, and zinc (Schirokoff & Saytzeff, *A* 196, 113) Oil Yields di ethyl ketone and propionic acid on oxidation With HCl it forms a compound converted by caustic potash into tri-oxy octane $KMnO_4$ yields $CH_3(OH) CH_2 CO_2H$ Dilute H_2SO_4 at 100° yields octenene (α 123°) (Reformatsky, *J pr* [2] 30, 217)

Octenyl alcohol $CMePr(C_2H_5)(OH)$ (160°). $SG \frac{1}{2}$ 8486, $\frac{2}{3}$ 8345 Formed from methyl propyl ketone, allyl iodide, and zinc (Semljanitzin, *J pr* [2] 23, 263, Reformatsky, *J pr* [2] 40, 412) Yields $CMePr(OH) CH_2 CO_2H$ on oxidation

OCTENYL CHLORIDE $C_8H_{15}CH CHCl$ (168°) $SG \frac{1}{2}$ 927 Formed from $C_8H_{15}Cl$, and alcoholic potash (Béhal, *A Ch* [6] 15, 278) Liquid, smelling like carrots

OCTINENE C_8H_{14} , *is* $CH_3 C \equiv C C_2H_5$, *Methyl-amylicetylene* (183°) $SG \frac{1}{2}$ 71 Formed by the action of alcoholic potash on octylene bromide (derived from octylene got by dehydrating octyl alcohol) (Rubien, *A* 142, 299, Béhal, *Bl* [2] 47, 33, 48, 704, 50, 359, 629, *A Ch* [6] 15, 274, 428) Mobile liquid Does not react with ammoniacal $CoCl_2$. On dissolving in cold H_2SO_4 , and pouring into ice-cold water it yields the ketone $C_8H_{14}O$ (171°), $SG \frac{1}{2}$ 885 which forms hexoic and acetic acids on oxidation.

Octinene C_8H_{14} , *is* $CH C \equiv C CH_2 C_2H_5$ (125°-133°) Obtained by heating the preceding isomeride with sodium at 110° (Béhal). Liquid Forms a yellow pp with ammoniacal cuprous chloride

Octinene C_8H_{14} . *Conylene* (125°) V D 55 6 (H=1). Obtained by dry distillation of dimethyl conline methylo-hydroxide and by heating 'azoonhydrine' with P_2O_5 (Wertheim, A 123, 170, Hofmann, B 14, 710). Oil. Forms a di-bromide (*v* Di-bromo-octylene).

Octinene C_8H_{14} . *Dusocrotyl* [5°] (125°-180°). Formed from OMe , $CHBr$ and sodium (Przybytek, J. R. 20, 506). Oil, rapidly absorbing oxygen.

Octinene OH , OMe CH_2 , CH_2 , OMe CH_2 . (114°). Formed from CH_2 , OMe CH_2 , Cl and sodium (Przybytek, B 20, 3240).

Octinene C_8H_{14} . (c 123°) S G § 7734, 154 7588 R₂ 62 12. Formed by heating octenyl alcohol CH_2 , CH CH_2 , CH_2 , OH with dilute H_2SO_4 at 100° (Beformatsky, J pr [2] 30, 217). Mobile liquid, absorbing oxygen from the air. Combines with bromine. Oxidised to acetic and propionic acids by chromic acid mixture. Yields C_8H_7Br .

Isomeride: Xylene tetrahydride

OCTINOIC ACID *v* Di ALLYL ACETIC ACID

OCTINYL ALCOHOL $C_8H_{17}O$

$(OH, CH, CH_2)_2OMe(OH)$ *Methyl-di-allyl-carbinol* (158° cor) S G § 864, 13 852 H C 1,201,400 (Longumme, A Ch [5] 23, 888). Formed from allyl iodide, acetic ether, and zinc (Saitzeff a Sorokin, B 9, 33, 277, A 185, 169). Yields oxy-methyl glutano acid on oxidation. *Acetyl derivative* (177° cor).

OCTODECANE *v* OCTADECANE

n **OCTOIC ACID** $C_8H_{16}O_2$ *v* $CH_3(CH_2)_6CO_2H$ *Caprylic acid* Mol w 144 [17°] (237° i V) S G § 927 (Zander, A 124, 71) 15 913, 25 908 M M 8 565 at 18.5° S 25 at 100° H C 1,138,694 (Longumme, A Ch [6] 11, 221), 1,145,600 (Stohmann, J pr [2] 43, 18). C.E. (0°-10°) 00092 S V 1978. Occurs as glyceryl ether in butter (Lerch, A 49, 214) in cocoa nut oil (Fehling, A 53, 399, Renesse, A 171, 380), and in Lumburg cheese (Ilyenko, A 55, 85). It occurs also in fusel oil from various sources. Formed by oxidation of *n* octyl alcohol (Zincke, A 152, 9) and by saponifying its nitrile which is formed by the action of bromine and NaOH on the amide of ennoic acid (Hofmann, B 17, 1408). Formed also by oxidising di-oxy stearic acid with alkaline $KMnO_4$ (Spiridonoff, J pr [2] 40, 248). White crystals, insol cold water.

Salts — BaA' . S 62 at 20° — CaA' , aq — ZnA' . [186°] — PbA' . [84°] — CuA' . [266°] — AgA' white curdy pp.

Methyl ether MeA' (193°) S G § 8942 S V 220 1 C.E. (0°-10°) 00094 (Gartenmeister, A 233, 286).

Ethyl ether EtA' Mol. w 172 (206°) S G § 8842 S V 245 9. C.E. (0°-10°) 00098.

Propyl ether PrA' (225°) S G § 8805. S V 270 3 C.E. (0°-10°) 00092.

Butyl ether C_4H_9A' (240.5°). S G § 8797 S V 295 9 C.E. (0°-10°) 00094.

Heptyl ether $C_7H_{15}A'$ (290°) S G § 8754 S V 377 0 C.E. (0°-10°) 00086.

n-Octyl ether $C_8H_{17}A'$ (306°) S G § 8755 S V 404 8. C.E. (0°-10°) 00084.

Phenyl ether PhA' . (800°)

Amide $C_8H_{15}ONH_2$. [106°]. S. 454 at 100°. Plates.

Anhydride $(C_8H_{15}O)_2O$. (c 285°). (Chiozza, A. 85, 229).

Nitrile $C_8H_{13}CN$. (195°) (F); (199°) (Hofmann, B 17, 1410). S G § 82 (Folletar, Z [2] 4, 665).

Iso-octic acid $C_8H_{16}O_2$. (219°) S G § 926; § 911 S 15 at 15°. Formed by oxidising iso-octyl alcohol (W C Williams, O J 31, 542, 35, 129). Liquid — NaA' — KA' — MgA' , 2aq — AgA' . Crystallises from hot water.

Ethyl ether EtA' (175°).

Iso-octyl ether $C_8H_{17}A'$ (c 280°)

Octoic acid $CH_3PrCH_2CHMeCO_2H$ (c 215°)

Formed by oxidation of isodibutyl $C_8H_{16}O$ (Butlerow, A 189, 70). Liquid. Probably identical with the preceding acid.

Octoic acid OMe , CMe , CO_2H ? (210°-230°)

Formed, in small quantity, by passing CO over a mixture of NaOMe and NaOAc at 200° (Geuther a. Fröhlich, A 202, 813).

Octoic acid Pr_2CHCO_2H

Di-n-propyl acetic acid (220°) S G § 9215

Obtained by boiling di propyl acetoacetic ether with alcoholic potash or by heating di propyl-malonic acid (Burton, Am 3, 389, Furth, M 9, 317) — CaA' , 2aq S 97 at 0°, 1 65 at 80° — BaA' — AgA' S 128.

Ethyl ether EtA' (183°)

Octoic acid $CH_3PrCMe_2CO_2H$ [18°] (215°)

Obtained from its ether, which is formed by the action of sodium on isobutyric ether (Brugemann, A 246, 149). Large hexagonal plates.

OCTOIC ALDEHYDE $C_8H_{16}O$ (171°)

Occurs among the products of the distillation of castor oil soap (Limpricht, A 93, 242, Bouis, A Ch [3] 48, 99, Stadelcr, J pr 72, 211, Dachauer, A 106, 270, Béhal, Bl [2] 47, 83, 163). Formed also by distilling a mixture of calcium octoate and calcium formate. Liquid. Combines with $NaHSO_4$. Yields a mirror with ammoniacal $AgNO_3$.

Oxim (222°)

Octoic aldehyde $C_8H_{15}CH=CHO$ (161°)

Formed by reduction of octenoic aldehyde (Rau penstrauch, M 8, 108). Oil. Volatile with steam. Reduces ammoniacal $AgNO_3$.

OCTO-ICOSOIC ALDEHYDE $C_{18}H_{36}O$

$C_8H_{17}CH(C_8H_{17})CH(C_8H_{17})CH(C_8H_{17})CHO$ (c 385°). A product of the action of alcoholic potash or of Ac_2O on heptoic aldehyde (ananthol) (Perkin, C J 43, 66). Oil. Yields a mixture of heptoic and hexoic acids on fusion with potash.

OCTONAPHTHENE $C_{18}H_{14}$ (119°)

Occurs in Russian petroleum (Markownikoff, B 20, 1851). Yields a very little tri nitro-*m*-xylene on treatment with H_2SO_4 and HNO_3 . H_2SO_4 yields a sulphonic acid, the amide of which crystallises in needles [220°].

OCTYL The radicle C_8H_{17} , which is also called CAPRYL.

Di octyl v HEXADECANE

OCTYL ACETATE *v* *Acetyl derivative* of

OCTYL ALCOHOL

DI-OCTYL-ACETONE *v* **METHYL HEPTADECYL**

KETONE

n-**OCTYL ALCOHOL** $C_8H_{17}O$. Mol. w 130 (195.5° i V). S G § 8375 (Z); § 8301, § 8249 (P). C.E. (0°-10°) 00080 M M 8 880 at 20° (Perkin) S V 190 6 (Zander, A 224, 84); 197 8 (Ramsay). Obtained from its acetyl derivative which occurs in the volatile oil of cow

paranep (*Heracleum Sphondylium*), and in the oil of *H. giganteum* (Zincke, *A* 152, 1, *B*, 4, 822, Möslinger, *A*, 185, 26) The butynyl derivative occurs in the ripe fruits of *Pastinaca sativa* (Renesse, *A* 166, 80). Heated with ammoniacal ZnCl_2 at 280° it yields a mixture of mono- di- and tri- octyl-amines, the yield of mixed bases amounting to 70 per cent of the alcohol used (Merz & Gasiorowski, *B* 17, 629)

Acetyl derivative $\text{C}_8\text{H}_{17}\text{OAc}$ (212° cor) SG § 8847 (G); § 8744, § 8678 (P) CE (0° - 10°) 00094. SV 2458 (Gartenmeister) MM 10 601 st 16.1° (Perkin, *O* J 45, 421)

Benzoyl derivative $\text{C}_8\text{H}_{17}\text{OBz}$ (306°)

Ethyl ether $\text{C}_8\text{H}_{17}\text{OEt}$ (183°) SG 11 79.

Sec Octyl alcohol $\text{C}_8\text{H}_{17}\text{CH(OH)CH}_3$ **Capryl alcohol** **Methyl-hexyl-carbinol** (179.5° cor) SG § 8236, § 8178 MM 9 004 at 12.4° (Perkin) SV 1913 (Sohlf, *A* 220, 103) $\mu_s = 1.4297$ $R_{20} = 65.57$ (Bruhl, *A* 203, 28)

Formed by distilling sodium nicoelate with NaOH (Bous, *A* 97, 34, Moschinn, *A* 87, 111, *A* *Ch* [3] 44, 140, Lumprecht, *A* 93, 242, Neison, *O* J 27, 837, Schorlemmer, *Pr* 16, 376, *O* J 27, 1029) Obtained also from *n*-octane via octyl chloride and octyl acetate (Schorlemmer, *A* 152, 152) Yields methyl hexyl ketone on oxidation

Acetyl derivative $\text{C}_8\text{H}_{17}\text{OAc}$ (193°)

Octyl alcohol $\text{C}_8\text{H}_{17}\text{O}$ **Di-isobutyl hydrate** (180°) SG § 841 Formed by chlorinating $\text{CH}_3\text{PrCH}_2\text{Pr}$, converting the resulting octyl chloride into octyl acetate, and boiling this with KOHAq (W C Williams, *O* J 35, 127) Yields an octoic acid on oxidation

Octyl alcohol $\text{C}_8\text{H}_{17}\text{O}$ (c 162°) SG § 820 Formed at the same time as the preceding (W) Yields on oxidation a ketone $\text{C}_8\text{H}_{15}\text{CO}$ C_8H_{15} (160°)

Octyl alcohol $\text{CHEt, CH(OH)C}_8\text{H}_{15}$ (164° - 168°) Formed from CH_2BrCOBr and ZnEt_2 followed by water (Winogradoff, *A* 191, 125) Sl sol water

Octyl alcohol $\text{C}_8\text{H}_{17}\text{O}$ (174° - 178°) SG § 811 Obtained from octylene, by treatment with HI, the resulting octyl iodide being converted into octylacetate by AgOAc (De Clermont, *C* R 66, 1211, *A* 149, 38, *Bl* [2] 12, 212) Yields a ketone $\text{C}_8\text{H}_{15}\text{O}$ on oxidation, and, on further oxidation, acetic and hexoic acids. Is probably identical with methyl-hexyl-carbinol

Acetyl derivative (c 178°)

Tert Octyl alcohol $\text{Pr}_3\text{CMe(OH)}$ (161.5° cor) SG § 8236, § 8151 Formed from di propyl ketone, MeI, and zinc, followed by water (Saytzeff, *J* pr [2] 81, 320, *Bl* [2] 46, 267) Yields acetic, propionic, and butyric acids on oxidation

Acetyl derivative (c 175°) SG § 8554 (160.5°) SG 12 838

Tert Octyl alcohol OEt, Pr(OH) Formed from butyryl chloride or EtOPr and ZnEt_2 , followed by water (Butlerow, *Bl* [2] 5, 17, Sokoloff, *J* R 1887, 595) Yields butyric, propionic, and acetic acids on oxidation

Acetyl derivative (177°)

Octyl alcohol $\text{CMe, CH}_2\text{CMe}_2\text{(OH)}$ **Isodibutol**, [c - 20°] (147°) SG § 842 Formed from 'di isobutylene' hydriodide and Ag_2O (Butlerow, *A* 189, 53) Yields $\text{CMe, CO}_2\text{H}$ and acetone on oxidation

Sec-Octyl alcohol (182° - 186°). Obtained from *n*-octane by chlorination and conversion of the mixed octyl chlorides into acetates (Schorlemmer) Yields propionic and valeric acids on oxidation

Acetyl derivative (200°)

OCTYL ALLOPHANATE $\text{C}_{10}\text{H}_{21}\text{N}_2\text{O}_2$ $\text{C}_8\text{H}_{17}\text{O CO NH CO NH}_2$ [166°] Formed from octyl alcohol and Cl CO NH_2 (Gattermann, *A* 244, 40) Silky needles, v. sol hot alcohol

n-OCTYLAMINE $\text{C}_8\text{H}_{17}\text{NH}_2$ (186°) (E), (180°) (H & D) Formed, together with di- and tri octyl amine, by heating octyl iodide with alcoholic NH_3 at 100° , or octyl alcohol with ammoniacal ZnCl_2 at 280° (Renesse, *A* 166, 85, Merz & Gasiorowski, *B* 17, 629) Formed also by reducing nitro octane (Eichler, *B* 12, 1885), and produced by decomposition of the urea $\text{C}_8\text{H}_{17}\text{NH CO NH CO NH}_2$ [100°], which is produced by the action of KBr on ennoic amide $\text{C}_8\text{H}_{17}\text{CO NH}_2$ (Hofmann, *B* 15, 773, Hoogewerf a. van Dorp, *R* T C 6, 387) Oil— B' HCl — $\text{B}'_2\text{H}_2\text{PtCl}_6$ —Picrate [113°] Plates (H & D)

Sec-Octylamine $\text{C}_8\text{H}_{17}\text{CH(NH}_2\text{)CH}_3$ **Caprylamine** (175°) (B), (163°) (Jahn, *M* 3, 172). Formed, together with the di- and tri-octylamines, by heating iso octyl chloride with aqueous NH_3 (Malbot, *C* R 105, 575, *A* *Ch* [6] 13, 507, cf Squire, *C* J 7, 108, Cahours, *A* 92, 399, *C* R 39, 254, Bous, *A* *Ch* [3] 44, 139) Formed also by heating the alcohol with ammoniacal ZnCl_2 at 260° (Merz & Gasiorowski, *B* 17, 634)— B' HCl — B' HI — B' HAuCl_4 — $\text{B}'_2\text{H}_2\text{PtCl}_6$ — B' HNO_3 — $\text{B}'_2\text{H}_2\text{SO}_4$: crystalline, v sol water

Di n-octyl-amine $(\text{C}_8\text{H}_{17})_2\text{NH}$ [37°] (298°) VD 8 27 (obs)— B' HCl — $\text{B}'_2\text{H}_2\text{PtCl}_6$ nearly insoluble yellow plates (M & G)

Di-iso-octyl-amine (260° - 270°) VD 8 49 (obs)— B' HCl — B' HAuCl_4 — $\text{B}'_2\text{H}_2\text{PtCl}_6$

Tri n-octylamine $(\text{C}_8\text{H}_{17})_3\text{N}$ (367°) Solid, sl. sol 90 p c alcohol— $\text{B}'_2\text{H}_2\text{PtCl}_6$

Tri-iso-octyl-amine $(\text{C}_8\text{H}_{17})_3\text{N}$ (c 370°) Oil— $\text{B}'_2\text{H}_2\text{PtCl}_6$ reddish brown mass

n-OCTYL-BENZENE $\text{C}_8\text{H}_{17}\text{C}_6\text{H}_5$ [-7°] (263°) SG § 849 Formed from bromobenzene, *n*-octyl bromide, and sodium (Schweinitz, *B* 19, 641, Ahrens, *B* 19, 2718)

Octyl-benzene $\text{Pr CH}_2\text{(CH}_2\text{)}_6\text{C}_6\text{H}_5$ (245° - 255°) Formed by heating phenacyl isomylmalonic acid with zinc-dust (Paal & Th Hoffmann, *B*, 23, 1502) Oil, with blue fluorescence

References—BROMO-, CHLORO-, and IODO-OCTYL-BENZENE and OCTYL-PHENYL AMINE

n-OCTYL-BENZENE SULPHONIC ACID $\text{C}_8\text{H}_{17}\text{C}_6\text{H}_4\text{SO}_3\text{H}$. Formed by sulphonation of *n*-octyl-benzene (Schweinitz, *B* 19, 642)— BaA' , aq— PbA' , aq— AgA' minute soluble prisms

n-OCTYL-BENZOIC ACID $\text{C}_8\text{H}_{17}\text{C}_6\text{H}_4\text{CO}_2\text{H}$ $\text{C}_8\text{H}_{17}\text{C}_6\text{H}_4\text{CO}_2\text{H}$ [14°] (189°) Formed by saponifying its nitrile which is obtained by distilling the formyl derivative of *p* octyl-phenylamine with zinc-dust (Beran, *B* 18, 133). Plates or needles— AgA'

Nitrils $\text{C}_8\text{H}_{17}\text{C}_6\text{H}_4\text{CN}$ (c 313° uncor.).

OCTYL BROMIDES Formed from the corresponding alcohols, Br, and P (Zincke, *A* 152, 5, Lachovitch, *A* 220, 181).

$\text{CH}_3(\text{CH}_2)_7\text{CH}_2\text{Br}$ (199°) (Z), (204° cor) (Perkin) S G $\frac{1}{2}$ 11180, $\frac{2}{3}$ 11099
 $\text{C}_8\text{H}_{17}\text{CH}_2\text{Br}$ (188°) S G $\frac{2}{3}$ 1099

OCTYL CARBAMATE $\text{NH}_2\text{CO}_2\text{C}_8\text{H}_{17}$, [55°]. (281°) Formed from *sec* octyl (capryl) alcohol and ONCl at 100° (Arth, *Bl* [2] 45, 703, *A Ch* [6] 8, 480) Crystals, v sol alcohol

***n* OCTYL CHLORIDE** $\text{CH}_3(\text{CH}_2)_7\text{CH}_2\text{Cl}$ Mol w 148.5 (180°) (Zincke, *A* 152, 4), (183° cor) (Perkin) S G $\frac{1}{2}$ 8786, $\frac{2}{3}$ 8719 M M 10 128 at 18° Formed from *n* octyl alcohol

Sec Octyl chloride $\text{C}_8\text{H}_{17}\text{CHClCH}_3$ (172° cor) S G $\frac{1}{2}$ 8708, $\frac{2}{3}$ 8639 M M 10 248 at 18° (Perkin) Formed from *sec* octyl alcohol and HCl (Bouis, *A* 92, 398, Malbot, *Bl* [3] 3, 68) Obtained also, together with the preceding isomeride, by chlorinating *n* octane (Schorlemmer, *A* 152, 152)

Octyl chloride $\text{PrCH}_2\text{CH}_2\text{CHMeCH}_2\text{Cl}$ A mixture of this chloride with $\text{PrCH}_2\text{CHClPr}$ is formed by chlorinating $\text{PrCH}_2\text{CH}_2\text{Pr}$ (Williams, *C J* 35, 127, cf. Schorlemmer, *A* 144, 190)

Octyl chloride $\text{CMe}_2\text{CH}_2\text{CClMe}$, (145°-150°) S G $\frac{2}{3}$ 890 Formed from 'disobutylene' and HCl at 100° (Butlerow, *A* 189, 51)

Octyl chloride CETPrCl (155°) (Butlerow, *Bl* 5, 24)

***n*-OCTYLENE** C_8H_{16} Octene (123°) S G $\frac{1}{2}$ 722 Formed from *n* octyl alcohol, I, and P (Möslinger, *A* 185, 52)

Octylene C_8H_{16} (123°) at 750 mm S G $\frac{2}{3}$ 7294 (S), $\frac{2}{3}$ 7197 (Bruhl, *A* 235, 11) C E $\frac{1}{2}$ (9-123.4) 00138 μ_n 1.418 S V 177.2 (Schiff, *A* 220, 90) Formed by heating *sec*-octyl alcohol with H_2SO_4 or fused ZnCl_2 (Bouis, *A* 92, 396) Formed also, together with octyl bromide, by the action of P and Bi on *sec* octyl alcohol (Lachovitch, *A* 220, 185) It is also a product of the action of NH_4Aq on *sec* octyl iodide at 150° (Malbot, *A Ch* [6] 13, 514) Oil, with unpleasant odour Not affected by cold alcoholic KOH Is perhaps identical with the preceding octylene

Octylene $\text{CMe}_2\text{CHCMe}_2$ Disobutylene (108° i v) (B), (112°) (Malbot, *A Ch* [6] 19, 370, *C R* 108, 957) S G $\frac{2}{3}$ 734 H C 1,252,500 H F 51,500 Formed by polymerisation of isobutylene by heating with H_2SO_4 (1 pt) and water (1 pt) at 100° (Butlerow, *B* 8, 1683, 9, 1687, *A* 180, 245, 189, 44, *J R* 1882, 190, Konovaloff, *Bl* [2] 34, 334) Yields acetone, $\text{CMe}_2\text{CO}_2\text{H}$, and oxy octoic acid on oxidation

Octylene CMePrCHET (120.4° cor) S G $\frac{2}{3}$ 7814 Formed in the action of MeI and Zn on di propyl ketone (Sokoloff, *J pr* [2] 39, 444)

Octylene CETPrCHMe or CETCHET (119°) S G $\frac{2}{3}$ 7865 A product of the action of EtI and zinc on ethyl propyl ketone (Sokoloff, *J pr* [2] 39, 440) Oil Yields acetic, propionic, and butyric acids on oxidation

Octylene PrCHCHPr (116°-120°) Formed from PrCH(OH)CH(OH)Pr and HI at 140°, followed by alcoholic potash (Fossek, *M* 4, 673)

Octylenes of undetermined composition have been prepared by Schorlemmer (*A* 125, 118), Cahours (*J* 1850, 402, 1863, 529), Renard (*Bl* [2] 59, 541), Wurtz (*A* 128, 230), Cloëz (*B* 7, 823), Williams (*B* 10, 908), Thorpe & Young (*A* 165, 14), and Fittig (*A* 117, 77)

OCTYLENE BROMIDE v DI-BROMO OCTANE
OCTYLENE GLYCOL v DI-OXY-OCTANE

OCTYLENE OXIDE $\text{C}_8\text{H}_{16}\text{O}$ (145°) S G $\frac{1}{2}$ 831 Formed by the action of KOH at 180° on the chloro octyl alcohol formed by union of octylene with HOCl (Clermont, *C R* 68, 1823)

***n*-OCTYL IODIDE** $\text{C}_8\text{H}_{17}\text{I}$ (220°) (Möslinger, *B* 9, 998), (225.5°) (Dobner, *A* 243, 29) S G $\frac{2}{3}$ 13583 (D), $\frac{1}{2}$ 13407, $\frac{2}{3}$ 13316 (Perkin). C E (0°-10°) 00089 (D) M M 16 197 at 20.7° S V 222.6 Formed from the alcohol and HI

Sec-Octyl iodide $\text{C}_8\text{H}_{17}\text{CHMeI}$ (211°) S G $\frac{1}{2}$ 131 (B), $\frac{2}{3}$ 1355 (Kraft, *B* 19, 2222) Formed from the alcohol (Bouis, *A Ch* [3] 44, 181, Squire, *C J* 7, 108)

Octyl iodide $\text{C}_8\text{H}_{17}\text{I}$ (120° in vacuo) S G $\frac{1}{2}$ 1314 Formed from octylene and HI (De Clermont, *Bl* [2] 12, 212)

DI-OCTYL-MALONIC ACID $\text{C}_{16}\text{H}_{32}\text{O}_4$ (c) (C_8H_{17}) $_2\text{C}(\text{CO}_2\text{H})_2$ [75°] Crystals, insol water (Conrad & Bischoff, *B* 18, 597, *A* 204, 163)

Ethyl ether EtA (338°) S G $\frac{2}{3}$ 896

OCTYL NITRITE $\text{C}_8\text{H}_{17}\text{ONO}$ (176°) S G $\frac{1}{2}$ 862 Formed from octyl alcohol and HNO_3 (Eichler, *B* 12, 1887)

Sec Octyl nitrite $\text{C}_8\text{H}_{17}\text{CHMeONO}$ (165°)

S G $\frac{2}{3}$ 881 Formed by the action of glyceryl trimrite on *sec*-octyl alcohol (Bertoni, *G* 16, 520) Oil, v sol ether

OCTYL OXIDE (C_8H_{16}) $_2\text{O}$ (292°) S G $\frac{2}{3}$ 8204 C E (0°-10°) 00088 S V 403.6 Formed from $\text{C}_8\text{H}_{17}\text{ONa}$ and $\text{C}_8\text{H}_{17}\text{I}$ (Möslinger, *A* 185, 56, Dobner, *A* 243, 10)

***o* OCTYL-PHENYL-AMINE** $\text{C}_8\text{H}_{17}\text{C}_6\text{H}_5\text{NH}_2$ -B'HCl -B' H_2SnCl_4 (Ahrens, *B* 19, 2725)

p Octyl-phenyl-amine (20°) (810° cor) Formed by heating octyl alcohol with aniline and ZnCl_2 at 280° (Beran, *B* 18, 132) -B'HCl -B' H_2SO_4 -B' $\text{H}_2\text{C}_2\text{O}_4$ white plates

Formyl derivative [56°] Plates

Acetyl derivative $\text{C}_8\text{H}_{17}\text{NHAc}$ [93°]

Benzoyl derivative [117°] Plates

The corresponding derivative of *sec*-octyl phenyl amine melts at 109°

***n*-OCTYL-PHOSPHINE** $\text{C}_8\text{H}_{17}\text{PH}_2$ (c 186°) S G $\frac{1}{2}$ 821 Formed by heating octyl iodide with PH_3 and ZnO (Möslinger, *A* 185, 65) -B'HI crystalline

OCTYL SULPHATES

n Octyl-sulphuric acid $\text{C}_8\text{H}_{17}\text{OSO}_3\text{H}$ (Möslinger, *A* 185, 62) Forms a sparingly soluble Ba salt, and an easily soluble K salt

Sec Octyl-sulphuric acid $\text{C}_8\text{H}_{17}\text{OSO}_3\text{H}$ (Bouis, *C R* 33, 144, 88, 935) -BaA' 9aq -KA' aq pearly crystals

***n* OCTYL SULPHIDE** (C_8H_{17}) $_2\text{S}$ (above 810°) S G $\frac{1}{2}$ 842 From the chloride and K_2S (Möslinger, *A* 185, 59)

Sec OCTYL SULPHOCYANIDE $\text{C}_8\text{H}_{17}\text{NS}$ (c) $\text{C}_8\text{H}_{17}\text{CHMeSCy}$ (142°) From the iodide and potassium sulphocyanide (Jahn, *B* 8, 805)

OCTYL-THIENYL METHYL KETONE

$\text{C}_8\text{H}_{17}\text{S}(\text{C}_6\text{H}_5)_2\text{COCH}_3$ (c 852°) Formed, together with only $\text{C}_8\text{H}_{17}\text{S}(\text{C}_6\text{H}_5)(\text{COCH}_3)$, (which yields an oxim [58°]), by the action of AcCl on octyl thiophene in presence of AlCl_3 (Schweinitz, *B* 19, 646) Oil, volatile with steam

***n*-OCTYL-THIOCARBIMIDE** $\text{C}_8\text{H}_{17}\text{NCS}$ (232°) Formed from *n*-octylamine (Jahn, *B* 8, 804, *M* 3, 173)

(a)-OCTYL-THIOPHENE $\text{C}_8\text{H}_{17}\text{C}_6\text{H}_4\text{S}$ (258°). S G $\frac{20.5}{274}$ 8118. Formed from octyl

bromide, (a) iodo thiophene and Na in ether (Schweinitz, *B* 19, 644) Oil

References — Baomx- and Iodo OCTYL THIO-
PHENE

OCTYL-THIOPHENE DICARBOXYLIC ACID $C_8H_7, C_8HS(CO_2H)_2$ [185°] Formed by oxidising $C_8HS(C_6H_5)(CO_2CH_3)$ with alkaline $KMnO_4$ (Schweinitz, *B* 19, 646) Needles — $BaA''13aq - CuA''23aq - Ag_2A''3aq$ yellow pp

Sec OCTYL-THIO-UREA $C_8H_{17}NHCSNH_2$ [114°] Formed from sec octyl thio carbimide and NH_3 (Jahn, *B* 8, 804, *M* 3, 173) Plates

OCTYL TOYL AMINE $C_8H_{17}Me(C_6H_5)_2NH_2$ (325°) Formed by heating α toluidine with octyl alcohol and $ZnCl_2$ at 280° (Beran, *B* 18, 145) Oil — $B'HCl - B'_2H_2SO_4 - B'_2H_2CO_4$

Acetyl derivative [81°] Needles

Benzoyl derivative [117°] Plates

OCTYL UREA Ennoyl derivative $C_8H_{17}NHCO NH CO C_8H_{17}$ [97°] From ennoio amide, Br, and $NaOHAq$ (Hofmann, *B* 15, 760)

GENANTH DIACETONAMINE v. ACETON-AMINE

GENANTHIC ACID v. HEPTOIC ACID

GENANTHOL v. HEPTOIC ALDEHYDE

GENANTHYLAMINE v. HEPTYLAMINE

GENANTHYLIC ACID v. HEPTOIC ACID

GENANTHYLIDENE v. HEPTENENE

GENOGLUCIN $C_8H_{16}O$ [208.5°] A substance resembling phloroglucin prepared from enolin, the red colouring matter of wine, by potash fusion (Gautier, *Bl* [2] 33, 583) Tables (containing 2aq), m sol water Gives no colour with $FeCl_3$

GENOLIN $C_8H_{16}O_2$? A colouring matter ppd by adding lead subacetate to red wine (Glénard, *C* 47, 268, Gautier, *Bl* [2] 32, 103) or by adding lime (Varenne, *Bl* [2] 29, 109) Its composition is variable

OIAZTHIOLES Derivatives of $\begin{smallmatrix} N & CH \\ | & | \\ N & CH \end{smallmatrix} S$

OILS Liquids nearly or quite insol water In a more restricted sense, the term oil is applied to neutral liquids derived from plants or animals Oils are said to be 'fixed' when they cannot be distilled either alone or with steam without undergoing decomposition, oils that can be so distilled being termed volatile or essential oils Most of the fixed oils are glycerides of stearic, palmitic, and oleic acids (v GLYCERIN) Fatty oils that absorb oxygen from the air and thus become slowly converted into varnishes are termed drying oils, e.g. linseed, hazel nut, hemp, and poppy oils Drying oils contain glycerides of linoleic and similar unsaturated acids (v LINOIC ACID and FAT) Essential oils consist either wholly of hydrocarbons or of mixtures of hydrocarbons with compounds of carbon, hydrogen, and oxygen These oxygenated compounds may be compound ethers (oil of chamomile), phenols (oil of thyme, oil of caraway), ketones (oil of rue), aldehydes (oil of cinnamon), or acids (oil of valerian) Many essential oils deposit a solid (stearoptene) on cooling strongly, leaving a liquid portion (elaeoptene) Most of the essential oils contain terpenes or at any rate hydrocarbons of the formula $(C_5H_8)_n$ (v TERPENE) The individual oils are described in their alphabetical places See also FAT

OLEANDRINE. A poisonous alkaloid [70°-75°], in the leaves of the oleander (Lukomski, *J.*

1861, 546, Betelli, *J* 1875, 783) Minute crystals (by sublimation)

OLEFINES Hydrocarbons, C_nH_{2n} , homologous with ethylene, so called from their property of combining with chlorine and bromine, even in the dark, forming oily dichlorides and dibromides (v vol u p 716)

OLEIC ACID $C_{18}H_{34}O_2$ Mol w 282 [8°] (Schön, *A* 244, 262), [14°] (Gottlieb) (223° at 10 mm), (286° at 100 mm) (Krafft a Noerdlinger, *B* 22, 819) SG 12 808 Occurs as glyceryl ether (triolein, vol u p 622) in most fixed oils and fats (Chevreul, *Recherches sur les corps gras*, p 205, Varrentrapp, *A* 35, 196, Laurent, *A* Ch [2] 65, 149, Gottlieb, *A* 57, 40, Heintz, *P* 83, 555, 89 583, 90, 143, Berthelot, *A* Ch [3] 41, 243) Prepared by saponifying olive or almond oil with potash, decomposing the soap with tartaric acid, heating the separated fatty acids with PbO , extracting lead oleate with ether, shaking the extract with $HClAq$, decanting and evaporating the ethereal solution

Propertus — White needles or (above 14°) oil. Insol water, v sol alcohol, miscible with ether May be distilled by superheated steam at 250° Neutral to litmus, but when impure it absorbs oxygen, becoming acid and rancid Gives a crimson colour (Pettenkofer's reaction) when heated with sugar or furfuraldehyde and H_2SO_4 (Mylus, *H* 11, 492)

Reactions — 1 Yields, on distillation, sebacic, acetic, and hexoic acids, CO_2 , CH_4 , C_2H_4 , and CO (Engler, *B* 22, 593) — 2 Potash fusion gives acetate, palmitate, and hydrogen — 3 Nitrous acid converts it into the isomeric elaidic acid, which is solid — 4 Nitric acid oxidises it to acetic, propionic, butyric, valeric, hexoic, heptonic, octoic, ennoic, decolic, suberic, pimelic, adipic and azelaic acids — 5 Glycerin yields on heating mono and tri olein — 6 Bromine combines, forming di-bromo stearic acid (Overbeck, *J* pr 97, 159) — 7 $HIAq$ and red P at 205° in sealed tubes yield stearic acid (Goldschmidt, *Sitz W* 72, 366, Muter, *An* 2, 63) — 8 Alkaline $KMnO_4$ gives azelaic acid and di oxy stearic acids [137°] (A Savtzeff, *J* pr [2] 31, 541, *Bl* [2] 46, 255) 9 Iodine (1 p.c) in sealed tubes at 270° forms stearic acid (Wilde a Reyehler, *Bl* [3] 1, 295) 10 Conc H_2SO_4 , followed by water, yields oxy-stearic acid [79°], oxy stearic lactone [48°], and oily $C_{18}H_{32}CH(O SO_2H) CO_2H$ (Sabaneeff, *Bl* [2] 46, 819, Geitel, *J* pr [2] 37, 74)

Salts — KA' S 25 in the cold Deliquescent Its solution deposits a gelatinous acid salt when greatly diluted — NaA' S 10 at 12° S (alcohol of SG 82) 5 at 13° S (boiling ether) 1 — LiA' — CaA' , granular pp — SrA' , — BaA' , crystalline — BaH_2A' , flocculent (Gossmann, *A* 86, 322) — MgA' , — PbA' , [c 80°]. White powder, sol boiling ether, turpentine oil, and ligroin — PbA' O [100°] — $Pb_2A'_2O_2$ (Schön), — MnA' , sl sol alcohol, v sol ether — FeA' , insol alcohol, v sol ether — AlA' , sl sol ether — HgA' , grey flocks, insol water, sol alcohol and ether — ZnA' , — CuA' , — AgA'

Methyl ether MeA' SG 12 88 Oil

Ethyl ether EtA' (307°) at 307 mm SG 12 875, 12 870 MM 21.9 at 15° (Perkin)

Glyceryl derivatives v vol u p 622

Amide $C_{18}H_{35}O(NH_2)$. [78°] (R), [75°]

(C) Formed by the action of alcoholic ammonia on oil of almonds or oil of hazel-nuts (Bowney, *C. J.* 7, 200, Carlet, *Bl.* 1859, 78).

Elaïdic acid. [47°] A polymeride of oleic acid produced by the action of nitrous acid on oleic acid. It may also be got by saponifying its glyceryl derivative, elaïdin, which is formed when olive oil is solidified by nitrous acid gas or by mercurous nitrate (Boudet, *A. Ch.* [2] 50, 391, Laurent, *A. Ch.* [2] 65, 149, Meyer, *A.* 85, 174, Gottlieb, *A.* 57, 54) Laminæ (from alcohol), m sol ether May be distilled Acid to litmus On fusion with potash it yields acetate and palmitate Alkaline KMnO_4 forms di-oxy stearic acid Combines with bromine forming di-bromo-stearic acid [27°] (Burg, *Bl.* [2] 8, 191).

Salts— NaA' plates— NaHA' — AgA'

Methyl ether MeA' . SG ¹⁸ 872 Oil

Ethyl ether EtA' . SG ¹⁸ 869

Glyceryl ether $\text{C}_2\text{H}_5\text{A}'$, *Elaïdin* [32°] Nodules, almost insol alcohol.

Amide [94°] Formed from elaïdin and alcoholic NH_3 . Needles

Iso-oleic acid $\text{C}_{18}\text{H}_{34}\text{O}_2$ [45°] Formed by the action of alcoholic potash at 120° on iodo-stearic acid obtained from oleic acid and HI (Michael, *A. Saytzeff*, *J. pr.* [2] 35, 886, 37, 277, Benedikt, *M.* 9, 520). Got also by distilling, at 100 mm pressure, oxy-stearic acid (v **OLEIC** acid, *Reaction* 10) Trimetric plates (from ether), insol water, v sol alcohol, m sol ether

Reactions—1 Yields acetate and palmitate by potash fusion.—2 KMnO_4 forms di-oxy-stearic acid [78°]—3 Bromine forms a liquid dibromide—4 HI yields an oily iodostearic acid.

Salts— NaA' — ZnA' .

OLEIN v **GLYCERIN**, *Oleyl derivative*.

OLIBANUM *Incense* A gum resin exuding from *Boswellia papyrifera*. It contains 7 p.c. essential oil, 72 p.c. of resin soluble in alcohol, and 20 p.c. of gum (arabin) (Stenhouse, *A.* 35, 306, Kurbatoff, *Z.* [2] 7, 201, *A.* 173, 1, of Bracconnot, *A. Ch.* [2] 58, 60) The essential oil contains a terpene, olubene $\text{C}_{10}\text{H}_{16}$ (157°), SG ¹² 863, and an oxidised substance Olubene forms a hydrochloride $\text{C}_{10}\text{H}_{15}\text{HCl}$ [127°]

OLIVE OIL SG ²⁰ 918 (Long, *Am.* 10, 892) An oil expressed from olives (*Olea europæa*) It contains the glyceryl ethers of oleic and palmitic acids. The glycerides of unsaturated acids are present in greater quantity (87 p.c.) than those of saturated fatty acids (13 p.c.) (Hazura, *G.* Grissner, *M.* 10, 248)

OLIVIL $\text{C}_{18}\text{H}_{34}\text{O}_2$ [120°] A neutral substance occurring, together with resin and a little benzoic acid, in the gum of the olive tree The resin is removed by ether, and the residue crystallised from alcohol (Pelletier, *A. Ch.* [2] 3, 105, 51, 196, Sobrero, *A.* 54, 67, Amato, *G.* 8, 88) Anhydrous crystals (from alcohol) or prisms containing Aq (from water) Reduces silver salts. Alkaline KMnO_4 yields vanillin Conc HClAq yields MeI and EtI .

OMICHOLIN $\text{C}_{18}\text{H}_{34}\text{NO}_2$? A red resin obtained, together with the similar omicholic acid from urine (Thudichum, *C.* R 106, 1803)

OMPHALOCARPIN A neutral substance in the fruit of *Omphalocarpum Procerum* (Naylor, *Ph.* [3] 12, 478) Needles (from alcohol)

ONOCERIN $\text{C}_{18}\text{H}_{34}\text{O}_2$. Occurs, together with ononin, in the root of *Ononis spinosa* (Hlasiwetz, *J. pr.* 65, 142). Hair-like crystals (from alcohol) Chlorine forms resinous $\text{C}_{18}\text{H}_{32}\text{Cl}_2\text{O}$.

ONONIN $\text{C}_{18}\text{H}_{32}\text{O}_{11}$ [235°] A glucoside occurring in the root of the spinous rest-harrow (*Ononis spinosa*) (Reinsch, *Rep. Pharm.* [2] 26, 12, Hlasiwetz, *Sitz. W.* 15, 142) Tasteless needles (from alcohol), sl. sol boiling water, nearly insol ether Its alcoholic solution is ppd by lead subacetate

Reactions—1 Boiling baryta-water splits it up into formic acid and crystalline onospin $\text{C}_{18}\text{H}_{34}\text{O}_{12}$ [162°]—2 Dilute H_2SO_4 yields glucose and formonetin $\text{C}_{18}\text{H}_{32}\text{O}_{10}$, which is further split up by baryta-water into formic acid and ononetin $\text{C}_{18}\text{H}_{32}\text{O}_8$ [120°] Formonetin and ononetin crystallise from alcohol Ononetin and onospin are coloured red by FeCl_3 .

OPHIOXYLIN $\text{C}_{18}\text{H}_{34}\text{O}_8$? [72°] S 15 at 100°, S (alcohol) 33 at 78° Occurs in the root of *Ophioxylon serpentinum* (Bethink, *R. T. C.* 8, 819) Dimetric needles Conc H_2SO_4 colours it blood-red and, on warming, indigo blue Its alkaline solution is violet

OPILANIC ACID $\text{C}_{18}\text{H}_{34}\text{O}_8$, i.e.

$\text{C}_6\text{H}_5(\text{OMe})_2(\text{CHO})\text{CO}_2\text{H}$ Mol w. 210. [146°]

An acid obtained by the oxidation of narcotine (*q. v.*) (Liebig, *A. Wöhler*, *A.* 44, 126, 50, 1, Blyth, *A.* 50, 29, Anderson, *Tr. E.* 20 [2] 347, Matthiessen, *A. Foster*, *C. J.* 16, 845, 21, 357) It is also formed by the action of KMnO_4 and dilute HNO_3 on hydrastin (Freund, *A. Will*, *B.* 19, 2799) Purified by passing a rapid current of nitrous acid gas (which does not affect it) through its boiling aqueous solution (Prinz, *J. pr.* [2] 24, 355) It may also be purified by etherification, the ether being subsequently decomposed by boiling with water (Liebermann, *A. Kleemann*, *B.* 20, 881)

Properties—Needles or prisms, v sol alcohol, ether, and hot water Tastes bitter

Reactions—1 Readily oxidised to hemipic acid—2 Reduced by sodium-amalgam to meconin—3 Boiling conc KOHAq yields hemipic acid and meconin—4 Conc HClAq at 100° yields MeCl and $\text{C}_6\text{H}_5(\text{OMe})(\text{OH})(\text{CHO})\text{CO}_2\text{H}$. HIAq acts in like manner Dilute HClAq at 170° forms isovanillin $\text{C}_6\text{H}_5(\text{OMe})(\text{OH})\text{CHO}$ and protocatechuic aldehyde—5 HNO_3 forms nitro-opilanic acid, nitro-hemipic acid, and a compound $\text{C}_{18}\text{H}_{32}\text{N}_2\text{O}_{11}$ —6 PCl_5 yields a chloride reduced by zinc and HCl to meconin—7 Heating with malonic acid, HOAc , and NaOAc forms

$\text{C}_6\text{H}_5(\text{OMe})_2<\frac{\text{CO}}{\text{CH}(\text{CH}_2\text{CO}_2\text{H})}>$ (Liebermann, *B.* 19, 2284)—8 H_2SO_4 (80 pts) at 180° forms rufopin, a red colouring matter (Anderson, *C. J.* 9, 277, Liebermann, *A. Chojnacki*, *B.* 4, 194)—9 Distillation with soda-lime yields methyl-vanillin. 10 *Tolylene-o-diamine* forms a crystalline compound [248°] (Bistrzycki, *B.* 21, 2528)—11 Benzidine yields $\text{C}_{18}\text{H}_{32}\text{N}_2\text{O}_8$ (above 320°)—12 Urea forms $\text{C}_{11}\text{H}_{12}\text{N}_2\text{O}_8$ [259°]—13 *Hydrasobenzene* produces $\text{C}_6\text{H}_5(\text{OMe})_2<\frac{\text{CO}}{\text{CH}(\text{NPh NHP})}>$ [188°]

14 An alcoholic solution of *hydroxylamine hydrochloride* forms, on boiling, hemipimide, but in the cold it yields the isomeric oxim-anhydride $\text{C}_6\text{H}_5(\text{OMe})_2<\frac{\text{COO}}{\text{CH N}}>$ [115°] (Liebermann, *B.* 19,

2928). This body suddenly changes, at its melting point, to hemipimide, with great rise of temperature.—15 *Aniline* in HOAc forms, on boiling, $C_6H_5(OMe)_2(CO_2H)CH NPh$ [187°].—16 *Ammonia* forms opiammon $C_{22}H_{21}NO_3$, a yellowish powder, decomposed by water at 150° into NH_3 and opianic acid.—17 Aqueous *sulphurous acid* forms crystalline $C_{17}H_{15}O_3H_2SO_4$, whence the salts BaA' , 8aq and PbA' , 6aq may be obtained.—18 H_2S forms yellow amorphous $C_{16}H_{15}SO_4$.—19 *Phenol* and H_2SO_4 in the cold form amorphous opiaurin $C_{20}H_{19}O_3$, which forms a purple solution with potash—20 *Phenylhydrazine acetate* forms $C_{16}H_{15}N_2O_3$ crystallising in needles [175°] (Liebermann, *B* 19, 763)—21 *Salicylic acid* and $ZnCl_2$ form $C_{17}H_{15}O_3$ [140°], whence $C_{17}H_{15}AcO_3$ [152°] (Michael, *Am* 5, 95)

Salts— $KA'2aq$ (Wegscheider, *M* 8, 348)— KA' aq triclinic plates— BaA' , 2aq efflorescent prisms— PbA' , 2aq sparingly soluble crystals [150°]— PbA' , tufts of silky prisms— AgA'

Methyl ether MeA' [85°] (W), [102°] (L) Formed from the silver salt and MeI, or by boiling the acid with MeOH. Monoclinic

Ethylether EtA' [92°]

Propylether PrA' [103°]

Acetyl derivative $C_{16}H_{15}AcO_3$ [121°]

Propionyl derivative [111°] Needles

Anhydride $\{C_{16}H_{15}(OMe)_2(CHO)CO\}_2O$

Tropianide [234°] Formed by heating the acid for two hours in a current of dry air Needles (from acetone)

Oxim $C_6H_5(OMe)(CO_2H)CH NOH$ [83°] Formed from the acid (1 mol), dilute KOH (4 mols), and hydroxylamine hydrochloride (1½ mols) in the cold (Perkin, *jun*, *C J* 57, 1071) Long slender needles, v sol alcohol At 130° it suddenly becomes solid owing to the formation of hemipimide

Di-phenyl-hydrazide

$C_6H_5(OMe)_2(CO_2H)CH N NPh_2$ [172°] CaA' 8aq

Chloro-opianic acid

$C_6HCl(OMe)(CHO)CO_2H$ [211°] Needles (Prinz, *J pr* [2] 24, 366)

Bromo-opianic acid [192°] (P), [204°] (Wegscheider, *M* 4, 267) Needles— BaA' , aq

Nitro-opianic acid

$C_6H(NO_2)(OMe)_2(CHO)CO_2H$ [166°] Formed by nitration of opianic acid (Prinz) Yellow prisms (from water) With acetone and dilute NaOHAq it yields the tetra-methyl derivative of tetra oxy indigo dicarboxylic acid (Liebermann, *B* 10, 352) Aniline in HOAc reacts, forming $C_6H(NO_2)(OMe)_2(CO_2H)CH NPh$ [184°]— KA' 8aq prisms— BaA' , 3aq yellow needles.

Ethylether EtA' [96°]

Acetyl derivative $C_{17}H_{15}AcNO_3$. Crystals

Phenyl-hydrazide

$C_6H(OMe)_2(NO_2)(OOH)CH N_2HPh$ [184°] Red needles (Liebermann, *B* 19, 764) By boiling with HOAc it is converted into 'nitro-opianyl-phenyl-hydrazine' $C_{16}H_{15}N_2O_3$ [173°]

Di-phenyl-hydrazide $C_{22}H_{21}N_2O_3$ [217°] Prisms, sl. sol ether (Bastrzycki, *B* 21, 2520)

Nitroso-opianic acid?

$C_6H(OMe)_2(NO)(CHO)CO_2H$ [5 5 3 4 1] [176°] Formed by reducing nitro opianic acid with NaOMe (Kleemann, *B* 20, 875) Long yellowish-green needles.— AgA' . felted needles.

Amido-opianic acid

$C_6H(NH_2)(OMe)_2(CHO)CO_2H$. Formed by reduction of nitroso-opianic acid (K_2).— HA' . HCl.

Acetyl derivative $C_{17}H_{15}N_2O_3$. [233°].

Phenyl-hydrazide

$C_6H(NH_2)(OMe)_2CO NPh$ [148°]. Needles (Liebermann, *B* 19, 2276)

Noropianic acid $C_6H_4(OH)_2(CHO)CO_2H$ [171° cor] Formed by heating opianic acid with HIAq Crystals (containing 1½aq)

Methyl derivative

$C_6H_4(OMe)(OH)(CHO)CO_2H$ [154°] (P), [142°] (Wegscheider, *M* 8, 790) Formed by heating opianic acid with HClAq at 100° Prisms (containing 2aq) Chlorine forms the chloro-derivative $C_6H_4ClO_2$ [206°] On nitration it yields $C_6H(NO_2)(OMe)(OH)(CHO)CO_2H$ [208°], which crystallises in needles (containing aq), and yields an oxim decomposing at 252° and converted by HOAc into the anhydride $C_6H_4N_2O_3$ [252°], the nitro-acid also yields a phenyl-hydrazide [179°] converted by HOAc into an anhydride [191°] (Elbel, *B* 19, 2306)— KA' 2aq— $BaC_6H_4O_3$ aq

Iso opianic acid v Methyl derivative of ALDEHYDO VANILLIC acid

Iso-noropianic acid $C_6H_4(OH)_2(CHO)CO_2H$ [4 3 5 1] [above 240°] Formed by heating aldehyde vanillic acid with HClAq at 175° (Tiemann a Mendelssohn, *B* 10, 400) Needles

Pseudopianic acid

$C_6H_4(OMe)_2(CHO)CO_2H$ [1 2 3 4] [122°] A product of the action of boiling dilute H_2SO_4 on berberol (Perkin, *jun*, *C J* 57, 1065) Slender needles (from hot water) When warmed with phenol and H_2SO_4 it gives a green colour, changing to violet brown— KA' — AgA' needles

Oxim $C_{10}H_9O_4(NOH)$ [124°] Needles

OPIANYL ACETIC ACID $C_{17}H_{15}O_3$, i.e. $C_6H_4(OMe)_2(CO_2H)CH(OH)CH_2CO_2H$ Formed, as barium salt, by boiling meconin-acetic acid (p 198) with baryta water (Liebermann a Kleemann, *B* 19, 2292) The acid when liberated from its salts at once loses H_2O and is reconverted into its lactone meconin acetic acid.— $A'Ag$ crystalline pp— A' , $BaAq$ prisms

OPINIC ACID $C_6H_5O_3$ [148°] Formed by the action of HI on hemipic acid (Liechti, *Z* [2] 6, 196, *A Suppl* 7, 151, Beckett a Wright, *J* 1876, 809) Prisms (containing 2aq) Coloured violet by FeCl.

OPIONIN [227°] A substance present in small quantity in Smyrna opium (Hesse, *A* 228, 299) Small needles, v sol alcohol, ether, and alkalis, v sl sol water Neutral to litmus When fused with potash it yields opionic acid [126°]

OPIUM The dried juice obtained from the unripe capsules of *Papaver somniferum* It contains morphine, codeine, narcotine, narceine, thebaine, papaverine, meconic acid, and meconin The quantity of morphine varies from 8 to 15 p c Hesse (*A* 153, 47, *Suppl* 8, 299) obtained also codamine, lanthopine, laudanine, meconidine, cryptopine, protopine, laudanone, and hydrocotarnine Kauder describes tritopine (*Ar Ph* 238, 419) The three alkaloids morphine, codeine, and thebaine are strong bases, while papaverine, narcotine, and narceine are feeble bases The various constituents of opium are described in their alphabetical position

OPOPANAX The dried juice obtained from the roots of the *Pastinaca Opopanax*. It contains 33 p.c. of gum, 28 p.c. of malic acid, and 42 p.c. of a resin which yields protocatechuic acid on fusion with potash (Pelletier, Hlasiwetz & Barth, *J* 1866, 630, Hirschsohn, *C* 1877, 182).

ORANGE PEEL OIL SG $\frac{30}{100}$ 8435 The essential oil of orange peel consists mainly of a terpene (174°), but contains also an aldehyde $C_{10}H_{16}O$ (224°–228°) (Wright, *C* J 18, 1186, 20, 552, Semmler, *B* 24, 202). It contains no cymene (Hartley, *C* J 37, 677). Tanret (*Bl* [2] 46, 501) got from orange peel resinous 'aurantiamaric acid' $C_{16}H_{24}O_4$, $[\alpha]_D = -28^\circ$, and uncrystallisable 'aurantiamarine' $C_{22}H_{36}O_{12}$? $[\alpha]_D = -60^\circ$, a very bitter substance.

ORCEIN $C_{22}H_{22}N_2O_4$. A colouring matter obtained from orcein by the action of aqueous ammonia and air. Purified by exhausting with ether, and crystallising from dilute alcohol (Zulkowsky & Peters, *M* 11, 227, cf. Robiquet, *A* Ch [2] 42, 245, 58, 320, Dumas, *A* 27, 145, Laurent & Gerhardt, *A* Ch [3] 24, 315, Liebermann, *B* 7, 247, 8, 1649). Micro crystalline mass, with metallic lustre, insol. water, ether, and CS_2 . Its solutions are crimson, but are turned bluish violet by ammonia or potash.

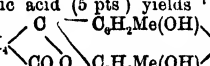
ORCENE DIALDEHYDE This name is given by Tiemann and Helkenberg (*B* 12, 1003) to two isomerides $C_8HMe(OH)_2(CHO)_2$, (a) [118°] and (b) [168°] formed, together with orceyl aldehyde $C_8H_8O_4$, by the action of chloroform and KOH upon orcein. The (a)-isomeride forms a di-anilide $C_8HMe(OH)_2(CH_2NPh)_2$, [281°].

ORCIN $C_8H_8O_4$ & $C_8H_8Me(OH)$ [135] *Di-oryz toluene Methyl resorcin*. Mol w 124 [108°] (Neville & Winther, *C* J 41, 417), [57°] (when containing aq.) (c 288) HFp 109,276 (Stohmann, *J* pr [2] 34, 315). Obtained by boiling orsellic acid with water or by action of alkalis on lecanoric, erythric, and evermic acids and picroerythrin, and also by dry distillation of the same bodies (Robiquet, *A* Ch [2] 42, 245, 58, 320, Liebig & Will, *A* 27, 147, Dumas, *A* 27, 140, Schunck, *A* 41, 159, 54, 269, Stenhouse, *Tr* 1848, 85, *A* 68, 93, 99, *Pr* 12, 263, *C* J 16, 327, 20, 223, De Luynes, *A* 128, 330, 130, 81, Lamparter, *A* 134, 256). Formed also by potash fusion from aloes (Hlasiwetz & Barth, *A* 134, 287), *p*-chloro toluene sulphonic acid (Vogt & Henniger, *C* R 74, 1107), toluene *m* disulphonic acid, *s*-bromo cresol, *s* di bromo toluene, and *s* bromo toluene sulphonic acid (Neville & Winther, *C* J 41, 417). Formed also from nitro-*m* cresol by reduction and application of the diazo reaction.

Synthesis — Acetone-*di*-carboxylic ether $OC(CH_3)_2CO_2Et$, when treated with sodium condenses to *di*-oxy phenyl acetic di-carboxylic ether $C_6H_4(OH)_2(CO_2Et)_2$, CH_2CO_2Et , the latter by the energetic action of alkalis is converted into *di*-oxy phenyl-acetic acid $C_6H_4(OH)_2CH_2CO_2H$, whose silver salt on dry distillation yields orceinol (Cornelius & Pechmann, *B* 19, 1446).

Properties — Monoclinic prisms (containing aq.), v. sol. water, alcohol, and ether. Sweet taste. Turns red in air. Its aqueous solution gives a white pp. with lead subacetate and a dark red pp. with $FeCl_3$. It reduces ammoniacal $AgNO_3$. Dry bromine forms tri-bromo-orcin

Bromine water yields penta-bromo-orcin (Stenhouse & Groves, *C* J 37, 463). Its solution is not rendered acid by borax. Nitrous acid gives an orange pink colour in dilute solution. (Lunde, *C* N 58, 1, 15). It forms a deliquescent compound with picric acid.

Reactions — 1. Air and ammonia yield orcein together with yellow $C_{10}H_8NO_3$ soluble in ether and an amorphous body, resembling litmus, in soluble in ether and alcohol (Zulkowsky & Peters, *M* 11, 227). Dry ammonia forms colourless crystals of $C_8H_8O_4NH_3$. — 2. Ammonium carbonate yields 'para orsellic' acid $C_8H_8O_4$ aq [151°] (Senhofer & Brunner, *C* J 40, 265). — 3. KOH and CO, form pseudo orsellic acid (Schwarz, *B* 13, 1643). — 4. Heating with HOAc (1½ pts) and $ZnCl_2$ (2 pts) forms crystalline $C_{22}H_{22}O_4$ as well as 'orcacetem' $C_{18}H_{18}O_4$, a yellow powder (Rasinski, *J* pr [2] 26, 56). On boiling orcein (9 g) with HOAc (135 g) and phosphorus oxychloride (18 g) there is formed $C_8HMe(OH)COCH_3$ [146°] crystallising in needles — 5. Orcin (3 pts) heated with acetoacetic ether (2 pts) and a little H_2SO_4 at 100° forms insoluble $C_{11}H_{12}O_4$ [249°] which yields $C_7H_7AcO_3$ [200°] and $C_{11}H_{12}Br_2O_3$ crystallising in plates — 6. HNO_3 saturated with HNO_2 acting upon an ethereal solution of orcein forms a scarlet dye 'azo orcein' $C_{11}H_7NO_3$ which forms an alkaline solution with orange red fluorescence (Kramer, *B* 17, 1832). — 7. H_2SO_4 saturated with nitrous acid forms a dye $C_{22}H_{22}NO_3$ which yields alkaline solutions with red fluorescence and $C_{22}H_{22}NO$, which does not yield fluorescent solutions (K, Brunner, *B* 21, 251). — 8. A mixture of HNO_3 and HBr yields $C_8H_8BrNO_3$ and $C_{10}H_{12}Me_2BrN_2O_{11}$, the alkaline solutions of these bodies exhibit brown and red fluorescence respectively (Brunner, *B* 21, 2494). — 9. A mixture of HNO_3 (10 c.c. of SG 1.39) and HCl (30 c.c. of SG 1.2) on the water bath yields $C_{22}H_{22}ClNO_3$, a greenish mass which yields $C_{11}H_{12}AcClNO_3$. — 10. Chloral hydrate and water on boiling form crystalline $C_8H_8O_4$ whence $C_7H_7Ac_2O_3$ [185°] (Michael & Ryder, *Am* 9, 135). — 11. Fusion with NaOH yields resorcin, tetra oxy di phenyl methane, and finally phloroglucin (Barth, *M* 3, 645). — 12. Benzoic aldehyde and a little HCl aq. give a white resin $C_{22}H_{22}O_4$ (M & R). — 13. Chloroform and dilute potash form, on boiling, two isomeric 'orcene dialdehydes' $C_8H_8Me(OH)_2(CHO)_2$, and 'orceyl aldehyde' $C_8H_8Me(OH)CHO$ [177°] (Tiemann, *B* 12, 999). Another product of the action of potash and chloroform on orcein is homofluorescin (vol. II p. 558) which is probably orcein aurin $C_{22}H_{22}O_4$, a body which is formed by heating orcein with formic acid and $ZnCl_2$ at 100° (Nencki, *J* pr [2] 25, 277, Grimaux, *Bl* [3] 3, 850). — 14. Phthalic anhydride (3 pts) heated with orcein (5 pts) and sulphuric acid (5 pts) yields 'orcun-phthalein' $C_{22}H_{22}O_4$  which crystallises from acetone in colourless needles, dissolving in alkalis with red colour, and yielding the acetyl derivatives $C_{22}H_{22}AcO_4$ [150°] and $C_{22}H_{22}Ac_2O_4$ [220°]. Orcin phthalein is reduced by zinc dust and NaOH aq. to orcein phthalin $C_{22}H_{22}O_4$ which yields $C_{22}H_{22}Ac_2O_4$ [211°] (E. Fischer, *A* 183, 72). — 15. Concentrated sulphuric

acid at 70° produces osmin disulphonic acid $C_8H_8Me(OH)_2(SO_3H)_2$ which yields the crystalline salts $Pb_2C_8H_8S_2O_6$, 6aq and $Pb_2(C_8H_8S_2O_6)_2$, 8aq (Hesse, *id* 117, 324)—16 Fuming HNO_3 , acting on osmin dissolved in ether forms orosurfin $C_{14}H_{11}NO_3$, crystallising in needles [225°] which are blue by reflected light and form a crimson solution in alkalis exhibiting yellow fluorescence. Orosurfin yields an acetyl derivative [204°] and an ethyl ether [269°] (Nietzki & Maackler, *B* 23, 720)—17 *Aldehyde* dissolved in alcohol forms on addition of a few drops of $HClAq$, a compound $C_{16}H_{18}O_2$, crystallising in tables (Michaela & Comey, *Am* 5, 349)—18 *Chloro-acetic acid* and caustic soda forms $C_8H_8Me(OCH_2CO_2H)_2$ [217°] crystallising in needles. This acid forms the salts NaA'' 8aq, K_2A'' 8aq, and CaA'' 2aq, the ether Et_2A'' [107°], and two nitro-derivatives one of which melts at 140° (Saarbach, *J pr* [2] 21, 162)—19 *Di chloro quinonumide* in alcoholic solution forms brown needles of 'orosurifamine' $C_{16}H_{11}N_2O_2$, which dissolves in acids with reddish violet colour and orange fluorescence. It dyes silk, and yields a crystalline mono acetyl derivative (N & M)

Acetyl derivative $C_8H_8Me(OAc)_2$ [25°]

Needles, nearly insol water

Benzoyl derivative $C_8H_8Me(OBz)_2$, [88°] Needles (Rasinski, *J pr* [2] 26, 65)

Orthocarbonyl derivative

$(C_8H_8MeO)_2C$ [195°] Formed by distilling $C_8H_8Me(O_2CO_2Et)_2$, which is produced by the action of $ClCO_2Et$ on sodium osmin (Wallach, *A* 226, 86, Bender, *B* 13, 700)

Methyl ether $C_8H_8Me(OMe)(OH)$ (c 273°) Liquid, sol water (Tiemann & Streng, *B* 14, 2001)

Di methyl ether $C_8H_8Me(OMe)_2$ (244°) V D 76.2 ($H=1$) (obs) Mobile liquid

Di ethyl ether $C_8H_8Me(OEt)_2$ [165°] (252° cor) Needles Yields $C_8H_8Br_2Me(OEt)_2$ [144°] (Herzig & Zeisel, *M* 11, 315, cf De Luyne & Lionet, *C R* 65, 213)

Nitroso-derivative $C_8H_8(NO)O_2$ [157°] Yellow needles, detonating at its melting point (Nietzki & Maackler, *B* 23, 723)

References — TRI AMIDO-, AMIDO DI IMIDO-, DI BROMO NITRO, TRI CHLORO, and IODO-OSMIN
M Osmic, I Iso-osmin, Cresosmin, and Lutocin
v DI OXY TOLUENE

β Osmic v BETORCIN

OSMIAL ALDEHYDE v DI OXY-TOLUIC ALDEHYDE

OREOSELIN $C_{16}H_{11}O_4$, *Oreoselone* [170°]

Formed by the action of alcoholic potash or acids on peucedanin (Wagner, *J pr* 62, 275, Hlasiwetz, *A* 174, 70, Heut, *A* 176, 73) Needles (from alcohol), v sl sol water. Its solution in H_2SO_4 exhibits bluish-green fluorescence. Potash fusion yields resorcin and acetic acid. It yields an acetyl derivative [123°] and an isovaleryl derivative [97°]

OREOSELONE $C_{16}H_{11}O_4$ [190°] Formed by passing HCl over dry athamantun (Schneidermann & Winckler, *A* 51, 320) Needles (from alcohol), insol water

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ORNITHINE $C_6H_{11}N_2O_2$. Obtained, together with benzoic acid, by boiling ornithuric acid
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with hydrochloric acid (Jaffé, *B* 10, 1925) — B_2H_7Cl , $-B'HCl$ — $B'_2H_7C_2O_2$, $-B'HNO_2$

Benzoyl derivative $C_8H_{11}BzN_2O_2$ [225°-230°]

ORNITHURIC ACID $C_6H_{11}N_2O_4$ [182°] Excreted by birds after a dose of benzoic acid (Jaffé, *B* 10, 1925, 11, 406) Needles, v sl sol hot water — CaA' , $-BaA'$, powder, v e. sol water

ORSEILLE v ARCHIL

ORSELLIC ACID v DI OXY TOLUIC ACID and LECANORIC ACID

OSAZONES v HYDRAZONES

OSMATES v OSMIUM, Salts of oxyacids of, p. 646

OSMIAMIC ACID v OSMIUM, Nitrogen containing acid of, p. 644

OSMIRIDIUM v IRIIDIUM, Alloys of, p. 47

OSMIUM Os At w 190.3 Mol w unknown Infusible at highest temperature attained by O H flame S G 22.477 at 17.5° (Deville & Debray, *P M* [4] 50, 651) S H 19° to 98° 0.3113 (Regnault, *P M* [4] 23, 103) C E 00000657 (Fizeau, *C R* 68, 1125)

Occurrence — As metal, alloyed with Ir, Pt, Rh, Ru, and Pd. The residues obtained by heating various Pt ores with *aqua regia* contain from 17 to 50 pct Os (v Deville & Debray, *A Ch* [3] 56, 431, Berzelius, *P* 13, 436, 527, 15, 208). In 1804 Tennant showed that the metallic residue remaining after treating Pt ore with *aqua regia* contained two new metals (*T* 1804 411), to one of these metals he gave the name *iridium*, because of the colours of its oxides (*iris*=rainbow), and to the other, the name osmium, because of the peculiar smell of its volatile oxide (*osmum*=smell)

Formation — 1 By ppg K_2OsO_4 by NH_4Cl , and heating the pp in $H-2$ By heating $(NH_4)_2OsCl_4$ or $(NH_4)_2OsCl_6$ — 3 By strongly heating any of the sulphides of Os in absence of air — 4 By warming K persulfate in $HClAq$, with a formate, or by digesting the same salt with Hg, and heating the Os Hg amalgam thus obtained — 5 By passing vapour of OsO_3 mixed with H or CO through a red hot tube — 6 By passing OsO_3 vapour along with N through a hot tube lined with C (obtained by previously passing C_2H_4 vapour through the tube)

Preparation — When Pt ores are treated with *aqua regia*, osmium remains, partly as lustrous tablets, and partly as a black powder. Sand and gangue are removed by fusing the osmium with Na_2CO_3 , boiling with water, then with $HClAq$, and again with water. Os is obtained from this residue by various methods, which are based on the ready oxidation of Os to OsO_3 and the volatility of this compound. The lustrous tablets of osmium yield much more Os than the black powder. OsO_3 may be obtained by the following methods. — 1 The osmium is heated to redness in a tube of porcelain, or Pt, while a current of air or O (previously passed through H_2SO_4) is sucked through the tube, the exit end of the tube is connected with a dry flask, or a series of dry Woulff's bottles, kept at a low temperature, the connecting tubes must be wide, else they may get choked with crystals of OsO_4 , a vessel containing KOHAq is placed between the receiver and the aspirator, to absorb the last traces of OsO_3 ,
T T

(Fremy, *J pr* 63, 342) — 2 The osm-iridium is very well mixed with an equal weight of dry NaCl, and the mixture is heated to low redness in a rapid stream of moist Cl, in a glass or porcelain tube, the apparatus is connected with a wide necked tubulated receiver, from which a tube passes into KOHAq or NH₄Aq. OsCl₃ is formed, and decomposed by the H₂O present to OsO₃, HCl, and Os which again combines with Cl to form OsCl₃, which OsCl₃ is again decomposed by H₂O, and so on (Wohler, *P* 31, 161, 104, 368, 107, 357) Schneider (*A Suppl* 5, 261) places the osm iridium in a large Hessian crucible, the lid of which is cemented on by CaSO₄ and carries a porcelain tube passing into a receiver — 3 The osm-iridium may be treated with molten KOH and KClO₄ (Fritzsche, *J pr* 37, 483), or with KOH and KNO₃ (Claus, *C O* 1862 129, Gibbs, *A* 120, 108), or with BaO₂ and Ba(NO₃)₂ (Deville a Debray, *A Ch* [3] 56, 481), in each case the K persmate obtained is distilled with *aqua regia* and OsO₃ is obtained. The preparation of OsO₃ from osm iridium is easily accomplished, but the process is extremely unpleasant, and also very dangerous, inasmuch as the vapour of OsO₃ rapidly attacks the eyes and lungs.

Os is obtained from OsO₃ by reducing the vapour by H or CO, or by C (Deville a Debray, *lc*, cf *Formation*, Nos 5 and 6), or a solution of OsO₃ in KOHAq may be reduced by warming with alcohol, the violet red crystalline pp of K₂OsO₄ may be dissolved in water, ppd by NH₄Cl, and the pp reduced to Os by heating in H (Fremy, *lc*). Or the OsO₃ may be dissolved in NH₄Aq, warmed for some hours till blackish brown (N is evolved, and an ammoniacal solution of Os₂O₃ remains), and evaporated until a brown pp of impure Os₂O₃ forms, the pp may then be washed, dissolved in HClAq, NH₄Cl added, the solution evaporated to dryness, and the residue heated in a retort as long as HCl is evolved and NH₄Cl sublimes. Os remains as a compact porous mass (Berzelius).

Os may be obtained in crystals by dissolving the metal in Sn, by heating to redness in a carbon crucible with 8 parts Sn, and allowing the mass to cool (Deville a Debray, *C R* 82, 1076). The crystals are purified by dissolving out Sn, and heating the residue in a stream of HCl. Os is also obtained in crystals by passing the vapour of OsO₃, mixed with N, through a hot porcelain tube lined with C. The lining of C is obtained by passing C₂H₄ vapour through the tube at a temperature high enough to decompose the C₂H₄. By alloying Os with Zn, and treating the alloy with HClAq, Os is obtained as a black powder which is easily ignited (D a D, *A Ch* [8] 56, 399).

Properties and Reactions — As obtained by reducing its compounds at high temperatures, Os is a lustrous, blue-white metal, resembling Pt or Sn. Prepared by reduction at lower temperatures, Os is a greyish-black powder, without metallic lustre. From solution in Sn, Os separates in hard, bluish crystals, probably belonging to the regular system, *S G* 22 477. Deville a Debray heated Os to the temperature at which Rh melts (c 2000°) without fusing it. The Os was placed in a carbon-crucible placed in a cylinder of lime, resting on a block of

lime, and surrounded by three other similar blocks, a powerful O H flame was caused to play over, and around, the lime cylinder (*A Ch* [3] 56, 386, 61, 5).

Os is easily oxidised to OsO₃. The finely divided metal obtained at low temperatures smells of OsO₃ at the ordinary temperature, when heated in air, it burns and is completely volatilised. A compact mass of Os ignites when strongly heated in air, but ceases to burn when the source of heat is removed, Os prepared at very high temperatures may be heated to c 225° without change. Finely divided Os, which has not been strongly heated, is oxidised to OsO₃ by heating with conc HNO₃ or *aqua regia*, if the metal has been exposed to a very high temperature, it cannot be oxidised in the wet way. Os is oxidised, to K₂OsO₄, by molten KOH, the oxidation is hastened by addition of KNO₃ or KClO₄. Heated in Cl, OsCl₃ is produced. Os forms a phosphide when heated in P vapour. The spectrum of Os has been examined by Huggins (*T* 154, 139), and Thalén (*v B A* 1884 431).

The atomic weight of Os has been determined (1) by heating in O and weighing the OsO₃ produced (Berzelius, *P* 13, 546, Fremy, *A Ch* [3] 12, 514), (2) by reducing KOsCl₂ in H, washing out KCl, and weighing Os (Berzelius, *P* 13, 530, Seubert, *B* 21, 1839, *A* 261, 257), (3) by determining S H of Os (Regnault, *P M* [4] 23, 103). The older values were too high. Seubert's analyses of K₂OsCl₆ and (NH₄)₂OsCl₆, conducted in 1888, gave the mean value 191.12, but this number was regarded by S as rather too high, in 1891 S re determined the at w very carefully and obtained the value 190.3.

Os is distinctly metallic in its physical characters, but some of its chemical properties are those of a non metal, *eg* acidic character of OsO₃, and existence of salts the negative radicle of which is OsO₃. Os belongs to the group of Pt metals, to which group Cu, Ag, and Au are more or less closely related (*v* COPPER GROUP OF ELEMENTS, 11 250, and NOBLE METALS, this vol p 628). Os is more closely related to Ru than to the other Pt metals, the analogy is shown in the existence of volatile oxides MO, capable of existing as gases, in the salts K₂MO₄, and probably KMO₄, and in other respects. The existence of K₂OsO₄, the fact that this salt is oxidised probably to KO₂Os, which salt is reduced to K₂OsO₃ by KOHAq, and also the existence of H₂OsCy₂, and salts of this acid, emphasise the analogies between Os on the one hand, and Mn and Fe on the other hand.

Detection and Estimation — Compounds of Os heated with a little Na₂CO₃ on Pt foil give the peculiar, chlorine like, odour of OsO₃, at the same time the non-luminous flame becomes luminous from separation of Os. Salts of OsO₃ are easily oxidised, *eg* solution of OsCl₃, which is deep blue, becomes violet in air, and then dark red (OsCl₂), and finally yellow (OsCl₄). Salts of OsO₃, *eg* K₂OsCl₆, give a brown red pp (Os₂O₃.2H₂O) with KOHAq, a grey brown pp sol in NH₄Aq, with AgNO₃Aq, and a blue colour (reduction to OsCl₂) on warming with tannic acid. Salts of OsO₃, *eg* K₂OsCl₆, give no pp with KOHAq in the cold, but on warming the liquid goes blue, and then black Os₂O₃.2H₂O.

suddenly separates Borax produces no change at ordinary temperatures, but a black pp forms on warming, this distinguishes OsO_2 salts from those of IrO_2 , which become blue on heating with borax, and after a little give a blue pp. Tannic acid does not react in the cold, on warming, the liquid becomes blue. Alkali salts of OsO_2 form violet solutions in water. HNO_3 oxidises to salts of OsO_4 , NH_4Aq followed by NH_4ClAq ppts yellow $\text{OsO}_2(\text{NH}_4, \text{NH}_4\text{Cl})_2$, which gives Os on heating OsO_4 , or OsO_4Aq , is recognised by its smell, addition of KOH Aq to dilute solutions of OsO_4 removes the smell. A delicate test for Os consists in fusing with KOH and a little KNO_3 , dissolving K_2OsO_4 formed in water, adding NH_4Cl , washing the yellow pp of $\text{OsO}_2(\text{NH}_4, \text{NH}_4\text{Cl})_2$ with very dilute HClAq , dissolving it in slightly warm water, and adding $\text{K}_2\text{FeCy}_6\text{Aq}$, when a splendid violet colour is produced (Gibbs, *Am J*, 233). Os is usually estimated as the metal. Os is separated from other metals by dissolving in *aqua regia*, adding HNO_3 , and distilling. The OsO_4 may be led into excess of NH_4Aq , the solution is saturated with H_2S and boiled, the pp of OsS_2 is collected, washed, dried, and placed in a small carbon crucible which is placed inside a Hessian crucible, the space between being filled with sand, and heated for some hours to c. 220° – 250° , compact Os is thus obtained.

References—Tennant, *T* 1804 411 Berzelius, *P* 13, 435, 527, 15, 208 Wöhler, *P* 31, 161 Claus a Jacobi, *J pr* 85, 142, 90, 65, A 63, 355 Deville a Debray, *A Ch* [3] 56, 393 Fremy, *A Ch* [3] 12, 522, 44, 391 Fritzsche a Struve, *J pr* 41, 97

Osmium, acids of No oxyacid of Os has been isolated, but salts of the hypothetical acids H_2OsO_4 and HOsO_4 are known (*v Osmium, salts of oxyacids of*, p 646). A peculiar acid, $\text{H}_2\text{N}_2\text{Os}_2\text{O}_8$, generally known as *Osmamic acid*, exists (*v Osmium, nitrogen containing acid of*, p 644). *Osmocyanhydric acid*, H_2OsCy_6 , and salts of this acid have been obtained (*v CYANIDES*, vol II p 343). Salts of the hypothetical acids H_2OsCl_4 and HOsCl_4 are known (*v Osmium, chlorides of*, p 644).

Osmium, alloys of Alloys with copper and gold were described by Tennant (*T* 1804 411) as very ductile, insoluble in *aqua regia*. An amalgam with mercury is obtained by the reaction of Hg with OsO_4Aq (*v Claus, J pr* 90, 65). The alloy with iridium occurs native, composition varies between c. Os, Ir and Os, Ir, this alloy usually contains Ru, Rh, and Au, besides Os and Ir (*v Iridium, Alloys of*, this vol p 47). Alloys of Os and Ir were obtained by Deville by melting Os and Ir with Cu_2S , and treating the fused mass with HClAq and then with HNO_3Aq (*M S* 1882 1228).

Osmium, ammonio-salts of, or Osm ammonium salts (*Ammoniacal osmium bases*) Not many of these compounds have been prepared. Those which are known seem best looked on as two or more NH_4 groups with H replaced by the radicle OsO or OsO_2 . The constitutional formulæ and the nomenclature of the osm ammonium salts are not settled, in this article compounds supposed to contain the radicle OsO are called *Osmosyl-ammonium compounds*, and those supposed to contain the radicle OsO_2 are called

Osmyl ammonium compounds (after the analogy of NO and NO_2 compounds).

OSMOSYL DIAMMONIUM HYDROXIDE

$\text{OsO}(\text{NH}_4, \text{OH})_2$ (*Ammoniated oxyosmous hydroxide*) *Oxyosmiumammonium hydroxide* Simplest empirical formula = $\text{OsO}_2 \cdot 2\text{NH}_4 \cdot \text{H}_2\text{O}$. Obtained by dissolving OsO_4 in excess of conc NH_4Aq , and warming the red solution in a closed flask to 40° – 60° until a black pp begins to form when the flask is opened (Claus a Jacobi, *J pr* 79, 28). N escapes during the process ($73\text{OsO}_4 + 10\text{NH}_4 = 8\text{OsO}(\text{NH}_4, \text{OH})_2 + 2\text{N}_2 + 3\text{H}_2\text{O}$). Forms a

brown black powder, decomposed by heat to Os with evolution of N and H_2O , insol water, sol acids, and reppd by KOH Aq or NH_4Aq . Amorphous basic salts (not fully examined) are obtained by evaporating the acid solutions, these salts are decomposed by water to neutral and more basic salts (Claus a Jacobi, *J pr* 90, 65, cf Berzelius, *P* 13, 435, 527). The base dissolves in KOH Aq , on boiling, NH_3 is evolved and a pp, $\text{OsO}_2 \cdot \text{H}_2\text{O}$ containing NH_3 , separates.

OSMOSYL TETRAMMONIUM HYDROXIDE

$\text{OsO}_2(\text{NH}_4, \text{NH}_4, \text{OH})_2\text{Aq}$ (*Ammoniated oxyosmic hydroxide*) *Osmidrammonium hydroxide* *Oxyosmiumdiammonium hydroxide* *Osmyl-ditetramine hydroxide* Simplest empirical formula = $\text{OsO}_2 \cdot 4\text{NH}_4 \cdot \text{H}_2\text{O Aq}$. Known only in solution. A solution of this base is obtained by decomposing the chloride by Ag_2O and H_2O , or the sulphate by BaO Aq (*v infra*), and filtering, the solution is yellow, has an alkaline reaction, and is easily decomposed with formation of OsO_4 , NH_3 , and a black pp (Claus, *J pr* 79, 28).

OSMIL TETRAMMONIUM CHLORIDE

$\text{OsO}_2(\text{NH}_4, \text{NH}_4\text{Cl})_2$ (*Oxyosmiumdiammonium chloride*) The formula is sometimes written $\text{OsO}_2\text{Cl}_2 \cdot 4\text{NH}_4$, and the salt is called *ammoniated osmium oxychloride*, the formula is also written $\text{OsO}_2 \cdot 4\text{NH}_4\text{Cl}$, and the salt is called *osmyle ditetramine chloride* (Gibbs, *Am J*, 233), Fremy (*A Ch* [3] 12, 522) wrote the formula $\text{OsO}_2(\text{NH}_4)_2 \cdot 2\text{NH}_4\text{Cl}$, calling the salt *osmiumdiammonium chloride*, Claus and Jacobi (*J pr* 79, 28, 85, 142, 90, 65) gave the formula $\text{Os}(\text{NH}_4)_2\text{Cl}_2 \cdot 2\text{H}_2\text{O}$, and the name *osmio diammonio chloride*. The investigations of Gibbs (*Am J*, 233) confirmed the empirical formula $\text{OsO}_2\text{Cl}_2 \cdot 4\text{NH}_4$. Obtained as a yellow crystalline pp by adding NH_4Cl to $\text{K}_2\text{OsO}_4\text{Aq}$ (Fremy, *A Ch* [3] 12, 522) ($7\text{K}_2\text{OsO}_4\text{Aq} + 4\text{NH}_4\text{Cl} = \text{OsO}_2(\text{NH}_4, \text{NH}_4\text{Cl})_2 + 2\text{KClAq} + 2\text{H}_2\text{O}$). Slightly sol cold water, more sol hot water, insol NH_4ClAq . May be crystallised from water containing HCl , ppd from solution by conc HClAq . Aqueous solution quickly decomposes, giving off OsO_4 , on evaporation a brown solid is obtained, which behaves like $\text{OsO}(\text{NH}_4, \text{OH})_2$, the solution gives a fine violet colour with $\text{K}_2\text{FeCy}_6\text{Aq}$ (Gibbs, *l c*). The compound is decomposed by heat, giving a residue of Os.

The *chloroplatinate* is obtained, as orange-yellow crystals, slightly sol cold water, by boiling $\text{OsO}_2(\text{NH}_4, \text{NH}_4\text{Cl})_2$ with PtCl_4Aq (Gibbs, *l c*).

OSMIL TETRAMMONIUM SULPHATE, NITRATE, &c. The *sulphate* is obtained by pouring $\text{K}_2\text{OsO}_4\text{Aq}$ into conc cold $(\text{NH}_4)_2\text{SO}_4\text{Aq}$, small orange-yellow crystals, easily sol hot water (Gibbs, *l c*). The *nitrate* is obtained by a similar reaction, and the *oxalate* also, using cold conc $\text{NH}_4\text{NO}_3\text{Aq}$

and $(\text{NH}_4)_2\text{C}_2\text{O}_4\text{Aq}$ instead of $(\text{NH}_4)_2\text{SO}_4\text{Aq}$ (Gibbs, *l.c.*).

Osmium, chlorides of. Two chlorides of Os have been prepared, OsCl_2 and OsCl_3 . OsCl_2 is known in solution, no chloride has yet been gasified, these are the only haloid compounds of Os known at present, except OsCy , be classed as a haloid compound. OsCl_2 and OsCl_3 combine with alkali chlorides to form $\text{OsCl}_2 \cdot 3\text{MCl}$ and $\text{OsCl}_3 \cdot 2\text{MCl}$ respectively. OsCl_2 also probably forms double salts, but they have not been isolated satisfactorily. OsCy , forms an acid, H_2OsCy_2 , from which various salts (osmocyanides) are obtained. OsCl_2 and OsCl_3 are formed by direct union of Cl with Os.

Osmous chloride OsCl_2 (*Osmium dichloride Osmochloride*) Obtained by heating finely divided Os in a long tube in a stream of dry Cl. Two sublimates are obtained, the less volatile (green) is OsCl_2 , the more volatile (red) is OsCl_3 . As thus prepared OsCl_2 forms dark green deliquescent needles, which probably contain H_2O , by heating again in perfectly dry Cl, OsCl_2 is obtained free from moisture as a nearly black sublimate (Claus a Jacobi, *J pr* 90, 65). An indigo-blue solution of OsCl_2 may be prepared by dissolving $\text{OsO}_2 \cdot x\text{H}_2\text{O}$ in HClAq , the solution soon becomes violet, then dark red, and then yellow from formation of OsCl_2 and OsCl_3 , the yellow solution again becomes blue by the action of reducers (C a J, *l.c.*) OsCl_2 dissolves in water, forming a green liquid, which is decolourised and decomposed on dilution with formation of Os, OsO_4Aq , and HClAq . Solutions of OsCl_2 in alkali chloride solutions are not decomposed by dilution, double salts are probably present (C a J). Berzelius said that double salts of OsCl_2 are formed by the action of alcohol on the salts M_2OsCl_4 and M_2OsCl_6 (P 13, 435, 527). OsCl_2 is sol alcohol and ether. The aqueous solution conducts electricity, but the alcoholic and ethereal solutions do not (*v* Hampe, Z 11, 1549, 12, 23).

Osmoso-osmic chloride OsCl_3 (*Osmium trichloride Osmochloride Osmium sesquichloride*) This chloride is only known in solution, and in combination with alkali chlorides. The brown to purple solution obtained by the action of air on $\text{OsO}_2 \cdot x\text{H}_2\text{O}$ in HClAq probably contains OsCl_3 (C a J). By treating a solution of OsO_3 in HClAq with Hg, and evaporating *in vacuo* over conc H_2SO_4 , Berzelius (P 13, 435, 527) obtained a purple, varnish-like substance, which perhaps was OsCl_3 .

Double Salts.—Ammonium-osmium trichloride (*Ammonium chlorosmiate*) $2(\text{OsCl}_3 \cdot 2\text{NH}_4\text{Cl}) \cdot 3\text{H}_2\text{O}$ Reddish-brown crystals, obtained by passing H_2S into OsO_3 in much HClAq till the solution is red, adding NH_4Cl , and evaporating (Claus, *J pr* 79, 28). The salt dissolves in water with a dark purple colour, which is unchanged by addition of alkali, but reduction begins on warming. On heating, Os and NH_4Cl are formed (Berzelius, *l.c.*)

Potassium-osmium trichloride (*Potassium chlorosmiate*) $2(\text{OsCl}_3 \cdot 3\text{KCl}) \cdot 6\text{H}_2\text{O}$. Formed by strongly heating in Cl a mixture of equal parts KCl and powdered Os, dissolving in water, crystallising out K_2OsCl_6 , and evaporating the mother-liquor, also by dissolving KOH in conc OsO_4Aq , adding NH_4Aq , and when the solution

is yellow and before K_2OsO_4 separates, saturating with HClAq and evaporating to dryness on a water-bath, when the salt is obtained on the bottom of the basin, while KCl and NH_4Cl are deposited on the upper parts. Forms dark brownish red crystals, easily sol water, with deep cherry-red colour, easily sol alcohol, insol ether. The salt loses $3\text{H}_2\text{O}$ in air, and the rest at 150° – 180° . The aqueous solution easily decomposes with separation of $\text{Os}_2\text{O}_7\text{Cl}_2$.

Osmic chloride OsCl_4 (*Osmium tetrachloride Osmichloride*) Obtained, as a dark red powder, by heating finely powdered Os in dry Cl, OsCl_2 is formed at the same time. OsCl_4 is the more volatile of the two chlorides. In presence of moisture, becomes cinnabar red, and then forms yellow needles ($? \text{OsCl}_4 \cdot x\text{H}_2\text{O}$). Dissolves in a little water to form a yellow solution, which is decomposed and decolourised on dilution (Berzelius, *l.c.*) OsCl_4 is a non conductor of electricity (Hampe, Z 11, 1549, 12, 23). $\text{OsO}_2 \cdot 2\text{H}_2\text{O}$ dissolves in conc HClAq , forming a dark greenish brown solution, which probably contains OsCl_4 .

Double salts—Potassium osmium tetrachloride (*Potassium chlorosmate*) K_2OsCl_6 ($= \text{OsCl}_4 \cdot 2\text{KCl}$) Obtained by heating an intimate mixture of equal parts KCl and finely powdered Os in dry Cl to low redness, removing excess of KCl by a little water, dissolving the residue in hot water, and allowing to crystallise (Berzelius, *l.c.*) Claus a Jacobi (*l.c.*) obtained the salt by adding KCl to OsO_3 in HClAq , then adding alcohol and evaporating. Dark-brown lustrous octahedra, cinnabar red when powdered. Sol water, forming a yellow solution, from which alcohol ppts the salt, incompletely, as a red crystalline powder. Not decomposed at low redness, but at a higher temperature Os and KCl are formed. OsO_3 is produced by distilling HNO_3 over K_2OsCl_6 (Berzelius, *l.c.*) Is not acted on by SO_2Aq at ordinary temperatures (K_2IrCl_6 is reduced). KOHAq does not decompose a cold solution of K_2OsCl_6 , but on warming the solution becomes blue, and then suddenly ppts black $\text{OsO}_2 \cdot 2\text{H}_2\text{O}$ (Claus, *l.c.*, other reactions of $\text{K}_2\text{OsCl}_6\text{Aq}$ are given).

Ammonium chlorosmate, silver chlorosmate, and sodium chlorosmate are described by Claus a Jacobi (*J pr* 79, 28, 85, 142, 90, 65, A 63, 855).

Supposed compound of Osmium hexachloride Berzelius (P 13, 435, 527, 15, 203) obtained a brown salt, which he supposed to be a compound of NH_4Cl and OsCl_6 , by saturating OsO_3 with NH_4Aq , after a time adding excess of HClAq , digesting for some days with Hg, filtering, and evaporating. According to Claus (*l.c.*), the salt is $2(\text{OsCl}_6 \cdot 2\text{NH}_4\text{Cl}) \cdot 3\text{H}_2\text{O}$ (*v supra*, *Ammonium-osmium trichloride*).

Osmium, cyanide of, OsCy_2 , and osmocyanhydric acid and its salts, H_2OsCy_2 , *v* CYANIDES, vol II, p 343

Osmium, hydrated oxides or hydroxides of, *Os*, Osmium, oxides and hydrated oxides of, p 645

Osmium, nitride of The brass-yellow substance, obtained by passing H over the pp. produced by adding HNO_3 to $\text{K}_2\text{OsO}_4\text{Aq}$, is probably a nitride of Os (Claus, P. 65, 202).

Osmium, nitrogen-containing acid of, and

its salts $\text{H}_2\text{N}_2\text{Os}_2\text{O}_4\text{Aq}$ or $?\text{H}_2\text{N}_2\text{Os}_2\text{O}_4\text{Aq}$ This acid is generally known as *osmic acid*. An aqueous solution of this acid is obtained by decomposing the Ba salt by the equivalent quantity of $\text{H}_2\text{SO}_4\text{Aq}$, or by treating the moist Ag salt with HClAq , and filtering. The clear yellow solution remains unchanged for some days if dilute, but if it be concentrated, OsO_4 and a black pp containing Os are formed, and gas is evolved. The aqueous solution of the acid evolves CO_2 from carbonates, and decomposes KCl , Zn dissolves in it, with evolution of gas and partial decomposition of the acid. The solution is decomposed by acids on warming, with production of OsO_4 (Fritzsche a Struve, *J pr* 41, 97). F a S gave the formula $\text{H}_2\text{N}_2\text{Os}_2\text{O}_4$, this was upheld by Claus (*J pr* 90, 65). Gerhardt (*J Ph* [3] 12, 304) gave the formula $\text{H}_2\text{N}_2\text{Os}_2\text{O}_4$. The K salt is obtained by the action of NH_4Aq on OsO_4 in excess of KOHAg , the reaction is expressed thus by F a S $2\text{OsO}_4 + 2\text{NH}_3 + \text{Aq} = \text{H}_2\text{N}_2\text{Os}_2\text{O}_4\text{Aq} + \text{O} + 2\text{H}_2\text{O}$, Claus gives the equation $6\text{OsO}_4 + 8\text{NH}_3 + \text{Aq} = 3\text{H}_2\text{N}_2\text{Os}_2\text{O}_4\text{Aq} + \text{N}_2 + 9\text{H}_2\text{O}$. According to F a S no gas is evolved during the reaction. If this is correct neither of the equations can express the change which occurs. The salts of osmic acid explode when heated, some explode when struck.

POTASSIUM OSMIATE $\text{K}_2\text{N}_2\text{Os}_2\text{O}_4$ or $?\text{K}_2\text{N}_2\text{Os}_2\text{O}_4$. Prepared by dissolving solid OsO_4 in conc KOHAg , to which conc NH_4Aq has been added. The solution is clear yellow, and deposits yellow crystals of the salt. Claus a Jacobi (*lc*) dissolve KOH in very dilute OsO_4Aq , add $\frac{1}{2}$ vol NH_4Aq , and evaporate rapidly until the salt begins to crystallise out, after which crystallisation is allowed to proceed. Citron yellow, tetragonal crystals. Sol warm water, also in alcohol, sl sol cold water, insol ether. Explodes at c 180° . Decomposed by conc HClAq , with evolution of Cl and formation of two kinds of red crystals (*v* F a S, *lc*).

SILVER OSMIATE $\text{Ag}_2\text{N}_2\text{Os}_2\text{O}_4$ or $?\text{Ag}_2\text{N}_2\text{Os}_2\text{O}_4$. A citron yellow crystalline powder, sl sol water or dilute HNO_3Aq , sol NH_4Aq , forming therewith a crystalline compound, blackens in light, explodes at 80° , also when struck by a hammer. Prepared by dissolving OsO_4 in an ammoniacal solution of a salt of Ag, and adding excess of HNO_3Aq , or by adding excess of HNO_3Aq , followed by solution of a salt of Ag, to OsO_4 , dissolved in NH_4Aq .

Ammonium, barium, mercurous, sodium, and ammoniated zinc, osmiates have been prepared (F a S, *lc*).

Osmium, oxides and hydrated oxides of Four oxides of Os are known, OsO , Os_2O_3 , Os_2O_4 , and OsO_4 . Salts in which OsO_4 acts as the negative radicle have been prepared (*v infra*, *Osmium, salts of oxyacids of*). A hydrate of OsO has been obtained, but not pure, it rapidly absorbs O , the hydrate $\text{OsO}_2 \cdot 3\text{H}_2\text{O}$ is known, as are also the hydrates $\text{OsO}_3 \cdot \text{H}_2\text{O}$ and $\text{OsO}_2 \cdot 2\text{H}_2\text{O}$. No oxy-salts have been obtained directly from the oxides of Os, a few oxy salts corresponding with OsO are known. OsO_4 acts as a weak acidic anhydride, this oxide is easily reduced to OsO_3 , or to Os . OsO_4 is the only oxide prepared by the direct union of Os with O . Chlorides corresponding with OsO and OsO_2 ,

have been obtained, and the chloride OsCl_4 , which corresponds with OsO_4 , probably exists in solution (*v Osmium, chlorides of*, p 644).

Osmous oxide OsO (*Osmium monoxide*). A greyish black solid, insol acids, obtained by heating in a stream of CO_2 , a mixture of Na_2CO_3 and OsSO_4 , or $\text{OsO}_2 \cdot 2\text{K}_2\text{SO}_4 \cdot 2\text{KH}_2\text{SO}_4 \cdot 4\text{H}_2\text{O}$, and washing the residue with water (Claus a Jacobi, *J pr* 85, 142, 90, 65). The salt OsSO_4 is formed by evaporating OsO_4Aq with excess of SO_2 , the salt $\text{OsSO}_4 \cdot 2\text{K}_2\text{SO}_4 \cdot 2\text{KH}_2\text{SO}_4 \cdot 4\text{H}_2\text{O}$ is formed by heating $\text{K}_2\text{OsCl}_6\text{Aq}$ with K_2SO_4 , and washing with water the powder which separates. *Hydrated osmous oxide*, $\text{OsO} \cdot x\text{H}_2\text{O}$, was obtained by C a J by the action of warm, very conc KOHAg on OsSO_4 , in absence of air, it forms a black blue pp which very rapidly absorbs O from the air. Berzelius obtained a greenish black pp by boiling OsCl_4 in KClAq with KOHAg , this pp was supposed to be $\text{OsO} \cdot x\text{H}_2\text{O}$, but C a J say it cannot be washed free from alkali. A few salts are known corresponding with OsO , e.g. OsSO_4 , but none is obtained directly from the oxide.

Osmo osmic oxide Os_2O_3 (*Osmium sesquioxide*). A black powder, insol acids. Obtained by gently heating, in a stream of CO_2 , a mixture of $\text{K}_2\text{OsCl}_6 \cdot 3\text{H}_2\text{O}$ and Na_2CO_3 , and washing with water (C a J, *lc*). Deville a Debray (*C R* 82, 1076) say that this oxide is often formed, as copper red crusts, in the preparation of Os, by passing OsO_4 vapour mixed with N through a hot tube lined with C , the lining of C being formed by passing C_2H_2 vapour through the tube heated to redness. Berzelius supposed that a compound of Os_2O_3 with NH_3 was formed by the action of NH_4Aq on OsO_4Aq , but the pp thus formed was shown by C a J to be $\text{OsO}(\text{NH}_4\text{OH})$, (*v Osmosyl diammonium hydroxide*, p 645). No oxy-salts are known corresponding with Os_2O_3 , a solution of OsCl_4 has been prepared.

Hydrated osmo-osmic oxide $\text{Os}_2\text{O}_3 \cdot 3\text{H}_2\text{O}$, a brown red pp, somewhat sol. cold KOHAg , sol acids even after drying, obtained by adding KOHAg to $\text{K}_2\text{OsCl}_6\text{Aq}$ (C a J, *lc*).

Osmic oxide OsO_2 (*Osmium dioxide*). Obtained by Berzelius (*P* 13, 435, 527, 15, 208) by heating a mixture of K_2OsCl_6 and Na_2CO_3 in CO_2 , washing the residue with water, and then with HClAq , as thus prepared OsO_2 forms a greyish-black powder. Claus a Jacobi (*lc*) obtained OsO_2 in the form of a copper red, metallic solid, by strongly heating $\text{OsO}_2 \cdot 2\text{H}_2\text{O}$ in a covered crucible (OsO_3 , H , and H_2O are also formed). $\text{OsO}_2 \cdot 2\text{H}_2\text{O}$ is obtained by the action of acids on $\text{K}_2\text{OsO}_4\text{Aq}$. OsO_2 may be heated to redness in absence of air (Berzelius), but Claus says that OsO_2 and Os are formed. Heated in air, or acted on by HNO_3 and HClAq , OsO_3 is produced. Reduced by H at ordinary temperatures. Detonates when heated with combustible bodies. A sulphate corresponding with OsO_2 is said to be prepared by the action of cold HNO_3Aq on OsS_2 , but little is known of its properties. The corresponding chloride, OsCl_2 , is known.

Hydrated osmic oxide $\text{OsO}_2 \cdot 2\text{H}_2\text{O}$, a black pp obtained by adding very dilute $\text{H}_2\text{SO}_4\text{Aq}$ to $\text{K}_2\text{OsO}_4\text{Aq}$, OsO_2 being formed at the same time (C a J, *lc*). The moist hydrate dissolves in HClAq , HNO_3Aq , or $\text{H}_2\text{SO}_4\text{Aq}$, oxidation begins

very quickly, with change of colour from purple to yellow-brown, green, and brownish-yellow on heating. From the solution in HClAq , Zn ppts Os (Wöhler, *A* 140, 256). By heating $\text{OsO}_2 \cdot 2\text{H}_2\text{O}$ to 200° in dry N, Fremy obtained the hydrate $\text{OsO}_2 \cdot \text{H}_2\text{O}$ (*A Ch* [3] 12, 515).

OSMIUM TETROXIDE OsO_4 , (*Perosmic anhydride*). Often called *perosmic acid*, and formerly generally known as *osmic acid*. Mol w 254.84. V D 128.5 at c 280° (Deville & Debray, *O R* 82, 1076).

Formation—1 By heating Os in air or O—2 By heating Os, or any of the lower oxides, with HNO_3 or *aqua regia*—3 By the action of dilute acids on $\text{K}_2\text{OsO}_4\text{Aq}$, $\text{OsO}_2 \cdot 2\text{H}_2\text{O}$ being formed at the same time—4 According to Claus (*J pr* 79, 28) by the action of Os on steam, H being evolved.

Preparation—1 Finely powdered Os is gently warmed in a stream of dry O, the Os is placed in one bulb of a two bulb tube, and the OsO_4 condenses in the second bulb—2 Three parts osmium are fused with 1 part KNO_3 , the cold mass is dissolved in water, the solution is neutralised by $\text{H}_2\text{SO}_4\text{Aq}$, and OsO_4 is distilled off into a cold receiver.

Properties—Long, colourless, translucent, monoclinic, needles (Mallet, *Am S* [2] 29, 49). Melts considerably under 100° , and boils a few degrees above its melting point. OsO_4 is very volatile, the vapour smells abominably, and is extremely poisonous, attacking the eyes and lungs, Deville nearly lost his eyesight while working with OsO_4 (*v D* a *D*, *A Ch* [3] 56, 400). Claus says the best antidote is H_2S gas. The V D was determined by D a D at 246° – 285° , it agrees with the formula OsO_4 . OsO_4 may be vapourised in H, if the mixed vapours are passed through a hot tube, Os is deposited. OsO_4 is slowly dissolved by water, the solution does not react acid, sol alcohol or ether, solutions are reduced with ppn of Os.

Reactions—1 Reduced to Os by heating with hydrogen, also by heating in a gas flame, an aqueous solution of OsO_4 is readily reduced by most metals, the solution is also reduced by phosphorus, ferrous sulphate, stannous chloride, sulphurous acid (a blue liquid is produced by SO_2Aq , perhaps containing an oxide which has not been isolated), and by many carbon compounds (e.g. $\text{H.CO}_2\text{H}$, tannin, sugar, alcohol, &c.) (*v* Tennant, *T* 1804 411, Berzelius, *P* 13, 435, 327, 15, 208, Döbereiner & Weiss, *A* 14, 17, 251, Butlerow, *A* 84, 278, Claus & Jacobi, *J pr* 90, 65)—2 A large excess of ammonia solution evolves N, and ppts $\text{OsO}(\text{NH}_4)\text{OH}$, (*v* *Osmosyl diammonium hydroxide*, p 643)—3 OsO_4 dissolves in potash, forming a red yellow liquid which probably contains K perosmate (*q v* under *Osmium, salts of oxyacids of, infra*)—4 Hydrogen sulphide ppts an oxy-sulphide (*q v*) from OsO_4Aq , on saturating OsO_4Aq with H_2S , and adding an acid, OsS_2 is ppd (*v* *Osmium, sulphides of*, p 647)—5 Hydrochloric acid does not react with OsO_4 , but on addition of KOH Aq , K_2OsO_4 is produced. OsO_4 is a very weak acidic anhydride, an aqueous solution does not affect the colour of litmus, nor does it decompose carbonates; KOH Aq probably forms K perosmate.

Osmium, oxyacids of No oxyacid of Os has been isolated, salts of the hypothetical H_2OsO_4 (osmates) are known, and some of the reactions of OsO_4 indicate the existence of unstable salts derived from the oxide OsO_4 (*v infra*, *Osmium, salts of oxyacids of*).

Osmium, oxychloride of The black pp obtained by boiling dilute $\text{K}_2\text{OsO}_4\text{Aq}$ is probably an oxychloride (Claus & Jacobi, *J pr* 90, 65).

Osmium, oxysulphides of The pp obtained by passing H_2S into OsO_4Aq is said by von Meyer to have the composition $\text{Os}_2\text{O}_3\text{S}$, $3\text{H}_2\text{O}$, and to be changed to $2\text{OsO}_3\text{S}$, $3\text{H}_2\text{O}$ by the action of air (*J pr* [2] 16, 77). Dried at 180° – 200° , OsO_3S is obtained (von M., *lc*).

Osmium, phosphide of A phosphide of Os is said to be formed by heating Os in P vapour, it appears black if prepared at a moderate temperature, but white and metal like if produced by heating strongly, it burns in air to OsO_4 , and osmious phosphate (Berzelius, *P* 13, 435, 527, 15, 208).

Osmium, salts of Very few salts have been obtained by replacing the H of oxyacids by Os. Osmium sulphate, OsSO_4 , is formed by evaporating OsO_4Aq with SO_4Aq , a complex salt $\text{OsSO}_4 \cdot 2\text{K}_2\text{SO}_4 \cdot 2\text{KHSO}_4 \cdot 4\text{H}_2\text{O}$ is obtained by heating K_2OsO_4 with $\text{K}_2\text{SO}_4\text{Aq}$, osmious phosphate and sulphate are also said to exist, but they have not been analysed (Berzelius). An osmic sulphate is stated to be produced by the action of cold HNO_3Aq on OsS_2 (Berzelius). No salt of Os has been formed by the direct reaction of Os or any of its oxides with an oxyacid (*v* SULPHITES, SULPHATES).

Osmium, salts of oxyacids of Some salts of the hypothetical *osmic acid*, H_2OsO_4 , have been isolated, and there are indications of the existence of alkali perosmates, i.e. salts derivable from OsO_4 .

OSMATES M^1OsO_4 (formerly called *Osmites*). Neither the acid H_2OsO_4 , nor the anhydride OsO_4 , has been isolated (*cf* Mallet, *Am S* [2] 29, 49). Alkali osmates are obtained by the reducing action of alcohol on OsO_4 , in KOH Aq , a Ba salt is formed by the prolonged reaction of warm BaO Aq on OsO_4 in a closed vessel, osmates of Ca, Pb, and Sr are obtained as insoluble pps from the alkali salts.

Potassium osmate $\text{K}_2\text{OsO}_4 \cdot 2\text{H}_2\text{O}$. A violet coloured, crystalline pp of this composition is formed by adding alcohol to OsO_4 , dissolved in KOH Aq , the solution is red, then becomes colourless, and deposits the salt. Fremy obtained the salt in large octahedra by adding KNO_3Aq to OsO_4 in large excess of KOH Aq (*J pr* 33, 411). According to Claus (*J pr* 34, 178, 424) $\text{K}_2\text{OsO}_4 \cdot 2\text{H}_2\text{O}$ is obtained by warming OsO_4 in an excess of KOH Aq , this reaction is similar to the reduction of KMnO_4Aq to $\text{K}_2\text{MnO}_4\text{Aq}$ by KOH . Violet red octahedra. Sol water, insol alcohol or ether, slightly sol conc salt solutions (Gibbs, *Am S* [2] 31, 70). An aqueous solution decomposes rapidly on warming to K perosmate and $\text{OsO}_2 \cdot 2\text{H}_2\text{O}$. $\text{H}_2\text{SO}_4\text{Aq}$ forms OsO_4 from $\text{K}_2\text{OsO}_4\text{Aq}$, and separates $\text{OsO}_2 \cdot 2\text{H}_2\text{O}$ (Claus & Jacobi, *lc*). HNO_3Aq produces a similar change (Fremy, *lc*). SO_4Aq produces a blue solution. $\text{NH}_4\text{Cl Aq}$ forms $\text{OsO}_4(\text{NH}_4)_2$, and $(\text{NH}_4)_2\text{SO}_4\text{Aq}$ produces $\text{OsO}_4(\text{NH}_4)_2\text{SO}_4$ (*v* *Osmyl-tetrammonium*).

chloride and sulphate, p 643) NH_4Aq is said to form $\text{OsO}_2(\text{NH}_4)_2$ (Freymy, *lc*)

Barium osmate $\text{BaOsO}_4 \cdot \text{H}_2\text{O}$ (Claus, *J pr* 4, 170, 224), **Calcium, lead, sodium, and strontium osmates** have been obtained (Freymy, *J pr* 33, 411)

PEROSMATES (formerly called *osmates*) These salts have been prepared pure Freymy (*J pr* 33, 409) says a perosmate of K is formed by dissolving OsO_4 in large excess of KOH Aq , and the salt is decomposed on dilution, the solution in excess of KOH Aq is colourless, but absorbs O and becomes brown, on boiling, K_2OsO_8 is formed, and an oxide of Os containing more O than OsO_4 is volatilised. On the other hand, Claus (*J pr* 85, 142, 90, 65) says that the greater part of the OsO_4 can be distilled off from a solution of this oxide in excess of KOH Aq , but that some decomposes to $\text{K}_2\text{OsO}_8 \text{ Aq}$ and O. Claus also says that OsO_4 distils off, at 60° , when Cl is passed into $\text{OsO}_4 \cdot \text{H}_2\text{O}$ suspended in a large excess of KOH Aq . From a conc solution of OsO_4 in KOH Aq , $\text{HNO}_3 \text{ Aq}$ ppt OsO_4 mixed with $\text{OsO}_3 \cdot \text{H}_2\text{O}$, according to Wöhler (*A* 140, 256). Wöhler (*lc*) melted Os with KOH and KNO_3 , and obtained a black solid which formed a deep red solution in water. This solution was supposed by Wöhler to contain K perosmate, only a very little OsO_4 was obtained by distilling, and the residual liquid deposited K_2OsO_8 on cooling. The mother liquor from the K_2OsO_8 was yellow, on treatment with CO_2 it became colourless, and then violet, and deposited OsO_4 and a greyish violet pp.

Osmium, sulphides of A sulphide of Os is formed by heating Os with S (Beizelus, *P* 13, 435-527, 15, 208)

Osmic sulphide OsS (*Osmium disulphide*) A dark brownish yellow solid, slightly soluble in water. Produced, according to Freymy (*J pr* 33, 409), by passing H_2S into $\text{K}_2\text{OsO}_8 \text{ Aq}$, according to Claus (*J pr* 79, 28), also by passing H_2S into $\text{K OsCl}_4 \text{ Aq}$.

Osmium tetrakisulphide $\text{OsS}_4 \cdot \text{H}_2\text{O}$ Obtained by saturating OsO_4 in HCl Aq with H_2S , on attempting to dry the pp it is partly oxidised, above 100° , the pp burns to OsO_4 and SO_2 , easily oxidised by $\text{HNO}_3 \text{ Aq}$, heated in absence of air, $\text{OsS}_4 \cdot \text{OsS}_2$ is said to be produced (Berzelius, Claus). The pp obtained by passing H_2S into $\text{OsO}_4 \text{ Aq}$ is an *oxysulphide* (q v)

M M P M

OSMOCYANIDES and *Osmocyanhydric acid* v CYANIDES, vol II p 343

OSMOSE When two liquids are separated by the intervention of a porous diaphragm, a flow of liquid takes place from one side of the septum to the other, or sometimes an unequal flow of the two liquids in opposite directions, so that the quantity of liquid increases on one side of the septum and diminishes on the other. This phenomenon was originally designated by the correlative terms *Endosmose* and *Exosmose*, but it is better expressed by the shorter word *Osmose* (from *ὄσμος*, impulsion), which includes the two former. For the application of osmotic methods to chemical questions v. MOLECULAR WEIGHTS, p 417, also PHYSICAL METHODS; section *Electrical methods* M M P M

OSOTRIAZOLE $\text{C}_4\text{H}_4\text{N}_4$, $\text{NH} \angle \text{N CH}$

[22 5°] (204°) at 715 mm Formed by heating its carboxylic acid Sol water and alcohol, insol ligroin (Pechmann, *A* 262, 320) — B' HCl — B' HgCl glittering needles

Benzoyl derivative $\text{NBz} \angle \text{N CH}$ [100°]

OSOTRIAZOLE CARBOXYLIC ACID

$\text{NH} \angle \text{N CH}$ [211°] Formed by the action of KMnO_4 on amido phenyl osotriazole carboxylic acid, which is got by reducing the nitro-acid formed by nitrating the acid produced by the action of soda on the phenyl-hydrazide of di-nitroso acetone (Pechmann, *A* 262, 315) Crystalline powder, v sol boiling water — CaA' , 2aq

OSOTRIAZONES and OSOTETRAZONES v HYDRAZONES

OSSEIN v PROTEIDS, Appendix C

OSTRUTHIN $\text{C}_{14}\text{H}_{11}\text{O}_2$ (Görup-Besanez, *A* 183, 321), $\text{C}_{14}\text{H}_{11}\text{O}_2$ (Jassoy, *C C* 1890 [1] 766) [115°] Occurs in the roots of *Imperatoria Ostruthium*. Trichinic crystals, v sl sol boiling water, v sol alcohol and ether. Its alcoholic solution exhibits blue fluorescence. Forms a crystalline compound with dry HCl

Acetyl derivative [78°] Plates

OTOBITE $\text{C}_8\text{H}_8\text{O}_2$ [133°] Occurs in the fruit of *Myristica otoba* (Uruciochea, *A* 91, 370). Pearly prisms (from ether), insol water

OUABAIN $\text{C}_{29}\text{H}_{44}\text{O}_{12}$ [α 185°] δ 65 at 11° , S (alcohol) 3.75 at 11° [α]_D = -33° . A poisonous glucoside present in Ouabain root, used as arrow poison, and in the seeds of the glabrous *Strophantus* of Gabon (Arnau, *C R* 106, 1011, 107, 348, 1162, *Bl* [δ] 1, 10). Rectangular plates (containing 7aq) v sl sol water. Its solution is ppd by tannin. It yields a sugar and a resin on boiling with acids — BaA' deliquescent pp

OXALACETIC ACID $\text{C}_4\text{H}_4\text{O}_6$ *Oxalozyl acetic acid*

Oxim $\text{CO}_2\text{H C}(\text{NOH})\text{CH}_2\text{CO}_2\text{H}$ [125°] Formed by the action of alcoholic NaOH on the ether $\text{CO}_2\text{Et C}(\text{NOH})\text{CH}_2\text{CO}_2\text{H}$ [107°], which is obtained by the action of water on the dihydride of di-nitroso di-oxo terephthalic ether (Cramer, *B* 24, 1204). An isomeric oxim $\text{CO}_2\text{Et C}(\text{NOH})\text{CH}_2\text{CO}_2\text{H}$ [54°] is formed from oxalacetic acid and hydroxylamine. It yields $\text{CO}_2\text{H C}(\text{NOH})\text{CH}_2\text{CO}_2\text{H}$ [88°], whence AcCl forms an anhydride $\text{C}_4\text{H}_4\text{O}_6$ [105°] yielding $\text{C}_4\text{H}_4\text{AcO}$ [105°]

Phenylhydrazide of the methyl ether $\text{CO}_2\text{Me CH}_2\text{C}(\text{NHPH})\text{CO}_2\text{Me}$ [118°] Mol. w 257 (obs.), (calc 250). Formed from di-methyl acetylene dicarboxylate and phenyl-hydrazine (Buchner, *B* 22, 2930). Plates. Yields oxo-phenyl pyrazole carboxylic acid on saponification

Mono ethyl ether $\text{CO}_2\text{Et CH}_2\text{CO}_2\text{H}$ [97°] Obtained from the di-ethyl ether and cold dilute KOH (Wislicenus, *B* 19, 8226, *A* 246, 828). Needles, v sol water. Its aqueous solution is coloured deep red by FeCl_3

Anilide of the ethyl ether $\text{CO}_2\text{Et CO CH}_2\text{CO NHPh}$ [88°] Formed from oxalic ether, acetanilide, and NaOEt (Wislicenus, *B* 24, 1250). Needles, v sol alcohol

p-Toluide of the ethyl ether [135°]

Ethyl anilide of the ethyl ether $\text{CO}_2\text{Et CO CH}_2\text{CO NEtPh}$ [69°] Formed from

NaOEt, oxalic ether, and NEtAcPh (Wislicenus a Ba^{++} er, *B* 24, 1254) — $\text{Cu}(\text{C}_2\text{H}_3\text{NO}_2)_2$ [189°]

Di-ethyl ether $\text{CO}_2\text{EtCH}_2\text{CO CO}_2\text{Et}$ 182° at 24 mm) SG $\frac{22.5}{4}$ 1159 Formed by the action of sodium on an ethereal solution of oxalic and acetic ethers (Wislicenus, *B* 19, 3225, 20, 3392, *A* 246, 315) Oil FeCl_3 colours its alcoholic solution red

Reactions — 1 Decomposed by dilute acids into acetic and oxalic acids — 2 Dilute (10 p c) H_2SO_4 yields pyruvic acid — 3 Sodium forms the compound $\text{CO}_2\text{EtCHNa CO CO}_2\text{Et}$, which reacts with alkyl iodides, forming alkyl oxal-acetic ethers $\text{CO}_2\text{EtCHX CO CO}_2\text{Et}$ The copper compound $\text{Cu}(\text{C}_2\text{H}_3\text{NO}_2)_2$ crystallises from alcohol in green needles [163°] — 4 *Phenyl hydrazine* reacts forming a phenyl-hydrazide $\text{CO}_2\text{EtCH}_2\text{C}(\text{N}(\text{HPh})\text{CO}_2\text{Et})$ [78°], crystallising in plates, and forming $\text{CO}_2\text{EtC} \begin{smallmatrix} \text{CH}_2\text{CO} \\ \text{N NPh} \end{smallmatrix}$

[182°] on heating — 5 *Aniline* at 0° forms $\text{CO}_2\text{EtCH}_2\text{C}(\text{NPh})\text{CO}_2\text{Et}$ (c 200° at 10 mm), which gives no colour with alcoholic FeCl_3 . The condensation products include $\text{C}_2\text{H}_3\text{NO}_2$, [108°] and $\text{C}_2\text{H}_3\text{NO}_2$, [218°] (Wislicenus a Spiro, *B* 22, 3349) — 6 Aqueous KOAc yields $\text{CO}_2\text{EtCH}(\text{CO}_2\text{H})\text{CH}(\text{CO}_2\text{Et})\text{CO CO}_2\text{Et}$, which forms the crystalline salts BaA'_2 , 2aq and PbA'_2 (Claisen a Hori, *B* 24, 124) — 7 Its oxim is reduced by sodium amalgam to mono ethyl aspartates $\text{CO}_2\text{HCH}(\text{NH}_2)\text{CH}_2\text{CO}_2\text{Et}$ [201°] and $\text{CO}_2\text{HCH}_2\text{CH}(\text{NH}_2)\text{CO}_2\text{Et}$ [165°] (Piutti, *C* 1888, 68, 1459) — 8 *Benzamidine hydrochloride* and dilute (10 p c) aqueous caustic soda yields $\text{CO}_2\text{EtCO CH}_2\text{CO NH CPh NH}$ [180°] (Finner, *B* 22, 1628, 2615)

Reference — AMIDO-OXALOXYL-ACETIC ACID

OXALAMIDO-BENZOIC ACID v CARBOXY-

PHENYL OXAMIC ACID

OXALANTIN $\text{C}_2\text{H}_3\text{N}_2\text{O}_4$ Formed by reducing alloxanic acid with zinc and HClAq (Lumprecht, *A* 111, 133, cf Schlieper, *A* 56, 2) Small crystals, sl sol water Decomposed by alkalis into NH_3 and oxaluric acid

OXALBUTYRIC ETHER

$\text{CO}_2\text{EtCO CHEtCO}_2\text{Et}$ (188° at 20 mm) Formed from oxalic ether, butyric ether, and NaOEt (Arnold, *A* 246, 337) Oil Gives a red colour with FeCl_3

OXALETHYLAMYLINE v ETHYL BUTYL-GLYOXALINE

OXAL-ETHYL-ETHYLENE v. METHYL-ETHYL-GLYOXALINE

OXALETHYLENE v METHYL ETHYL GLYOXALINE

OXALETHYLENANTHYLINE v. ETHYL-HEXYL-GLYOXALINE

OXAL ETHYL PROPYLENE v DI-ETHYL-GLYOXALINE

OXALIC ACID $\text{H}_2\text{C}_2\text{O}_4$, 2aq, s c $\text{CO}_2\text{H CO}_2\text{H}$ 2aq or $\text{C}(\text{OH})_2$, $\text{C}(\text{OH})_2$, [187°] (when anhydrous) (Staub a Smith, *B* 17, 1742 note) SG (hydrated) $\frac{15}{4}$

1 653 (Clarke, *Am. Z*, 174) S (of $\text{H}_2\text{C}_2\text{O}_4$) 95 at 14.5° (Nichols, *C* N 22, 14), (of $\text{H}_2\text{C}_2\text{O}_4$) 53 at 10°, 10.2 at 20°, 120 at 90° (Alluard, *O* R 59, 500), 55 at 10°, 88 at 20°, 68.8 at 70° S (alcohol) 23.5 at 15° (Burgoin, *Bl.* [2] 29, 248) S (ether) 1.27 at 15° (Miczyski, *M.* 7, 257) B_p 36.74 (in a 7.41 p c aqueous solution) (Kanonnikoff, *J* pr [9] 81, 847). H.C.v 61,100,

H C p 60,200 H.F 196,800 (Stohmann, *J* pr. [2] 40, 204), 198,000 (von Rechenberg) S.H. (0°–50°) 3359 (Hess, *A* Ch [2] 35, 410)

Heat of neutralisation 28,100

Occurrence — As acid potassium salt in sorghol (*Oxalis acetosella* and *Rumex acetosa*) (Savary, *A.D* 1773, Wiegler, *A.D* 1779), as sodium salt in *Salsola* and *Salsicornia*, as calcium salt in the root, bark, and leaves of many plants, (often found as crystals, *raphides*), in urine, urinary calculi, and in the Malpighian vessels of the caterpillar of *Sphinx convolvuli*. Calcium oxalate is found also in many lichens, and as the mineral Whewellite. Ferrous oxalate occurs in lignite beds, and ammonium oxalate in guano. Free oxalic acid occurs in some fungi (e.g. *Fistulina hepatica* (Hamlet a. Plowright, *C* N 36, 93)

Formation — 1 By the oxidising action of fused potash or of nitric acid on alcohol, glycol, sugars, starch, cellulose, citric, tartaric, myristic, and other fatty acids. Also by oxidising acetic acid and phenol with alkaline KMnO_4 (Japp, *C* J Proc 4, 91) — 2 Ammonium oxalate is among the products of the decomposition of cyanogen by water — 3 By heating sodium or potassium formate above 400° (Merz a Weith, *B* 15, 1507) — 4 By the action of NaOEt on di bromo acetic ether — 5 By passing CO_2 over a mixture of sodium and sand at about 360° (Drechsel, *Z* [2] 4, 120) — 6 By heating CCl_4 with dry KOH for some days at 220° (Geuther, *A* 111, 174) CCl_4 with dry KOH at 200° forms oxalate and hydrogen — 7 By the spontaneous decomposition of $\text{CH}_3\text{CH}(\text{ONO})\text{CO}_2\text{H}$ (Henry, *B* 12, 1837) — 8 By the action of damp oxygen upon CCl_4 , $\text{CCl}_4(\text{OMe})$ (Henry) — 9 A product in the preparation of picric acid from phenol (Perkin, *C* J Proc 4, 91) — 10 By oxidising chloranilic acid (Groves, *C* J Proc 4, 91) — 11 A by product in the preparation of nitrous ether (Frickhinger, *Ar* Ph. [8] 24, 1065)

Preparation — By heating sawdust with a mixture of NaOH and KOH at 240°–250°, the yield being less when NaOH is used alone (Thorn, *D* P J 210, 24) The product is boiled with water, and the filtrate evaporated. Sodium oxalate separates in granules, which are then boiled with milk of lime. The calcium salt is then decomposed by H_2SO_4 . Oxalic acid may be freed from the last trace of alkaline oxalates by crystallisation from boiling dilute (10 p c) HClAq . Octahedral crystals of anhydrous oxalic acid $\text{H}_2\text{C}_2\text{O}_4$ may be obtained by allowing a solution of oxalic acid (1 pt) in warm H_2SO_4 (12 pts) to stand for some days. These crystals take up water (2aq) from the air, and fall to powder (Villiers, *O* R 90, 821, cf Riechardt, *J* 1864, 871) According to Lescœur (*C* R 104, 1799) dried oxalic acid takes up 2½aq from the air

Estimation — By precipitation as calcium oxalate, followed by conversion into CaCO_3 or CaO , or by titration with KMnO_4 or baryta-water

Properties — Large monoclinic crystals (containing 2aq), v sol water and alcohol. Gives off its water of crystallisation at 100°, and, in a few weeks, over H_2SO_4 (Erdmann, *J* pr 85, 218) Partially sublimes at 150°, but is partly split up into water, CO , and CO_2 . Its aqueous solution decomposes slowly in light, but the decomposition of dilute solutions may be prevented by

heating for half an hour at 70° Neubauer (*Fr.* 9, 892) When a gas is passed through a strong solution of oxalic acid at 100° CO₂ is given off (Charles, *C R* 71, 226) A 5 p.c. solution containing 1 p.c. of uranium nitrate is rapidly decomposed by sunlight into CO₂, formic acid, and CO (Seekamp, *A* 122, 113) Ammonium oxalate is also decomposed by sunlight, but the oxalates of K, Li, and Na are but slightly affected (Downes & Blunt, *Pr* 29, 219) Ferric chloride is reduced by oxalic acid in sunlight, and also by heating above 50° (Lemoine, *Bl* [2] 46, 289) Oxalic acid reduces At and Pt from their salts Oxalic acid expels HCl when heated with dry NaCl CaCl₂ ppt. solutions of alkaline oxalates, the ppt. CaC₂O₄ being insol. acetic acid and ammonia, but sol. HCl aq., and immediately reprecipitates by ammonia

Reactions—1 Heated with glycerin (1 pt.) it yields CO₂ and formic acid, but when a small quantity of glycerin ($\frac{1}{2}$ pt.) is used, decomposition takes place at a higher temperature, allyl alcohol being formed from the glycerin Oxalic acid also yields formic acid when distilled with glycol, mannite, dulcitol, erythritol, and quercitol, but not with sugars (Lorin, *C R* 77, 129, 363, 84, 1136)—2 On heating with conc. H₂SO₄ or with P₂O₅, it is resolved into water, CO, and CO₂—3 PCl₃ yields CO, CO₂, HCl, and POCl₃ PCl₅ acts in like manner—4 Boiling nitric acid slowly oxidises it to CO₂—5 PbO₂ and MnO₂ rapidly oxidise it in aqueous solution, yielding CO₂ and the corresponding oxalate Dried oxalic acid is oxidised by rubbing with PbO, great rise of temperature occurring—6 MnO₂ and dilute H₂SO₄ completely oxidise it to CO₂, KMnO₄, and CrO₃ also oxidise it—7 Chlorine water and HClO₄ yield HCl and CO₂—8 Potash fusion yields hydrogen and a carbonate—9 Liberates Cl, Br, and I when its saturated solution is boiled with KClO₃, KBrO₃, and KIO₃, respectively (Guyard, *Bl* [2] 31, 299)—10 The copper *sinc* couple forms small quantities of glycollic acid (Balbiano & Alessi, *G* 12, 190, cf. Pimpton, *B* 11, 516)—11 SbCl₃ in chloroform forms crystals of Sb₂Cl₂C₂O₄ [149°], which are decomposed by water with formation of oxalic acid (Anschütz & Evans, *A* 239, 285)—12 Primary alcohols heated with dry oxalic acid yield a mixture of mono- and di-alkyl oxalates, on distillation the mono-alkyl oxalates split up into CO₂ and alkyl formates Secondary alcohols give but little alkyl oxalate, while tertiary alcohols are wholly split up by oxalic acid into water and hydrocarbons (Cahours & Demarcay, *C R* 83, 668, 86, 991)—13 Electrolysis of potassium oxalate yields CO₂ at the positive pole (Burgoin, *A Ch.* [4] 14, 167)—14 A solution of molybdic acid in oxalic acid yields, on addition of HNO₃ and evaporating, monoclinic crystals of H₂C₂O₄MoO₄ aq., $a, b, c = 947.1, 1.078, \beta = 98^\circ 52'$ This 'oxalo-molybdic acid' yields the salts Ag₂C₂O₄MoO₄, BaC₂O₄MoO₄, and Na₂C₂O₄MoO₄ aq. (Péchar, *C R* 108, 1058)—15 Ammoniacal cupric oxide at 160° yields (NH₄)₂CO₃ and Cu₂O (Casseneuve, *Bl* [2] 32, 277)

Salts (Souchay & Lenssen, *A* 99, 81, 100, 808, 102, 85, 41, 108, 808, 105, 245)—K₂A^{aq} monoclinic crystals, $a, b, c = 875.1, 1.157, \beta = 69^\circ 5'$ (De la Provostaye, *A Ch.* [3] 4, 454) S. 88 at 16° (Nichols)—K₂A^{aq}—KHA^{aq} Salt of sorrel.

Monoclinic crystals (Marignac, *J* 1855, 462)—KHA^{aq}—KHA^{aq} trimetric crystals, $a, b, c = 459.1, 6.196, \beta = 88^\circ 8'$ —KHA^{aq} trimetric crystals (Rammelsberg, *P* 93, 24)—KH₂A^{aq} 2aq. Trichmic prisms $a, b, c = 2.100, 3.256, \alpha = 96^\circ 12', \beta = 79^\circ 29', \gamma = 97^\circ 5'$ (Wollaston, *Tr.* 1808, 99, Anderson, *C J* 1, 231, De la Provostaye) S. 16 at 13° (N)—Na₂A^{aq} S. 3 at 15° Neutral in reaction—NaHA^{aq} monoclinic crystals which redden litmus S. 17 at 15°—Li₂A^{aq} S. 8 at 19.5°—LiHA^{aq} S. 8 at 17°—(NH₄)₂A^{aq} S. 4.2 at 15° (N), 2.2 at 0° (Engel, *C R* 102, 365) Hemihedral efflorescent trimetric prisms, $a, b, c = 776.1, 733$ (Anschütz, *B* 18, 1394) Ppt. by adding NH₄Cl or NH₄OAc to its solution (Heintz, *J* pr 87, 809)—(NH₄)₂A^{aq} 2aq Occurs in guano (Tanner, *C N* 32, 162)—(NH₄)₂HA^{aq} trimetric prisms, $a, b, c = 458.1, 559$ —(NH₄)₂HA^{aq} 2aq trichmic prisms, isomorphous with KH₂A^{aq} 2aq S. 3.25 at 0° (Engel)—(NH₄)₂H₂SO₄ monoclinic crystals.—Rb₂A^{aq} (Piccard, *J* pr 86, 449)—RbHC₂O₄—RbH₂A^{aq} 3aq (Stolba, *C O* 1878, 331)—CaA^{aq} Crystalline powder, formed by ppg hot solutions (Schmid, *A* 97, 226)—CaA^{aq} 3aq occurs, mixed with CaA^{aq} in the ppt. formed from cold solutions—(CaA^{aq})₂CaCl₂ 2aq—CaA^{aq}CaCl₂ 7aq (Fritzsche, *P* 28, 121)—SrA^{aq} S. 5 at 100°—SrA^{aq} 3aq (Wicke, *A* 90, 101)—SrH₂A^{aq} 2aq—SrA^{aq}SrCl₂ 6aq—(SrA^{aq})₂SrCl₂ 16aq (Raney, *Pr.* 14, 144)—Ba₂C₂O₄ aq S. 0.4—BaH₂A^{aq} 2aq S. 3 at 15° (Clapton, *C J* 5, 223)—Be(NH₄)₂A^{aq} (Sénarmont, *J* 1857, 295, Shadwell, *J* 1881, 681)—BeK₂A^{aq}—BeK₂A^{aq}(OH)₂ 2aq (Philipp, *B.* 16, 752)—MgA^{aq} 2aq S. 0.7 at 16°, 0.8 at 100°.—Mg(NH₄)₂A^{aq} 9aq—Mg₂(NH₄)₂A^{aq} 24aq—Mg(NH₄)₂A^{aq} 8aq—Mg(NH₄)₂A^{aq} 8aq—Mg₂(NH₄)₂A^{aq} 24aq (Brandes, *Schw* *J* 27, 18)—MgK₂A^{aq} 6aq—ZnA^{aq} 2aq—Zn(NH₄)₂A^{aq} 8aq (Kayser, *P* 60, 140)—ZnK₂A^{aq} 4aq—CdA^{aq} 3aq S. 0.08 in the cold, 0.09 at 100°—CdA^{aq}(NH₄)₂ aq—Cd(NH₄)₂A^{aq} 9aq—Cd(NH₄)₂A^{aq} 11aq—Cd(NH₄)₂A^{aq} 8aq—CdK₂A^{aq} 2aq—CdNa₂A^{aq} 2aq—Al₂A^{aq}(OH)₂ (Mathieu Plessey, *C R* 97, 1033)—Al₂Na₂A^{aq}(OH)₂ 9aq (Lenssen, *J* pr 86, 314)—Al₂(NH₄)₂A^{aq}(OH)₂ 6aq—Al₂(NH₄)₂A^{aq}(OH)₂ 3aq—Al₂Na₂A^{aq}(OH)₂ 6aq—Al₂Na₂A^{aq}(OH)₂ 6aq—Al₂K₂A^{aq}(OH)₂ 2aq—Al₂K₂A^{aq}(OH)₂ 2aq—Al₂Mg₂A^{aq}(OH)₂ 5aq—Al₂Ba₂A^{aq}(OH)₂ 8aq (Collin, *B* 3, 316)—CeA^{aq} 9aq (Jolin, *Bl* [2] 21, 540)—La₂A^{aq} 9aq (Clève, *Bl* [2] 21, 202)—TiA^{aq} (Crookes, *O J* 17, 150, Kuhlmann, *C R* 55, 807)—TiHA^{aq}—Sm₂A^{aq} 10aq—SmK₂A^{aq} 2aq (Clève, *Bl* [2] 48, 171)—FeA^{aq} 1aq *Humboldtine*—FeA^{aq} 2aq yellow crystals S. 0.22 in the cold, 0.26 at 100°—K₂FeA^{aq} 2aq golden needles, sol. water—K₂FeA^{aq} aq—(NH₄)₂FeA^{aq} 3aq—Fe(NH₄)₂A^{aq} 8aq greenish crystals S. 48 at 17°—Fe(NH₄)₂A^{aq} 4aq—FeNa₂A^{aq} 4aq—FeK₂A^{aq} 8aq S. 7 at 17°—KFeA^{aq} 2aq S. 92 at 21°—Ba₂FeA^{aq} 7aq—K₂Cr₂A^{aq} 6aq Deep-blue monoclinic prisms; obtained by reducing K₂Cr₂O₇ with oxalic acid (Gregory, Rammelsberg, *P* 93, 24, Werner, *C J.* 53, 404, 602, *O J* Proc. 3, 142, *O A* Schumak, *C N* 51, 152, Hartley, *C J* Proc. 8, 4) S. 20 at 15°. The solution is red with green reflex, and gives no ppt. with CaCl₂—K₂Cr₂A^{aq} 8aq; red monoclinic crystals (Croft, *P M* [8] 21, 197) S. 10—Na₂Cr₂A^{aq} 8aq blue prisms—Na₂Cr₂A^{aq} 9aq—(NH₄)₂Cr₂A^{aq} 8aq blue scales. S. 75 at 15°—

(NH₄)₂Cr₂A''⁸, 8aq⁺ red crystals — Ag₂Cr₂A''⁹, 9aq blue needles S 11 at 100° — Ba₂Cr₂A''¹⁰, 12aq dark violet needles — Ba₂Cr₂A''¹¹, 8aq (Werner) — Ba₂Cr₂A''¹², 18aq — Ba₂Cr₂A''¹³, 6aq (Clarke, *B* 14, 1640) — Ca₂Cr₂A''¹⁴, 18aq — Ca₂Cr₂A''¹⁵, 36aq (Reece, *C* R 21, 1116) — Cr₂Cl₂A''¹⁶, 10NH₃ (Joergensen, *J* pr [2] 20, 143, 80, 28) — Cr₂A''(NH₃)₁₂, 4aq — K₂Cr₂A''¹⁷, 4aq Pleochroic, red, blue, and green (Hartley, *Pr* 21, 499) — K₂Ca₂Cr₂A''¹⁸, 6aq (Werner) blue prisms — K₂BaCr₂A''¹⁹, 3aq — Sr₂Cr₂A''²⁰, 18aq — K₂SrCr₂A''²¹, 6aq — Pb₂Cr₂A''²², 15aq — U₂A''²³, 6aq — U₂(NH₃)₁₂A''²⁴ — U₂O₂A''²⁵, 3aq — U₂O₂(NH₃)₁₂A''²⁶, 4aq — (U₂O₂)₂K₂A''²⁷, 3aq — Mn₂A''²⁸, 2½aq S 05 in the cold, 08 at 100° — Mn₂A''²⁹, 2aq Decomposes at 150° into CO₂, CO, and MnO (Castelaz, *Bl* [2] 50, 645) — Mn₂A''³⁰, 3aq S G (of Mn₂A'')²¹⁴ 2 444 (Clarke, *Am* 2, 174) — K₂Mn₂A''³¹, 6aq black monoclinic prisms (Kehrmann, *B* 20, 1594) — Mn(NH₃)₁₀A''³², 2aq — Mn(NH₃)₁₀A''³³, 4aq — Mn(NH₃)₁₀A''³⁴, 8aq — Mn(NH₃)₁₀A''³⁵, 8aq — Mn₂A''(OH)₆, 6aq — MnK₂A''³⁶, 2aq — Co₂A''³⁷, 2aq — Co₂A''(OH)₆, 6aq — CoK₂A''³⁸, 6aq — Co₂Cl₂ 5NH₃ (Krok, *J* pr [2] 18, 239) — CoA''Br 5NH₃ — Co(OH)₄(NH₃)₂SO₃NH₃ — Co₂A''(NO₂)₅NH₃ — CoA''(NO₂)₅aq — Co₂N₂(NH₃)₄, 4½aq (Rautenberg, *A* 113, 360) — Ni₂A''³⁹, 2aq — NiA''(NH₃)₃, 3aq (Winkelblech, *A* 13, 278) — K₂NiA''⁴⁰, 6aq — CuA''⁴¹, 3aq — Cu(NH₃)₄A''⁴², 2aq — CuA''(NH₃)₄, 2aq — CuA''NH₃ — CuLi₂A''⁴³, 2aq (Troost, *A* Ch [3] 51, 103) — CuK₂A''⁴⁴, 2aq — CuK₂A''⁴⁵, 4aq — CuNa₂A''⁴⁶, 2aq — PbA'' (Pelouze, *A* Ch [2] 79, 104, [3] 4, 104) — Pb₂A''O₂ — Pb₂A''(NO₂)₂, 2aq pearly plates (Johnston, *P* M [3] 13, 25, Dujardin, *J* pr 15, 308) — Pb₂A''(NO₂)₂O₂, 3aq — PbK₂A''⁴⁷, 2½aq (Reis, *B* 14, 1174) — Hg₂A''⁴⁸ white pp, insol water — Hg₂A''⁴⁹, 1aq — Hg(NH₃)₄A''⁵⁰, 2aq — HgK₂A''⁵¹, 2aq — Bi₂A''⁵², 7½aq — Bi₂A''⁵³(OH)₂, 3aq — Bi(NH₃)₄A''⁵⁴, 12aq — BiK₂A''⁵⁵, 12aq — BiK₂A''⁵⁶, 12aq — BiK₂A''⁵⁷, 2aq — Sb₂A''⁵⁸, 10aq — Sb₂K₂A''⁵⁹, 1aq — Sb₂K₂A''⁶⁰, 3aq — SbK₂A''⁶¹, 2aq (Kay, *C* N 57, 193) — SbK₂A''⁶², 4aq (Wagner, *Chem Zeit* 12, 1726) crystallises also with 1½aq, 4½aq, and 6aq — Sb₂K₂H₂A''⁶³, 3aq — SbNa₂A''⁶⁴, 9aq (Svenssen) — SbNa₂A''⁶⁵, 10aq — Sb(NH₃)₄A''⁶⁶, 2aq — Sb(NH₃)₄A''⁶⁷, 5aq (Svenssen, *B* 3, 314) — Sb₂(NH₃)₄A''⁶⁸, 16aq — SbH(NH₃)₄A''⁶⁹, 7aq — AsK₂A''⁷⁰, 3aq — Pd(NH₃)₄A''⁷¹, 2aq (Kane) — Pd(NH₃)₄A''⁷², 8aq — PtNa₂A''⁷³, 4aq — PtCl₂A''⁷⁴, 4NH₃ — PtK₂A''⁷⁵, 2aq (Clève, *Bl* [2] 45, 191) — Pt(NH₃)₄A''⁷⁶, 2aq — PtAg₂A''⁷⁷, 2aq — PtCaA''⁷⁸, 6½aq — PtCaA''⁷⁹, 4½aq — PtCaA''⁸⁰, 8aq — PtSrA''⁸¹, crystallises with 3½aq, 6½aq, and 8aq — PtBaA''⁸², 3aq — PtBaA''⁸³, 2aq — PtMgA''⁸⁴, 6aq — PtFeA''⁸⁵, 6aq — PtMnA''⁸⁶, 7aq — PtNiA''⁸⁷, 7aq — PtZnA''⁸⁸, 7aq — PtCoA''⁸⁹, 8aq — PtCdA''⁹⁰, crystallises with 5aq, 4½aq, and 4aq — PtCuA''⁹¹, 6aq — PtHg₂A''⁹², 1½aq — PtHg₂A''⁹³, 2aq — PtPbA''⁹⁴, 3aq — PtA''⁹⁵ 2NH₃ — PtA''(NH₃)₄, 1aq — PtA''(NH₃)₄, 1½aq — SnA''⁹⁶ — Sn(NH₃)₄A''⁹⁷, 2aq — SnK₂A''⁹⁸, 2aq — Sn₂A''O₂, 6aq — Th₂A''⁹⁹, 2O₂, 12aq — Ag₂A''¹⁰⁰ white pp — Ag₂A''¹⁰¹ 4NH₃.

Mono methyl ether MeHA'' (109° at 12 mm) Solid (Anschütz a Schönfeld, *B* 19, 1442, *A* 254, 8) — KMeA'' (Salomon, *B* 8, 1509)

Di-methyl ether MeA'' Mol w 118 [64°] (164°) (Dumas a Péligot, *A* Ch [2] 58, 44, Wöhler, *A* 81, 876, Erlenmeyer, *N Rep Pharm* 23, 624, Purdie, *C* J 51, 629) S V 116 7 H F 180,900 (Stohmann, *J* pr [2] 40, 353) Monoclinic tables, sol water, alcohol, and ether Its

aqueous solution slowly decomposes into oxalic acid and MeOH

Tetra-methyl ether CO₂Me C(OMe), (76° at 12 mm) S G ½ 1 1312 Formed from CO₂Me CCl₂(OMe) and NaOMe (Anschütz, *I* 254, 31) Converted by PCl₅ into Me C₂O₄

Mono ethyl ether EtHA'' **Ethylloxalic acid** (117° at 15 mm) S G ½ 1 2175 Formed from Et₂C₂O₄ (1 mol) and KOH (1 mol) in alcohol (Mitscherlich, *P* 33, 332) Formed also by heating anhydrous oxalic acid (1 pt) with absolute alcohol (1 pt) at 135°, decanting from unused oxalic acid, and distilling *in vacuo* (Anschütz, *B* 16, 2413, *A* 254, 9) Liquid When distilled under atmospheric pressure it yields formic acid and Et₂C₂O₄ — KEtA'' scales, decomposed by heat into CO and KEtCO₂ (Elite koff, *B* 6, 1259)

Di ethyl ether EtA'' **Oxalic ether** (186° cor) S G ½ 1 0793 μ 1 4156, n_D^{20} 1.4156 (Bruhl) S V 166 2 (Bruhl, *A* 203, 27), d_4^{20} 1.0856, d_4^{25} 1.0761 (Perkin, *C* J 45, 508) M M 6 654 at 12 8° S H 45 (R Schiff, *G* 17, 286) Formed by distilling dehydrated oxalic acid (11 pts) with absolute alcohol (1½ pts) (Bergmann, *Opuscula*, 1 256, Löwig, *J* pr 83, 129), or by heating KHC₂O₄ (1 pt) with alcohol (1 pt) and H₂SO₄ (2 pts) (Dumas a Boullay, *J* Ph 14, 113)

Preparation — Equal weights of dry oxalic acid and alcohol (97 p c) are boiled for 4 hours and distilled, as soon as the thermometer reaches 110° a quantity of alcohol equal to the weight of the distillate is added, and the mixture boiled again for 4 hours, the mixture is then distilled, 825 g oxalic acid give 750 g (or 56 p c) oxalic ether and 110 g formic ether (Schatzky, *J* pr [2] 34, 500)

Properties — Colourless oil with slight odour, v sol alcohol Decomposed by water, especially on heating Alcoholic potash yields a pp of KEtC₂O₄ With SnCl₂ it forms crystalline Et₂C₂O₄SnCl₂, decomposed by water (Lewy, *C* R 21, 371) TiCl₄ yields Et₂C₂O₄TiCl₄ and Et₂C₂O₄2TiCl₄ (Demarcay, *C* R 70, 1414)

Reactions — 1 Gaseous NH₃ yields CO₂Et CONH₂ — 2 Aqueous NH₃ forms oxamide 3 Ethylamine yields C₂O₄(NH₂)Et, diethyl amine forms CO₂Et CONEt, while triethyl amine has no action (Hofmann) Alanine forms two compounds C₂H₅N O₂ [127°] and [154°] — 4 Sodium and potassium decompose it, yielding CO and Et₂CO NaOEt acts in the same way (Geuther, *Z* [2] 4, 656) — 5 Sodium-amalgam yields des oxalic ether C₂H₅EtO₂, and, when alcohol is present, tartaric and glycolic acids — 6 ZnEt₂, followed by water, yields CO₂Et CEt₂(OH) (Frankland, *Pr* 12, 396) Other zinc alkyls act in the same way (Frankland a. Duppa, *Pr* 13, 140, 14, 17, 79, 83, 191) — 7 Acted on by Zn and a mixture of EtI and allyl iodide a mixture of HO CEt₂CO Et and HO C(CH₃)₂CO₂Et is formed, and not HO CEt(C₂H₅)CO₂Et (Barataeff, *J* pr [2] 35, 7) 8 Resorcin (1 mol) and NaOEt (2 mols) in alcohol slowly form C₂H₂O₄, which crystallises from alcohol in pale yellow prisms [to 256°], and yields the acetyl derivative C₂H₃AcO₄ [127°] (Michael, *J* pr [2] 35, 510) — 9 Phenylhydrazine forms C₂O₄(NH₂Ph)₂ [278°] and N₂H₂Ph CO CO₂Et [119°] (E Fischer, *A* 190,

181, Bulow, *A* 236, 197) In presence of alcoholic NaOEt the product is $N_2H_5PhCOCO_2H$ [170°] (Michael, *J pr* [2] 35, 458) — 10 Acetone in presence of NaOEt (1 pt) in alcohol (80 pts) forms $CH_3COCH_2COCO_2Et$ [18°] (214°) This body is converted by baryta water into oxotoluic acid, an intermediate body being $CH_3AcC(OH)(CO_2H)CHAcCO_2Et$ [90°] Acetone (2 mols) and NaOEt (2 mols) in ether yield $CH_3COCH_2COCO_2CH_2COCH_3$ [121°] crystallising in white prisms (Claisen a Stylos, *B* 20, 2188, 21, 1141, 22, 3271, 24 116) A mixture of acetone, NaOAc, KOAc, and Ac_2O yield a coloured product $C_8H_8O_4$. Another product of the action of alcoholic NaOEt on acetone and oxalic ether is $CO(CH_2COCO_2Et)_2$ [104°], which yields cheilidonic acid on warming with mineral acids On adding sodium to a mixture of oxalic ether and EtOAc dissolved in Et_2O , oxalacetic ether is formed — 11 Succinic ether and NaOEt yield oxalosuccinic ether $CO_2EtCH(CO_2CO_2Et)CH_2CO_2Et$ and a compound $C_8H_8O_4$ [90°] (Wishnecus, *B* 22, 889) — 12 Alcoholic acetophenone and NaOEt form benzoyl pyruvic acid $C_6H_5COCH_2COCO_2H$ [158°] Acetophenone and NaOEt in ether yield $BzCH_2COCO_2CH_2Bz$ [180°] (Claisen, *B* 21, 1131) — 13 Chloroacetic ether and zinc yield 'ketipic' ether $CO_2EtCH_2COCO_2CH_2CO_2Et$ [77°] (Fittig a Daimler, *B* 20, 202) — 14 Heating with oxalic, formic, or acetic acids at 140° yields formic ether, CO, and CO Benzoic acid has no action even at 240° (Lorin, *Bl* [2] 49, 344) — 15 Benzyl cyanide and NaOEt yield $CNCHPhCOCO_2Et$ [130°] whence boiling dilute H_2SO_4 forms phenyl pyruvic acid [155°] (Erlenmeyer *B* 22, 1483) — 16 A solution of urea in alcoholic NaOEt gives a pp of sodium parabanate $CONNa > CO$ (Michael, *J pr* [2] 35, 457) — CO NH

17 Phthalide and NaOEt in ether yield

$$C_6H_4 \begin{cases} \text{CH CO CO}_2\text{Et} \\ \text{CO O} \end{cases} \quad [121^\circ], \text{ whence phenyl}$$

hydrazine forms $C_{10}H_{16}N_4O_4$ [159°] (Wishnecus, *A* 246, 342) — 18 Chlorine in sunlight forms $(C_6Cl_5)_2O_4$ [144°], whence potash yields CCl_3CO_2K (Malaguti, *A Ch* [2] 74, 299)

Tetra-ethyl ether $CO_2EtC(OEt)_2$ (98° at 12 mm) $SG \frac{2}{2} 10020$ Formed from $CO_2EtCCl_2(OEt)$ and NaOEt (Anschutz, *A* 254, 32) Converted by PCl_5 into Et_2CO

Methylethylether CO_2MeCO_2Et (174°) $SG \frac{2}{2} 11557$ $SV 1391$ (Wiens, *A* 253, 297) Formed by distilling $KEtCO_2$ with $KEtSO_4$ (Chancel, *Compt Chim* 1850, 373, 403), and by the action of MeOH on $COClCO_2Et$ or of EtOH on $COClCO_2Me$ (Paul, *C J Proc* 2, 168) Converted by repeated distillation into a mixture of Me_2CO , and Et_2CO

Di methyl di ethyl ether $CO_2MeCO_2(OMe)_2$ (92° at 13 mm) Formed from $CO_2MeCCl_2(OMe)$ and NaOEt at 100° (Anschutz, *A* 254, 35)

Di-methyl di-ethyl ether $CO_2EtC(OEt)(OMe)_2$ (96° at 12 mm) Formed from $CO_2EtCCl_2(OEt)$ and NaOMe at 100°

n Propyl-ether $PrHA''$ (119° at 13 mm) $SG \frac{2}{2} 11578$ Liquid (Anschutz a Schönfeld, *B* 19, 1442, *A* 254, 6)

Di n-propyl ether Pr_2A'' (213 5°)

$SG \frac{2}{2} 10384$ $SV 2154$ (W, cf Cahours, *C R* 77, 749) $SH 451$

Tetra-n-propyl ether $CO_2PrC(OPr)_2$ (257°) $SG \frac{2}{2} 9566$ Formed from $CO_2PrCCl_2(OPr)$ and NaOPr (*A*)

Iso propyl ether CO_2PrCO_2H (111° at 13 mm) $SG \frac{2}{2} 11657$ Decomposes on boiling into Pr_2CO_3 , isopropyl formate, CO, CO, and water (Anschutz, *A* 254, 6) — CO_2PrCO_2K

Di-isobutyl ether $(C_4H_9)_2A''$ (225°) $SG \frac{2}{2} 1002$ $SH 457$ Yields $K(C_4H_9)_2C_2O_4$ (Cahours, *C R* 77, 1403)

Tetra-isobutyl ether $(C_4H_9O)_2CCO_2C_4H_9$ (146° at 10 mm) $SG 921$ Formed from $C_4H_9OCCl_2CO_2C_4H_9$ and $NaOCC_4H_9$ (Anschutz, *A* 254, 33)

Di n butyl ether $(C_4H_9)_2A''$ (243°) $SG \frac{2}{2} 10099$ $SV 2534$ (Wiens)

Isoamyl ether $(C_5H_{11})_2HA''$ Oil, smelling of bugs (Balard, *A Ch* [3] 12, 309) — CaA'_2 , 2aq — AgA' pearly scales

Di isoamyl ether $(C_5H_{11})_2C_2O_4$ (265°) $SG \frac{2}{2} 968$ (Delffs, *J* 1854, 26) $SH 464$ (Schiff) Oil, smelling of bugs (Balard)

Tetra amyl ether $(C_5H_{11}O)_2CCO_2C_5H_{11}$ (190° at 14 mm) $SG \frac{2}{2} 9141$ (Δ)

Ethyl heptyl ether $CO_2EtCO_2C_7H_{15}$ (263 7°) $SG \frac{2}{2} 9954$ $SV 2849$ (Wiens)

Propyl heptyl ether $Pr(C_7H_{15})_2A''$ (284°) $SG \frac{2}{2} 9814$ $SV 3157$

Propyl octyl ether $Pr(C_8H_{17})_2A''$ (291°) $SG \frac{2}{2} 9725$ $SV 3404$ (Wiens)

Allyl ether $(C_3H_5)_2A''$ (215 5° V) (Kekulé, *B* 6, 387) $SG \frac{2}{2} 1055$ Formed from $Ag_2C_2O_4$ and allyl iodide (Hofmann a Cahours, *A* 102, 288)

Di phenyl ether $Ph_2C_2O_4$ [130°] Formed by heating phenol (1 pt) with dry oxalic acid (1 pt) and $POCl_3$ (1 pt) at 115° (Nencki, *J pr* [2] 25, 283) Prisms (from alcohol), sl sol ether

Di phenyl ortho oxalate $(PhO)_2C(OH)_2$ [127°] A by product in the manufacture of aurin Formed by distilling phenol with dry oxalic acid, or by dissolving oxalic acid and excess of phenol in HOAc (Claparède a Smith, *C J* 43, 358, Staub a Smith, *B* 17, 1740) Thin white plates, distilling with decomposition at 150°-180° Sol water, but almost at once split up into phenol and oxalic acid Alcohol yields phenol and oxalic ether. On heating with H_2SO_4 it yields aurin

Di (a) naphthyl ortho oxalate $(C_{10}H_7O)_2C(OH)_2$ [163°] Formed by heating (a) naphthol with dry oxalic acid and HOAc (S a S) Crystalline powder

Di (b) naphthyl ortho oxalate $(C_{10}H_7O)_2C(OH)_2$ [167°] White crystalline powder (from HOAc), partly decomposed on distillation (Staub a Watson Smith, *C J* 45, 303)

Penta chloro-ethyl ether $CO_2HCO_2CCl_3$ Formed from $CO(NH_2)_2$, CO_2CCl_3 , and NH_4Aq (Malaguti, *A Ch* [2] 74, 308) Colourless deliquescent needles — NH_4A'

Chloride of the methylether $CO_2MeCOCl$ (120°) $SG \frac{2}{2} 13316$ Formed by heating $CO_2MeCCl_2(OMe)$ for 40 hours at 215° (Anschutz, *A* 254, 26) Liquid

Chloride of the ethylether $CO_2EtCOCl$ Chloroacetic ether. Chloro glyoxylic ether ($q v$) (186°) $SG \frac{2}{2} 12223$ PCl_5 , acting upon oxalic

ether first forms $\text{CO}_2(\text{OEt})\text{CO}_2\text{Et}$, which may be distilled under 15 mm pressure, when distilled under atmospheric pressure it is split up into EtCl and COClCO_2Et (Anschutz, *B* 19, 2158, *A* 254, 27) Formed also from oxalic ether and POCl_3 (Henry, *B* 4, 598, 5, 949) Decomposed by water Alcoholic NH_3 yields ethyl oxamate Aniline forms $\text{CO}(\text{NPhH})\text{CO}_2\text{Et}$ Mercaptan forms $\text{CO}_2\text{EtCOSEt}$ (Morley & Saint, *C J* 43, 400) ZnEt_2 followed by water forms $\text{OEt}(\text{OH})\text{CO}_2\text{Et}$ Carbamic ether yields $\text{CO}_2\text{EtNHCOCO}_2\text{Et}$ [45°] (Salomon, *J pr* [2] 9, 290) $\text{CO}(\text{NH}_2)\text{CO}_2\text{Et}$ at 180° yields $\text{NH}(\text{COCO}_2\text{Et})_2$ [67°]

Chloride of the propyl ether
 COClCO_2Pr (154°) $\text{SG } \frac{2}{3}$ 11670 Formed by treating $\text{Pr}_2\text{C}_2\text{O}_4$ with PCl_5 and heating the resulting $\text{CO}_2\text{PrCOCl}(\text{OPr})$ at 190° (Anschutz)

Chloride of the isobutyl ether
 $\text{COClCO}_2\text{C}_4\text{H}_9$ (165°) $\text{SG } \frac{2}{3}$ 11153 Formed in like manner

Chloride of the isoamyl ether
 $\text{COClCO}_2\text{C}_5\text{H}_{11}$ (185°) $\text{SG } \frac{2}{3}$ 10931 (*A*)

Oxamic acid $\text{CO}(\text{NH}_2)\text{CO}_2\text{H}$ Mol w 89 [210°] $\text{S } 14$ at 14° Formed by heating $\text{CO}_2(\text{NH}_2)\text{CO}_2\text{H}$ or by boiling oxamide with aqueous NH_3 (Balard, *A Ch* [3] 4, 93, Tous saint, *A* 120, 237) The NH_4 salt is also formed by passing NH_3 into a cold alcoholic solution of oxalic ether (De Coppet, *A* 137, 105) Excreted when animals are fed with oxamic ether Prepared by heating an aqueous solution of $\text{CO}(\text{NH}_2)\text{CO}_2\text{Et}$ to boiling, adding ammonia gradually till the liquid is alkaline (Oelkers, *B* 22, 1566) Prepared also by heating ammonium oxalate with NH_4NO_3 for four hours at 175° (Mathieu Plessy, *C R* 109, 653) Crystalline powder, sl sol alcohol Converted by boiling water into $(\text{NH}_4)_2\text{H}_2\text{O}_4 \cdot \text{NH}_4\text{A}' - \text{NH}_4\text{A}'_{1/2}\text{aq} - \text{NaA}'_{1/2}\text{aq} - \text{KA}'_{1/2}\text{aq}$ (Engström, *J* 1856, 453) — $\text{MgA}'_{1/2}\text{aq} - \text{CaA}'_{1/2}\text{aq}$ S (of $\text{CaA}'_{1/2}$) 16 at 13°, 4 at 100° — $\text{BaA}'_{1/2}\text{aq} - \text{PbA}'_{1/2}\text{aq} - \text{Ph}(\text{OH})\text{A}'_{1/2} - \text{FeA}'_{1/2}\text{aq} - \text{NiA}'_{1/2}\text{aq} - \text{CuA}'_{1/2}\text{aq}$ (Bacaloglio, *J pr* 81, 869) — $\text{AgA}'_{1/2}$ needles

Methyl ether MeA' Cubes

Ethyl ether $\text{CO}(\text{NH}_2)\text{CO}_2\text{Et}$ **Oxamethane** [115°] Formed from oxalic ether and dry or alcoholic NH_3 Plates (from alcohol) With COClCO_2Et it forms $(\text{CO}_2\text{EtCO})_2\text{NH}$ [67] With cyanic acid vapour at 130° it yields $\text{C}_3\text{H}_2\text{N}_2\text{O}_{12}$ [155°–160°] (Grmaux, *Bl* [2] 21, 154), crystallising from water in needles. Chloral forms $\text{CCl}_3\text{CH}(\text{OH})\text{NHCO}_2\text{Et}$ [121°] (Moscheles, *B* 24, 1804)

Penta-chloro-ethyl ether $\text{C}_2\text{Cl}_5\text{A}'$ [184°] Formed from $(\text{C}_2\text{Cl}_5)_2\text{C}_2\text{O}_4$ and NH_3 (Malaguti)

Isobutyl ether $\text{C}_4\text{H}_9\text{A}'$ [90°] Prisms (Oahours, *O R* 77, 1408, Wallach, *B* 18, 507)

Isoamyl ether $\text{C}_5\text{H}_{11}\text{A}'$ [93°]

Phenyl ether $\text{CO}(\text{NH}_2)\text{CO}_2\text{Ph}$ [182°] Formed from $\text{CO}_2\text{EtCOCl}(\text{NH}_2)$ and phenol

Acetyl derivatives of the ethyl ether $\text{CO}(\text{NHAc})\text{CO}_2\text{Et}$ [54°] Needles

Oxamide $\text{CO}(\text{NH}_2)\text{CO}(\text{NH}_2)$ S 037 at 78° (Henry, *C R* 100, 943) H.F. 169,000 (Berthelot) Formed from oxalic ether and NH_4Aq (Bauhot, *ad* 1817) Formed also by heating ammonium oxalate (Dumas, *A Ch* [2] 44, 129, 54, 240) and by the slow decomposition of cyanogen by water containing aldehyde (Laebig, *A* 118,

248), or by conc HClAq (Schmidt & Gluts, *B* 1, 66) It also occurs among the products of oxidation of HCy , cyanides, and ferrocyanides (Playfair, *Attfield, C J* 16, 94)

Properties — White powder, sl sol hot water, insol alcohol Cupric acetate forms the salt $\text{Cu}(\text{C}_2\text{H}_3\text{N}_2\text{O}_4)_2\text{aq}$

Reactions — 1 By passing through a red hot tube it is decomposed into CO , ammonium carbonate, HCy , and urea. — 2 P_2O_5 yields, on heating, cyanogen, CO_2 , and CO — 3 Boiling dilute acids yield oxalic acid — 4 Boiling aqueous alkalis also saponify it *Magnesia* acts in the same way (Berthelot, *Bl* [2] 47, 840) — 5 Water at 224° forms ammonium oxalate — 6 Heating with HgO yields urea, CO_2 , and Hg (Williamson) Boiling with water and HgO forms a compound $(\text{C}_2\text{H}_3\text{N}_2\text{O}_4)_2\text{HgO}$ (Dessaignes, *A* 82, 233) — 7 Ac_2O has no action at 160° Bz_2O at 200° forms benzamide — 8 Conc HNO_3 decomposes it in the cold

Oxalimide $\text{CO} \rangle \text{NH}$ Formed from oxamic acid, PCl_5 , and POCl_3 at 80°–90° (Ost & Meute, *B* 19, 3228) Monoclinic prisms, v sl sol cold water, sol NH_4Aq Boiling water produces oxamide and oxalic acid — $\text{C}_2\text{O}_2\text{NH}_2\text{HCl}$ crystal line powder, insol water

Methyl oxamic acid $\text{CO}(\text{NHMe})\text{CO}_2\text{H}$ [146°] (Hantzsch, *B* 17, 2919) Formed by heating methylamine acid oxalate (Wurtz, *A Ch* [3] 30, 443), and, as a subsidiary product, by oxidising caffeine with CrO_3 (Maly & Hinteregger, *M* 2, 128) Crystals (from hot water or by sublimation) — $\text{KA}' - \text{CaA}'_{1/2} - \text{CaA}'_{1/2}\text{aq} - \text{BaA}'_{1/2}\text{aq}$ monoclinic crystals, $a b c = 1.018 \ 1.1306$, $\beta = 87^\circ 13'$

Methyl ether MeA' [85°] Formed from $\text{Me}_2\text{C}_2\text{O}_4$ (76 g) and methylamine (20 g) in MeOH (30 g) (Franchimont & Klobbie, *R T C* 8, 305)

Ethyl ether EtA' (243°) Oil

Methyl oxamide $\text{CO}(\text{NH}_2)\text{CO}(\text{NHMe})$ [229°] Formed from oxamic ether and NMeH_2 (Wallach & West, *B* 9, 262) Minute needles

Di-methyl-oxamide $\text{C}_2\text{O}_2(\text{NHMe})_2$ [217°] (Mylius, *B* 17, 291) Needles Conc HNO_3 yields $\text{C}_2\text{O}_2(\text{NMeNO}_2)_2$ [124°] (Franchimont, *R T C* 2, 94, 4, 193) PCl_5 yields $\text{C}_2\text{H}_2\text{ClN}_2$

Di-methyl-oxamic acid $\text{CO}(\text{NMe}_2)\text{CO}_2\text{H} - \text{CaA}'_{1/2}$ crystalline (Duvillier, *A Ch* [5] 23, 315)

Ethyl ether EtA' (c 244°) Not attacked by HNO_3 (S G 15) (Franchimont & Klobbie, *R T C* 8, 304)

Ethyl-oxamic acid $\text{CO}(\text{NHEt})\text{CO}_2\text{H}$ [120°] Tables — $\text{CaA}'_{1/2}\text{aq}$ Prisms (Hantz, *A* 127, 48) — $\text{CaA}'_{1/2}\text{aq}$ S 817 at 175° — $\text{BaA}'_{1/2}\text{aq}$

Ethyl ether EtA' (245°) Oil (Wallach, *A* 184, 59) Decomposed by water

Ethyl-oxamide $\text{CO}(\text{NH}_2)\text{CO}(\text{NHEt})$ [208°] Needles (*W*)

s-Di-ethyl oxamide $\text{CO}(\text{NHEt})\text{CO}(\text{NHEt})$ [179°] (Schiff, *B* 17, 1084) Formed from oxalic ether and ethylamine (*W*)

u-Di-ethyl-oxamide $\text{CO}(\text{NH}_2)\text{CO}(\text{NEt}_2)$ [126°] (268° cor) From $\text{CO}_2\text{EtCO}(\text{NEt}_2)$ and ammonia (Wallach, *A* 214, 263) With PCl_5 it yields 'chloroxalethylene' $\text{C}_2\text{H}_2\text{ClN}_2$

Tri-ethyl-oxamide $\text{CO}(\text{NHEt})\text{CO}(\text{NEt}_2)_2$ (258°) Formed from diethyl oxamic ether and ethylamine (*W*).

Methyl ethyl-oxamide

$\text{CO}(\text{NHMe})\text{CO}(\text{NHEt})$ [157°] Formed from methylamine and $\text{CO}_2\text{Et CONHET}$ (W)

Di-phenyl oxamic acid $\text{CO}(\text{NEt}_2)\text{CO}_2\text{H}$ [101°]

Prisms PCl_5 yields $\text{CO}(\text{NEt}_2)\text{COCl} - \text{CaA}'_2, 2\text{aq}$

Ethyl ether EtA' (250°-254°)

Isopropyl oxamic acid $\text{CO}(\text{NHPr})\text{CO}_2\text{H} -$

CaA'_2 (Duvillier)

Di propyl oxamide $\text{C}_3\text{O}_2(\text{NHPr})_2$ [162°]

Plates, which feel fatty (Wallach, *A* 214, 312)

Di-isobutyl oxamide [167°] Plates, v sol.

alcohol (Malbot, *C R* 104, 228)

Isoamyl-oxamide [181°] Needles

Di isoamyl-oxamide [129°]

Di amyl-oxamide $\text{C}_5\text{O}_2(\text{NH CH}_2\text{CMe}_2)_2$ [165°]

Needles (Freund, *B* 23, 2868)

Di allyl oxamide $\text{C}_3\text{O}_2(\text{NHCH}_2\text{CH}_2)_2$ [154°]

(274°) Br yields $\text{C}_3\text{O}_2(\text{NHCH}_2\text{CH}_2\text{Br})_2$ (Wallach a

Strecker, *B* 13, 513)

Ethylene oxamide $\text{C}_2\text{O}_2\text{N}_2\text{H}_2(\text{C}_2\text{H}_4)$ Amor

phous precipitate formed, together with soluble

$(\text{CO}_2\text{Et CO})_2\text{N}_2\text{H}_2\text{C}_2\text{H}_4$ by the action of alcoholic

ethylene diamine on oxalic ether (Hofmann, *B*

5, 247) Similar products are obtained from

propylene-diamine (Strache, *B* 21, 2360)

Ethylidene oxamide $\text{C}_2\text{O}_2\text{N}_2\text{H}_2(\text{CHMe})$

Formed from cyanogen and crude aldehyde (Ber-

thelot, *A* 128, 338)

Phenyl oxamic acid $\text{CO}(\text{NHPh})\text{CO}_2\text{H}$ **Ox-**

amic acid [151°] Formed by heating dehy-

drated oxalic acid (20 g) with aniline (25 g) at

140° for an hour (Laurent, *A* 68, 15, Claus, *Z*

[2] 4, 158, Aschan, *B* 23, 1820) The product

is crystallised from water and the acid set free

by dilute H_2SO_4 . Needles (containing Aq) or

anhydrous scales (from ether), sl sol cold water,

v sol alcohol Yields CO , water, CO_2 , and di-

phenyl oxamide on heating strongly Yields on

nitration *p* nitro phenyl oxamic acid [310°]

crystallising in prisms containing aq (Aschan,

B 18, 2936) The isomeric *o* nitro phenyl-

oxamic acid [112°] is formed by heating oxalic

acid with *o*-nitro aniline (Hübner a Heoff, *A*

209, 367) ***m* Nitro phenyl oxamic ether** [150°]

is formed from oxalic ether and *m* nitro aniline

$-\text{NH}_2\text{A}' - (\text{NH}_2)\text{HA}'_2 - \text{KA}'\text{aq} - \text{NaA}'_3\text{aq}$

plates, v. sl sol cold water $-\text{PbA}'_2 - \text{CuA}'_2$,

(Anschütz, *B* 22, 736) $-\text{BaA}'_2 - \text{AgA}'_2$ white

tables $-(\text{NH}_2\text{Ph})\text{HA}'_2$, needles (from water)

Chloride $\text{CO}(\text{NHPh})\text{COCl}$ [82°]

Methyl ether MeA' [114°] Formed

from $\text{Me}_2\text{C}_2\text{O}$, and aniline (Anschütz, *A* 354, 10)

Plates (from alcohol) or needles (from ligroin)

Ethyl ether EtA' [67°] Converted by

AcCl into $\text{CO}(\text{NPhAc})\text{CO}_2\text{Et}$ [67°] PCl_5 yields

NHPH CCl , CO_2Et [72°], which splits up on melt-

ing into HCl and $\text{NPh CCl CO}_2\text{Et}$ [91°], whence

aniline forms $\text{NHC}_6\text{H}_5\text{ CO C(NPh)(NHPH)}$ [235°].

Propyl ether PrA' [92°] Needles

Isopropyl ether PrA' [52°] Needles.

Isobutyl ether C₄H₉A' [85°] Plates

Amyl ether C₅H₁₁A' [50°] Needles

References—Bromo-, Carboxy-, Di-chloro-

and Iodo- PHENYL OXAMIC ACID

Phenyl-oxamide $\text{CO}(\text{NHPh})\text{CONH}_2$. A pro-

duct of the action of HClAq on cyananiline

(Hofmann, *A* 73, 181) Formed also from

phenyl oxamic ether and NH_3 (Klinger, *A* 184,

279) Crystals (from water)

Di-phenyl-oxamide $\text{CO}(\text{NHPh})\text{CO}(\text{NHPh})$

Oxanilides [245°] (H), [241°] (F); [252 5°]

(Reissert, *B* 28, 2245) (320°) Formed by heating aniline oxalate at 170° (Gerhardt, *A Ch* [3] 14, 120, 15, 88) and by decomposing cyananiline with HClAq (Hofmann, *A* 65, 56, 73, 181, 74, 85) Formed also from ethyl camphor-oxalate and aniline (Tingle, *C J* 57, 655) Nacreous scales, insol water, sl sol hot alcohol

Reactions—1 **Nitrous acid** passed into its solution in glacial acetic acid forms the nitro-amine $\text{CO}(\text{NHPh})\text{CO}(\text{NPhNO})$ [86°] and di-*p*-nitro oxanilide (Fischer, *B* 10, 960, Sent, *J pr* [2] 35, 521)—2 **Chlorine** forms tetra chloro di-phenyl-oxamide [c 255°] (Dyer a Mixer, *Am* 8, 349)—3 **Bromine** yields $\text{C}_2\text{O}_2(\text{NHC}_6\text{H}_4\text{Br})_2$ [above 300°] whence HNO_3 yields the nitro-compounds $\text{C}_2\text{O}_2(\text{NH}_2\text{C}_6\text{H}_4\text{Br}(\text{NO}_2))_2$ [388°] and $\text{C}_2\text{O}_2(\text{NHC}_6\text{H}_4\text{Br}(\text{NO}_2))_2$ [c 287°] (Muxter a Willcox, *Am* 9, 362)—4 **Iodine** HNO_3 and HOAc yield $\text{C}_2\text{O}_2(\text{NHC}_6\text{H}_4\text{I}(\text{I} 4))_2$ —5 **Nitric acid** forms the compounds $\text{C}_2\text{O}_2(\text{NH C}_6\text{H}_4(\text{NO}_2))_2$ [260°], $\text{C}_2\text{O}_2(\text{NH C}_6\text{H}_3(\text{NO}_2)_2)_2$ melting at [300°], and $\text{C}_2\text{O}_2(\text{NH C}_6\text{H}_2(\text{NO}_2)_3)_2$ [300°] (Muxter a Walther, *Am* 9, 355, cf Hubner, *A* 209, 366) The hexa nitro oxanilide is converted by aqueous KHCO_3 into $\text{CO}(\text{NH}_2)\text{CO NH C}_6\text{H}_4(\text{NO}_2)_3$ [c 260°] and picric acid Di *o* and di-*m* micro oxanilides melt above 300° and 270° respectively (H) di *o* nitro oxanilide yields, on reduction by tin and HOAc , a base $\text{C}_6\text{H}_4\text{N}_4$ [above 300°] which forms the salts $\text{B}^+\text{H}_2\text{Cl}_2, 2\text{aq}$ and $\text{B}^+\text{H}_2\text{SO}_4, 2\text{aq}$

References—Di bromo and Tetra-chloro- PHENYL OXAMIDE

Phenyl oxamide carboxylic acid

$\text{CO}(\text{NH}_2)\text{CO NHC}_6\text{H}_4\text{CO}_2\text{H}$ Formed by boiling

the so-called carboxamido carbidamidamido benz

oic acid $\text{CO}(\text{NH}_2)\text{C(NH)}\text{NH C}_6\text{H}_4\text{CO}_2\text{H}$ with

water (Griess, *B* 18, 2411) Small white plates.

$-\text{BaA}'_2, 5\text{aq} - \text{AgA}'_2$

Di-phenyl-oxamide *m*-carboxylic acid

$\text{CO}(\text{NHPh})\text{CO NHC}_6\text{H}_3(\text{CO}_2\text{H})$ [1 3]

Amloxal-benzamic acid [300°-305°] Formed from

carboxy phenyl oxamic acid (or its mono ethyl

ether) and boiling aniline (Schiff, *A* 232, 135)

Plates (from alcohol)

Anilide $\text{NHPH CO CO NH C}_6\text{H}_4\text{CO NHPh}$.

[290°-295°] Small needles

Di-phenyl-oxamide di *m*-carboxylic acid

$\text{C}_2\text{O}_2(\text{NH C}_6\text{H}_3(\text{CO}_2\text{H}))_2$ Formed by heating

$\text{CO}_2\text{Et CO NH C}_6\text{H}_3(\text{CO}_2\text{H})$ (Schiff, *A* 232, 137)

Mono amide $\text{C}_6\text{H}_5\text{N}_2\text{O}_2(\text{NH}_2)$ Formed

by heating amido-benzamide with the compound

$\text{CO}_2\text{H CO NH C}_6\text{H}_3(\text{CO}_2\text{H})$ Crystalline At 200°

it yields theamide $\text{C}_6\text{H}_5\text{N}_2\text{O}_2(\text{NH})$, a powder,

sol ammonia

Di amide $\text{C}_2\text{O}_2(\text{NH C}_6\text{H}_3(\text{CO NH}_2))_2$ Formed

by heating $\text{CO}_2\text{Et CO NH C}_6\text{H}_3(\text{CO NH}_2)$ with

amido-benzamide Insoluble powder

***m* Amido-phenyl-oxamic acid**

$\text{C}_6\text{H}_4(\text{NH}_2)\text{NH CO CO}_2\text{H}$ Formed from *m*-phenyl-

ene-diamine and oxalic acid (Klusemann, *B* 7,

1263) Needles $-\text{AgA}'_2$

Di-methyl-amido-phenyl-oxamic acid v.

***p* 278.**

The **anhydrides** of the **methyl-**

hydroxides $\text{C}_6\text{H}_4\text{N}_2\text{O} < \text{NH CO} > \text{CO}$ are formed

by the action of MeI in MeOH upon the sodium

amido-phenyl oxamates, the products being de-

composed by Ag_2CO_3 (Griess, *B* 18, 2408) The

***m*-compound** crystallises with $3\frac{1}{2}$ aq and yields

the salts B'HIaq and B',H₂PtCl. The *p* isomeride crystallises (with 2½ aq) in needles, v sol hot water

m-Phenylene-oxamide C₆O₂(NH)₂C₆H₄. Formed from *m*-phenylene-diamine and oxalic ether (K) Amorphous

Di-phenyl-di-ethyl oxamide C₆O₂(NH C₂H₅Ph)₂ [180°] Crystals (Neubert, B 19, 1826)

Di-phenyl-di-methyl oxamide C₆O₂(NMePh)₂ (250°) Formed from methyl aniline and oxalic ether

o-Tolyl-oxamic acid CO(NHC₆H₄) CO₂H [137°] Formed by fusing KEtC₂O₄ with *o*-toluidine (Mauthner a Suida, M 7, 233, 9, 735) Needles (containing aq) Yields indole when heated with zinc dust —CaA', —BaA', aq —AgA'

p-Tolyl-oxamic acid [170°] Obtained from its ether CO(NHC₆H₄) CO₂Et [67°] which is formed by heating oxalic ether with *p*-toluidine

Nitro-tolyl-oxamic acid C₆H₄N₂O₅ 1e [1 4 3] C₆H₄Me(NO₂) NH CO CO₂H Formed by heating nitro-*m*-toluidine with oxalic acid (Hinsberg, B 15, 2691) Yellowish red plates (containing aq) —NaA' aq —BaA' aq

Ethyl ether EtA' [127°]

o-Tolyl oxamide CO(NH₂) CO(NHC₆H₄) (Bladin, Bl [2] 41, 129)

Di-*o*-tolyl oxamide C₂O₂(NHC₆H₄)₂ [200°] Formed by heating oxalic acid with *o*-toluidine at 220° (M a S) Crystals (from benzene) Converted by treatment with fuming HNO₃ into C₂O₂(NH C₆H₄Me(NO₂))₂ [2 1 3 5], which decomposes at about 270° (Mixer a Kleeberg, Am 11, 236) On oxidation by neutral KMnO₄ it yields C₂O₂(NH C₆H₄ CO₂H)₂, which forms the salts Cu₂A'O and Ag₂A'.

p-Tolyl-oxamide CO(NH₂) CO(NHC₆H₄) [237°] Formed from *p*-toluidine cyanide (C₆H₄NH₂)Cy₂ and HOAc (Bladin) Needles, v sol alcohol

Di-*p*-tolyl-oxamide C₂O₂(NHC₆H₄)₂ [269°] Crystals, v sol hot HOAc Yields on nitration C₂O₂(NHC₆H₄NO₂)₂ and C₂O₂(NHC₆H₄(NO₂)₂)₂

Amido-tolyl-oxamic ether C₆H₄N₂O₅ 1e C₆H₄Me(NH₂) NH CO CO₂Et Tolylene oxamethane [168°] Formed from tolylene *m*-diamine and oxalic ether in alcohol (Tiemann, B 8, 222) Plates (from alcohol) When dilute alcohol of 90 pc only is used the acid C₆H₄Me(NH₂) NH CO CO₂H [224°] is formed (Schiff, B 23, 1819) Phenyl thiocarbimide yields C₆H₄Me(NH CS NPh) NH CO CO₂Et [155°], a crystalline compound [138°], and C₆H₄Me<NH C₆O<NPh [198°] ClCO₂Et forms [1 2 4] C₆H₄Me(NH CO₂Et) NH C₂O₂Et [131°], whence alcoholic ammonia yields C₆H₄Me(NH CO₂Et) NH C₂O₂NH₂ [209°] (Schiff a Vanni, B 24, 687, 1315) The compounds [1 4 2] C₆H₄Me(NH CO₂Et) NH C₂O₂Et [128°] and [1 4 2] C₆H₄Me(NH CO₂Et) NH C₂O₂NH₂ [223°] are also crystalline The following bodies of like character are also crystalline:

[1 2 4] C₆H₄Me(NH CO NH₂) (NH C₂O₂Et) [218°], [1 2 4] C₆H₄Me(NH CO NH₂) (NH C₂O₂NH₂) [239°], [1 2 4] C₆H₄Me(NH NH C₂O₂Et) (NH C₂O₂NH₂) [210°], [1 2 4] C₆H₄Me(NH C₂O₂Et), [180°], and [1 2 4] C₆H₄Me(NH C₂O₂NH₂) [220°] The compounds [1.2.4] C₆H₄Me(NH CO₂Et) NH C₂O₂Et [170°] and

[1 2 4] C₆H₄Me(NH CONH₂) NH C₂O₂H [203°] have also been prepared.

Amide C₆H₄Me(NH₂) NH C₂O₂H [1 2 4]

[223°] From the ether and alcoholic NH₃

Anilide C₆H₄Me(NH₂) NH C₂O₂NPh [186°] (Schiff, B 24, 871)

Xylol-oxamic acid CO(NHC₆H₄Me)₂ CO₂H

[129°] Formed by heating *m*-xylidine with KEtC₂O₄ (Mauthner a Suida, M 9, 745) Needles (containing aq) On heating it yields di xylol-oxamide C₂O₂(NHC₆H₄Me)₂ [210°] —CuA' —AgA'

ψ-Cumyl-oxamic acid CO(NHC₆H₄Me)₂ CO₂H

[167°] Yellow needles which on heating yield di ψ-cumyl oxamide C₂O₂(NHC₆H₄Me)₂ [230°] —NaA' aq —KHA' —CaA' aq —AgHA' —AgA'

Di-propyl-di-benzyl-oxamide

C₂O₂(NH CH₂C₆H₅Pr)₂ [182°] (Goldschmidt a Gessner, B 22, 932) Crystalline

Tetra-phenyl di-propyl oxamide

C₂O₂(NH CH₂CHPh CH₂Ph)₂ [116°] Formed from CH₂Ph CHPh CH₂NH₂, and oxalic ether (Freund a Remse, B 23, 2862)

(a) Naphthyl-oxamic acid

CO(NHC₁₀H₇) CO₂H [180°] Formed from oxalic ether and (a) naphthylamine (Ballo, B 6, 247) Needles —KA' —CaA' —BaA' —(C₁₀H₇NH₂)HA' [154°] Needles (from water)

Ethyl ether EtA' [106°] Needles

Di-(a)-naphthyl-oxamide C₂O₂(NHC₁₀H₇)₂

[200°] Small scales (Zinn, A 108, 228)

Benzylidene-oxamide C₆H₄CH(NH) C₂O₂

Formed from oxamic ether and benzoic aldehyde (Medicus, A 157, 50) Plates (containing 'aq)

Piperidyl oxamic ether C₆H₄N CO CO₂Et

[290°] Formed from piperidine and oxalic ether (Wallach a Lehmann, A 237, 245) Oil

Oxamidoxim C(NH)(NH₂) C(NH)NH.

The hydrochloride of this base B'HClaq is formed by the action of alcoholic NH₃ on the hydrochloride of oximido ethyl ether (Pinner, B 16, 1656) It crystallises in plates, v sol water

Oxalamidoxim C(NOH)(NH₂) C(NOH)(NH₂)

[200°] Formed by the action of hydroxylamine on cyananiline or on cyanogen (E Fischer, B 22, 1932, Ephraim, B 22, 2305, Zinkeisen, B 22, 2946, Vorlander, B 24, 814) Colourless prisms, v sol hot water, sl sol alcohol ClCO₂Et forms C(NO CO₂Et)NH₂ C(NO CO₂Et)NH₂ [168°] Aldehyde produces CH₃CH<N>C<N>CHCH₃

[198°] Succinic anhydride yields the acid

CO₂H C₂H₄ C<N>C<N>C₂H₄ CO₂H

[200°] Chloral yields C₆H₅N₂O₂Cl [197°] —B'HClaq prisms, insol alcohol

Di-acetyl derivative [184°] Crystals

On heating with Ac₂O it yields

CMe<N>C<N>CMe [165°]

Di-benzoyl derivative [217°] Excess

of BzCl forms PhC<N>C<N>CPh

[246°], insol water

Di-ethyl ether

C(NOEt)(NH₂) C(NOEt)(NH₂) [115°] Formed

from oxalamidoxim, EtI, and NaOEt (Zinkeisen,

B 22, 2950) Needles, sl sol hot water

Oxalamidoxim

C(NOH)(NH CO NH₂) C(NOH) NH CO NH₂

[192°]. Formed from oxalamidoxim and conc.

aqueous potassium cyanate (Z) White needles, insol cold water

Phenyl-oxalamidoxim

$C(NO)(NHPh)C(NO)(NH)_2$ [c 180°] (Z), [148°] (Piemann, B 22, 1936) Formed from al cholic hydroxylamine hydrochloride and solid cyanamline (Zinckeisen, B 22, 2954) Plates

Ac_2O produces $C(NO)(NHPh)C\langle\begin{smallmatrix} N \\ NO \end{smallmatrix}\rangle CMe$ [172°]—B'HCl colourless needles

Di benzoyl derivative [189°] Needles

Hydroxylamide of oxalic acid $C_2O_2(NH OH)_2$ Formed from oxalic ether and hydroxylamine (Lossen, A 150, 814) Minute prisms (from water) Explodes at 105°— $NaHA''$ — KHA'' — CaA'' — $BaHA''$ — ZnA'' — Ag_2A'' — $NH_2(OH)H A''$

Oxalyl-tetra-methyl di hydrazine

$C_2O_2(NH NMe)_2$ [220°] Formed from di methyl hydrazine and oxalic ether (Renouf, B 13, 2172) Plates, sol alcohol

Oxalyl-di-ethyl di-hydrazine

$C_2O_2(NH NHet)_2$ [204°] Formed from ethyl-hydrazine and oxalic ether (Fischer, A 199, 297) Needles Gives rise to the nitrosamine $C_2O_2(N_2HPh NO)_2$ [145°], crystallising from water in prisms

Oxalphenylhydrazic acid $CO(NH_2Ph)CO_2H$ [170°] Formed from oxalic ether, phenyl hydrazine, and alcoholic $NaOEt$ (Michael, J pr [2] 35, 453)— NaA' — EtA' [119°]

Oxalyl di phenyl di-hydrazine

$C_2O_2(N_2H_2Ph)_2$ [278°] Formed from oxalic ether and phenyl hydrazine at 110° (E Fischer, A 190, 131) Plates With $COCl_2$ it yields $C_2O_2(N_2(CO)Ph)_2$ [above 300°]

Semi nitrile of oxalic acid v CYANOFORMIC ACID

Semi-nitrile of ortho oxalic acid *Tri ethyl ether* $C(OEt)_3CN$ (161°) SG 1.55 1.003 Polymerises on standing (Bauer, A 229, 178)

Tri propyl derivative $C(OPr)_3CN$ (216°-219°)

Nitrile of oxalic acid is CYANOGEN

OXALIMIDO ETHYL ETHER $C_2H_5N_2O_4$

$C(NH)(OEt)C(NH)(OEt)$ [c 25°] [c 170°] Formed from cyanogen and alcoholic HCl (Pin ner & Klein, B 11, 1481) Long prisms Benzylamine at 110° forms $(CH_2PhNH)_2C_2(NH)_2$ [150°] (Vorlander, B 24, 806) *p* Toluidine forms the isomeric $(C_2H_5MeNH)_2C_2(NH)_2$ [220°-230°], which is converted by hydroxylamine sulphate into $C(NO)(NH_2)C(NO)(NHC_2H_5)$ [175°], which yields a dibenzoyl derivative [194°], and by hydroxylamine hydrochloride into $C(NO)(NH_2)C(NH)NHC_2H_5$ [148°], whence may be got $C(NOEt)NH_2C(NH)NHC_2H_5$ [135°] and $C(NOCH_2Ph)(NH_2)C(NH)NHC_2H_5$ [165°]

OXALINES A name given to certain alkyl-glyoxalines (Wallach, A 214, 278, 325, Japp, C J 43, 197, Radziszewski, B 15, 2706) They are described as alkyl glyoxalines (v also Gly-oxalines)

OXAL-METHYL-ETHYLENE v DI-METHYL-GLYOXALINE

OXALMETHYLENE is identical with **METHYL-GLYOXALINE** (q v)

OXAL-METHYL PROPYLENE v METHYL-ETHYL-GLYOXALINE

OXALOXYL AMIDO-BENZOIC ACID v. CYN-AMIC ACID and CARBOXY-PHENYL OXAMIC ACID.

OXALPROPIONIC ETHER v METHYL OXAL-ACETIC ETHER

OXALPROPYLAMYLENE v PROPYL BUTYL-GLYOXALINE

OXALPROPYLBUTYLENE v DI PROPYL GLY-OXALINE

OXAL PROPYL ETHYLENE v METHYL PRO-PYL GLYOXALINE

OXALPROPYLENE v ETHYL PROPYL GLY-OXALINE

OXALSUCINIC ETHER

$CO_2Et CO CH(CO_2Et)CH_2CO_2Et$ (156° at 17 mm) Formed by the action of $NaOEt$ on an ethereal solution of oxalic and succinic ethers (W Wislicenus, B 22, 885) Oil, v sol alkalis $FeCl_3$ gives a deep red colour in alcoholic solutions Split up, in dilute solutions, on warming into oxalic and succinic acids and alcohol Yields a phenyl hydrazide

OXALURIC ACID $C_4H_4N_2O_8$

$NH_2CO NH CO CO_2H$ Mol w 132 HF 2,500 (Matignon, C R 113, 198) Occurs as ammonium salt in urine (Schunck, J 1866, 749) Formed by heating parabanic acid with aqueous alkalis (Liebig & Wohler, A 26, 287), and occurs among the products of the action of HCl and $KClO_4$ on guanine (Strecker, A 118, 151) Its ether is produced by the action of urea on $COCl CO_2Et$ in the cold (Henry, B 4, 644) White crystalline powder, sl sol water It reddens litmus Its aqueous solution is decomposed on boiling into urea and oxalic acid $POCl_3$ converts it into parabanic acid (Grimaux, C R 77, 1548) HNO_3 slowly decomposes it into CO_2 (54 cc), N_2O (38 cc), CO and N (15 cc together) (Franchimont, R T C 6, 216)

Salts— NH_4A' silky needles, v sol hot water— $KA'aq$ Trimetric crystals $abc = 1.601.539$ — $CaA'_2.2aq$ S 205 at 15°, 5 at 100° (P Waage, A 118, 301)— $BaA'_2.2aq$ S 158 at 9°, 18 at 100°— AgA' long silky needles

Ethyl ether EtA' [178°] (Salomon, B 9, 374) Silky needles

Amide $NH_2CO NH CO CO NH_2$ [155°-160°] Formed, together with dialuric acid, by the action of NH_4Aq and HCy on alloxan (Rösing a Schischkoff, A 106, 255, Strecker, A 113, 48) Formed also by the action of alcoholic ammonia on oxaluric ether, and of dry NH_3 on parabanic acid at 130° (Roudinsköia, Bl [2] 45, 250) Produced by heating urea with oxamic ether (Carstangen, J pr [2] 9, 143) Crystalline powder, insol cold water Converted into ammonium oxalurate by boiling water

Oxaluryl-hydrazine $CO(NH_2)CO CO N_2H_2Ph$ [215°] Formed by heating phenyl hydrazine parabanate (Skinner & Ruhemann, C J 53, 550)

Dimethyl-oxaluramide $C_4H_4N_2O_8$ [225°] Formed from di-methyl parabanic acid and alcoholic NH_3 at 100° (Menschutkin, A 178, 203) Needles

OXALYL-DI CHLORO-ACETIC ETHER v. TETRA-CHLORO TETRA-OXY-ADIPIC ETHER

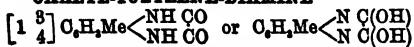
OXALYL-DI-p-DIMETHYLPHENYLENE

DIAMINE v TETRA-METHYL DI AMIDO-DI PHENYL-OXAMIDE

OXALYL-METHYL-THIO-UREA v. METHYL-THIO-PARABANIC ACID.

OXALYL-METHYL-UREA *v* *Methyl-Para*

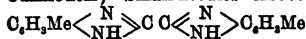
BANIC ACID

OXALYL-TOLYLENE-DIAMINE

Di-oxy-methyl-quinoxaline Formed by heating the acid oxalate of tolylene-diamine to 160° (Hinsberg, *B* 16, 1531) Converted by PCl_5 into $C_6H_4Me \begin{array}{c} \text{N CCl} \\ \text{N CCl} \end{array}$ [115°], crystallising in needles, insol water

Oxalyl-di tolylene tetramine

$C_6O_2(NH C_6H_4Me NH_2 [4 \frac{1}{3}])_2$ [above 300°] Formed by reducing di-nitro di tolyl oxamide (*v* OXALIC ACID) Small needles Above 300° it forms



Salts — $B'H_2SO_4 aq$ — $B'H_2Cl_2 aq$ — $B'H_2PtCl_6$, yellow amorphous pp

OXALYL-UREA *v* PARABANIC ACID

OXAMETHANE is the Ethyl ether of oxamic acid *v* OXALIC ACID

OXAMETHANE CHLORIDE *v*. Di CHLORO-

AMIDO ACETIC ETHER

OXAMIC ACID *v* OXALIC ACID**OXAMIDE** *v* OXALIC ACID**OXAMIDINE** *v* OXALIC ACID **ω -OXAMIDO ACETOPHENONE-OXIM**

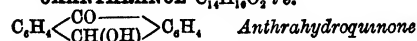
$C_6H_4C(OH)CH_2NH(OH)$ [163°] Formed by digesting a dilute alcoholic solution of ω -bromo-acetophenone with hydroxylamine hydrochloride for several hours at the boiling point (Schramm, *B* 16, 2183) White crystalline solid Sol alcohol and ether, insol cold water and ligroin Dissolves in alkalis — $C_6H_4N_2O_2Ag$

 m -OXAMIDO-CARBIMIDO-CARBOXAMIDO-BENZOIC ACID

$(OH)NH C(NH) CO NH C_6H_4 CO_2H$ Formed by the action of an aqueous solution of hydroxylamine upon cyanocarboxamido benzoic acid $NC CO NH C_6H_4 CO_2H$ (vol 1 p 157) (Griess, *B* 18, 2416) White needles SI sol hot water — BaA' , 4aq

OXANILIC ACID is Phenyl oxamic acid *v* OXALIC ACID

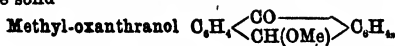
OXANILIDE is Di phenyl oxamide *v* OXALIC ACID

OXANTHRANOL $C_{12}H_{10}O_2$ *v* *se*.

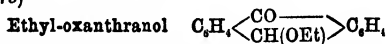
Prepared from anthraquinone (1 pt), zinc dust (2 pts), and NaOH (30 pts of 50 per cent solution) The filtrate is ppd by acids, but the ppd oxanthranol must be kept in a closed bottle under carbonic acid water, as it is reoxidised by air to anthraquinone (Graebe & Liebermann, *A* 160, 126, 212, 65) The red solution of oxanthranol in KOHAq is attacked by alkyl iodides forming alkyl-oxanthranols The alkyl-oxanthranols may be reduced to alkyl anthracenedi hydrides, which may be re-oxidised to the alkyl oxanthranols The alkyl oxanthranols give, with PCl_5 , alkyl-oxanthranyl chlorides In these respects methyl oxanthranol behaves differently from the others The alkyl-oxanthranyl chlorides are converted by water back to alkyl-oxanthranols Alkyl-oxanthranols (1 pt) are reduced by zinc-dust (2 pts) and ammonia (8 pts of SG 88) and water (5 pts) to alkyl hydro anthranols (Liebermann, *A* 212, 108)

Acetyl derivative $C_{12}H_8AcO_2$ Formed

from oxanthranol, NaOAc and Ac_2O Crystal line solid



or $C_6H_4 \begin{array}{c} \text{CO} \\ \text{CMe(OH)} \end{array} C_6H_4$, [187°] Formed by heating a mixture of anthraquinone, NaOH, zinc dust, watr, and MeBr (Liebermann & Landshoff, *B* 14, 456, *A* 212, 75) Colourless plates, sl sol alcohol, forming a solution with blue fluorescence Reduced by HI and P to anthracene dihydride Not attacked by PCl_5 An isomeric of methyl oxanthranol [98°] is sometimes formed by the action of NaOHAq and MeI on oxanthranol (Liebermann, *B* 21, 1175)



[107°] Formed from oxanthranol, NaOHAq, and EtI at 100° Formed also by oxidation of $C_6H_4 \begin{array}{c} \text{COEt} \\ \text{C(OEt)} \end{array} C_6H_4$, [77°] with CrO_3 in HOAc (Goldmann, *B* 21, 2507) Needles or trimetric prisms, $a b c = 741 \cdot 1 \cdot 495$ Its alcoholic solution exhibits strong blue fluorescence It is insol aqueous alkalis Reacts with hydroxylamine (E von Meyer, *J pr* [2] 29, 496) HI and P reduce it to ethyl anthracene dihydride Conc H_2SO_4 forms crystalline $C_{12}H_{12}O_2$ Yields a di bromo-derivative $C_{12}H_8Br_2O(OEt)$ [123°] and a di nitro derivative crystallising in small needles PCl_5 forms $C_6H_4 \begin{array}{c} \text{CO} \\ \text{C(EtCl)} \end{array} C_6H_4$, [89°]

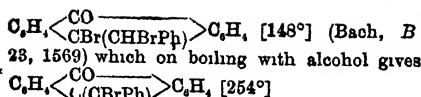
Isobutyl-oxanthranol

$C_6H_4 \begin{array}{c} \text{CO} \\ \text{C(C}_2\text{H}_5\text{)}_2\text{(OH)} \end{array} C_6H_4$, [130°] Prisms or needles (Liebermann & Walder, *B* 14, 462) PCl_5 yields the chloride $C_6H_4 C_2OCl(C_2H_5)_2 C_6H_4$, [78°], which reproduces isobutyl oxanthranol on boiling with water

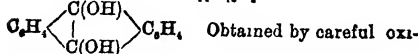
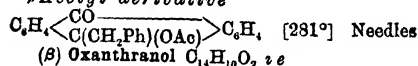
Isoamyl-oxanthranol

$C_6H_4 \begin{array}{c} \text{CO} \\ \text{C(C}_2\text{H}_5\text{)}_2\text{(OH)} \end{array} C_6H_4$, [125°] Monoclinic tables (from benzene ligroin) Conc H_2SO_4 removes H_2O , forming $C_{12}H_{10}O$ [72°] which separates from alcohol as yellow needles, and forms a dibromide $C_{12}H_8Br_2O$ [120°] Further action of H_2SO_4 forms a cherry coloured liquid, whence alcohol pppts yellow needles of $C_{12}H_{10}O$ [206°], a body which, on oxidation, yields anthraquinone carboxylic acid and a compound $C_{12}H_{12}O_2$, [157°] The compound $C_{12}H_{10}O$ is reduced by HI in HOAc to $C_{12}H_{12}$, [93°] which crystallises from alcohol in needles, and forms on nitration a compound $C_{12}H_8N_2O_5$ (Liebermann, *A* 212, 99) Phosphorus pentachloride converts isoamyl oxanthranol into the chloride $C_6H_4 \begin{array}{c} \text{CO} \\ \text{CCl(C}_2\text{H}_5\text{)}_2 \end{array} C_6H_4$, [85°] which forms monoclinic crystals, $a b c = 1266 \cdot 12752$, $\beta = 68^\circ 23'$ NaOAc converts the chloride into $C_{12}H_{10}AcO_2$, [73°]

Benzyl-oxanthranol $C_{12}H_{10}O_2$, [146°] Formed by boiling anthraquinone (5 pts), zinc dust (5 pts), KOH (7½ pts), benzyl bromide (5 pts), and water (100 pts), for a long time (Levi, *B* 18, 2153) White tables, v sol alcohol Conc H_2SO_4 at 70° forms a violet solution containing the anhydride $C_{12}H_8O$ which crystallises in yellow needles, oxidised by CrO_3 to anthraquinone. The anhydride yields a dibromide



Acetyl derivative



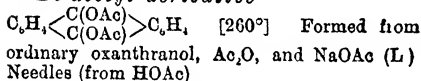
idation of anthracene by adding 56 grms of lead peroxide to a hot solution of 2 grms of pure anthracene in 50 cc of acetic acid (Schulze, *B* 18, 3036) Greenish yellow needles (from alcohol) The alkaline solution is red, but quickly becomes decolourised on shaking with air from oxidation to anthraquinone It is very oxidisable In its properties it closely resembles oxanthranol With ammoniacal AgNO_3 it gives a pp of metallic silver Cupric hydrate mixed with excess of NaOH is reduced to black cuprous oxide

Di methyl ether $\text{C}_{14}\text{H}_{10}(\text{OMe})_2$ [196°]

Di ethyl ether $\text{C}_{14}\text{H}_{10}(\text{OEt})_2$, crystals

Di benzyl ether $\text{C}_{14}\text{H}_{10}(\text{OC}_6\text{H}_5)_2$ [220°]; small colourless glistening crystals

Di acetyl derivative



*Di-oxy-(*B*)-oxanthranol* *Tri oxy anthranol*

OXATOLUIC ACID *v* **Di benzyl glycollic acid**

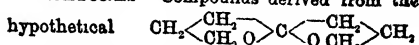
OXAZINES Compounds derived from the hypothetical oxazine $\text{NH} \begin{smallmatrix} \text{CH} & \text{CH} \\ \text{CH} & \text{CH} \end{smallmatrix} \text{O}$ Quin-

oxazine is $\text{C}_6\text{H}_5\text{N} \begin{smallmatrix} \text{O} & \text{CH} \\ \text{CH} & \text{NH} \end{smallmatrix} \text{CH}$

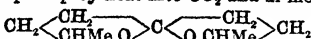
OXAZOLES Compounds derived from the hypothetical oxazole $\text{N} \begin{smallmatrix} \text{CH} & \text{CH} \\ \text{CH} & \text{N} \end{smallmatrix} \text{O}$ (Hantzsch, *B*

OXETHYL- *v* OXY ETHYL

OXETONES Compounds derived from the



They are formed by treating lactones with sodium and heating the products (Fittig, *A* 256, 57) Thus valerolactone $\text{C}_5\text{H}_8\text{O}_2$ yields $\text{C}_{10}\text{H}_{16}\text{O}_4$, which, when boiled with NaOH aq forms $\text{C}_{10}\text{H}_{14}\text{NaO}_4$, whence the acid $\text{C}_{10}\text{H}_{16}\text{O}_4$, which is split up by heat into CO_2 and di methyl oxetone



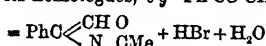
OXIDATION This term was used formerly to connote chemical changes wherein oxygen was added on to an element or compound, or a compound was decomposed by the action of O with formation of oxidised products The term was nearly synonymous with *combustion* in the earlier and more restricted meaning of that word For an account of the phlogistic theory of combustion *v* **COMBUSTION**, vol II p 241

The term oxidation has been widened until at present it is applied to all chemical changes which result in an addition of negative radicle, simple or compound, to elements or compounds, or a decrease in the relative quantity of the positive radicle of a compound, whether this is or is not accompanied by substitution of negative radicle Thus the following changes are classed together as oxidations $4\text{Fe} + 3\text{O}_2 = 2\text{Fe}_2\text{O}_3$, $2\text{Fe} + 3\text{Cl}_2 = 2\text{FeCl}_3$, $2\text{BaO} + \text{O}_2 = 2\text{BaO}_2$, $4\text{Cu} + \text{S}_2 = 2\text{Cu}_2\text{S}$, $2\text{Hgl} + \text{I}_2 = 2\text{HglI}_2$, $2\text{KNO}_3 + \text{O}_2 = 2\text{KNO}_2$, $\text{BaS} + 2\text{O}_2 = \text{BaSO}_4$, $4\text{FeSO}_4 + 2\text{H}_2\text{SO}_4 + \text{O}_2 = 2\text{Fe}_2(\text{SO}_4)_3 + 2\text{H}_2\text{O}$, $2\text{K}_2\text{FeCy}_4 + \text{Cl}_2 = 2\text{K}_2\text{FeCy}_4 + 2\text{KCl}$, $2\text{Cr}_2\text{O}_3 + 4\text{K}_2\text{O} + 3\text{O}_2 = 4\text{K}_2\text{CrO}_4$, $4\text{K}_2\text{MnO}_4 + \text{O}_2 = 4\text{KMnO}_4 + 2\text{K}_2\text{O}$, $\text{Bi}_2\text{O}_3 + 4\text{Cl}_2 + 2\text{H}_2\text{O} + 4\text{KOH} = \text{Bi}_2\text{O}_5 + 4\text{KCl} + 4\text{H}_2\text{O}$, $3\text{C}_2\text{H}_4\text{O} + 2\text{Cr}_2\text{O}_3 = 3\text{C}_2\text{H}_4\text{O} + 3\text{H}_2\text{O} + \text{Cr}_2\text{O}_3$

Processes of oxidation are accompanied by processes of reduction or deoxidation The following examples make this clear (*cf* **DEOXIDATION**, vol II p 377) —

Original element or compound		Oxidiser		Oxidised Product		Deoxidised Product
H_2	+	O	=	H_2O		H_2O
		(The H_2O may be regarded as		oxidised H, or as reduced O)		
2Hg	+	O_3	=	HgO		O_2
		(The O_2 may be regarded as reduced ozone [O_2])				
SbCl_3	+	Cl_2	=	SbCl_5		SbCl_3
		(The SbCl_3 is oxidised		SbCl_3 , or reduced Cl)		
Sn	+	2HNO_3	=	$\text{SnO}_2 + \text{H}_2\text{O}$	+	N_2O_3
$3\text{H C O}_4\text{Aq}$	+	$2\text{KMnO}_4\text{Aq}$	=	$6\text{CO}_2 + 3\text{H}_2\text{O}$	+	$2\text{MnO}_2 + \text{K}_2\text{OAc}$
PbO	+	$2\text{Cl} + 2\text{KOHAc}$	=	$\text{PbO}_2 + \text{H}_2\text{O}$	+	2KClAq
$\text{K}_2\text{FeCy}_4\text{Aq}$	+	Cl_2	=	$\text{K}_2\text{FeCy}_4\text{Aq}$	+	2KClAq
$4\text{K}_2\text{MnO}_4\text{Aq}$	+	O_2	=	$4\text{KMnO}_4\text{Aq}$	+	$2\text{K}_2\text{O}$
(In the three cases in the bracket, KCl and K_2O may be regarded as reduced Cl and O respectively						
$3\text{SO}_4\text{Aq}$	+	$2\text{HNO}_3\text{Aq} + 2\text{H}_2\text{O}$	=	$3\text{H}_2\text{SO}_4\text{Aq}$	+	2NO

21, 944) They are formed by the action of the halogen derivatives of ketones on acetamide and its homologues, *eg* $\text{Ph CO CH}_2\text{Br} + \text{Me CO NH}_2$,



Compounds derived from $\text{CH CH} \begin{smallmatrix} \text{CH} \\ \text{CH N} \end{smallmatrix} \text{O}$ may be called isoxazoles Thus phenyl isoxazole is formed by the action of AcCl on the oxim of benzoyl acetic aldehyde (Claissen *B* 24, 134)

VOL. III

Any element or compound which frequently reacts to produce substances containing relatively more negative radicle than the original substance acted on, is called an *oxidiser* or *oxidising agent* Among the substances commonly used in the laboratory to effect oxidations are oxygen, ozone, chlorine, hypochlorites, nitric acid, potassium permanganate, potassium chlorate, molten potash, and chromium trioxide The conditions under which oxidation occurs vary much, thus Hg is oxidised by ozone at the

U U

ordinary temperature, but by O only at temperatures near the B.P. of Hg, KMnO_4 oxidises $\text{H}_2\text{C}_2\text{O}_4$ completely only in the presence of H_2SO_4 , and at a moderately high temperature, O does not oxidise SO_2 under ordinary conditions, but if the gases are passed over hot spongy Pt, SO_3 is produced rapidly. When O is absorbed by charcoal, and the charcoal is then brought into contact with H_2S , PH_3 , $\text{C}_2\text{H}_2\text{O}$, &c., oxidation proceeds rapidly (*v. Calvert, C J [2] 5, 293*). The products of oxidation obtained from a specified substance often vary according to the oxidiser employed, thus ozone produces Hg_2O from Hg at ordinary temperatures, but HgO is formed by the action of O on hot Hg, carbohydrates generally give HCO_2H when oxidised by CrO_3 , but $\text{H}_2\text{C}_2\text{O}_4$ when oxidised by HNO_3 , phthalic acids, $\text{C}_6\text{H}_4(\text{CO}_2\text{H})_2$, are obtained by oxidising naphthalene, C_{10}H_8 , by CrO_3 in glacial acetic acid, but the chief product of the oxidation of the same compound by K_2CrO_4 and H_2SO_4 (which is essentially a solution of CrO_3) is naphthoquinone, $\text{C}_{10}\text{H}_6\text{O}_2$. There are some compounds the presence of which in a solution of an oxidisable body hastens the oxidation when O is passed into the solution, thus, $\text{H}_2\text{C}_2\text{O}_4$ is oxidised by CrO_3 in presence of MnSO_4 (*Harcourt, B A 1864 34*), SO_4 is rapidly oxidised by O in the presence of certain salts, especially MnSO_4 , CuSO_4 , FeCl_3 , CoCl_2 , &c. (*v. L. Meyer, B 20, 3058*, *Roessler, D P J 242, 278*, *cf. Mendelejeff, B 19, 2656*). M M P M

OXIDES Binary compounds of oxygen

For the purposes of this definition, those compounds of O with organic radicals which react similarly to oxides of elements must be classed among binary compounds (*eg* the ethers), in the present article, however, only the binary compounds of O with elements are considered. Oxides of all elements except Br and F have been isolated.

Oxides are frequently prepared by the direct union of O with other elements, O unites directly with all the elements except Br, Cl, F, I, Au, and Pt. Metallic oxides are formed by the action of heat on carbonates, nitrates, and other salts of volatisable acids, but the alkali oxides cannot be thus prepared. Most (?all) metals decompose water or steam, forming oxides or hydroxides, and evolving H. Many metallic sulphides yield oxides when roasted in air or O. Those oxides or hydroxides of metals which are not easily soluble in water are generally formed by the reaction of alkalis or alkaline oxides with solutions of metallic salts. The higher oxides of metals are often obtained by the action of hypochlorites, or Cl and KOH , on the lower oxides, sometimes by the action of conc. HNO_3 on the lower oxides or the metals, and sometimes by reacting on salts of the metals in solution with H_2O_2 in presence of an alkali.

Fusion of metallic oxides with KOH or KNO_3 generally results in the formation of alkali salts of metal-containing acids in the cases of those metals which are capable of forming such salts. The higher oxides of metals are generally more or less easily reduced to lower oxides, this reduction occurs sometimes by heating, in other cases by the action of such reducers as H, CO, SO_2 , As_2O_3 , or H_2S . The oxides of non-metals are frequently formed by combining O with the non-

metal, *eg* B_2O_3 , SO_2 , SO_3 , CO, CO_2 , H_2O , NO_2 , P_2O_3 , P_2O_5 , sometimes they are produced by such indirect methods as decomposing oxyacids or salts of oxyacids of the non-metals, *eg* N_2O_5 from HNO_3 , I_2O_5 from HIO_3 , Cl_2O from KClO_3 , sometimes they are formed by very indirect methods, *eg* Cl_2O by the reaction of Cl with HgO .

Oxides may be divided into classes in accordance with their empirical composition, thus, monoxides, M_2O and MO , sesquioxides, M_2O_3 , dioxides, MO_2 , trioxides, MO_3 , tetroxides, MO_4 , pentoxides, M_2O_5 , heptoxides, M_2O_7 .

A better classification is that based primarily on chemical properties, on this system, oxides are classified as *basic*, *acidic*, *indifferent* or *neutral*, and *peroxides*. None of these terms can be defined with strictness. The term *basic* is applied to those oxides which react with acids or with oxides more negative than themselves to form salts. *Acidic oxides* are those which react with water to produce acids, or are formed by removing water from acids, or react with oxides more positive than themselves to form salts. *Acidic oxides* are sometimes called *anhydrides*. *Peroxides* react with acids to form salts which correspond with oxides containing less O than the peroxides. Some peroxides also form acids when dissolved in water, or react with acidic oxides to produce salts. Peroxides which exhibit acidic functions may be called *acidic peroxides*, *eg* CrO_5 , peroxides which exhibit no acidic functions may be called *basic peroxides*, *eg* BaO_2 . Certain other oxides are sometimes included in the class peroxides (*v. infra*). The class of *indifferent* or *neutral* oxides includes all oxides not belonging to one or other of the three preceding classes.

Basic oxides. Most of the lower oxides of metals belong to this class. The characteristic reactions of the class are shown by the following two typical changes — $\text{BaO} + \text{H}_2\text{SO}_4 = \text{BaSO}_4 + \text{H}_2\text{O}$, $\text{PbO} + \text{SO}_3 = \text{PbSO}_4$. None of the oxides of any undoubted non-metal is distinctly basic. Oxides of non-metals, however, exist which form salts by reacting with certain strong acids, or the anhydrides of certain strong acids, and which also form salts by reacting with oxides more basic than themselves, thus, B_2O_3 reacts with H_2SO_4 containing SO_3 to form $\text{B}(\text{HSO}_3)_2$, and B_2O_3 also reacts with K_2O to form $\text{K}_2\text{B}_2\text{O}_4$, similarly As_2O_3 reacting with SO_3 forms compounds belonging to the salt type, $\text{xAs}_2\text{O}_3 \cdot \text{ySO}_3$, and with K_2O it forms KAsO_2 .

Some metallic oxides are basic, and nevertheless also form compounds with water which react as weak acids towards the more positive oxides, thus Al_2O_3 is distinctly basic, $\text{Al}_2\text{O}_3 + \text{H}_2\text{O}$ is also basic, yet it reacts with K_2O to form the unstable salt $\text{K}_2\text{Al}_2\text{O}_4$, similarly Au or $\text{3H}_2\text{O}$ dissolves in HNO_3 to form the salt $\text{Au}(\text{NO}_3)_3$, and $\text{Au}_2\text{O}_3 + \text{3H}_2\text{O}$ also dissolves in K_2O to form $\text{K}_2\text{Au}_2\text{O}_7$. The term basic oxide is sometimes widened to include oxides which correspond with salts, although these salts may not be formed directly from the oxides, thus, no salts have been obtained by the action of oxyacids on OsO_3 , but a few salts corresponding with this oxide are produced by indirect methods, *eg*, Os_2SO_5 is formed by reacting on OsO_3 with SO_2 . The *alkali-forming oxides* constitute a

division of the basic oxides, these oxides are distinctly and markedly basic, they also dissolve in water to form alkalis (*v* ALKALIS, vol 1 p 111, *v* also BASES, vol 1 p 445)

Acidic oxides or anhydrides The greater number of the oxides of non-metals belong to this class. SO_2 is a typical acidic oxide, it reacts with water to form the acid H_2SO_3 , it is obtained by removing H_2O from H_2SO_4 , by heating the acid, it reacts with basic oxides to form salts, *eg* with BaO it produces BaSO_4 . All acidic oxides do not exhibit the three characteristic reactions, some yield acids with water, but are not obtained by removing water from their acids, *eg* P_2O_5 , some are obtained by removing water from acids, but do not react with water to form acids, *eg* Sb_2O_3 , some do not form acids with water, are not obtained by removing water from acids, but react with oxides more positive than themselves to produce salts, *eg* As_2O_3 . Those acidic oxides which do not form corresponding acids by reacting with water generally show basic functions when they react with strong acids. Should an oxide form no acid with water, nor be obtained by removing water from an acid, but yet react with basic oxides to form salts, this oxide, although classed as acidic, will be found, almost certainly, to react as a basic oxide towards strong acids, or the anhydrides of strong acids. Thus, As_2O_3 does not form an acid with water, nor is it obtainable from a corresponding acid, it does, however, react with strongly basic oxides, *eg* with K_2O , to form salts, now As_2O_3 combines with SO_2 to form $\text{As}_2\text{O}_3 \cdot 2\text{SO}_2$, a compound in which As_2O_3 acts as a basic oxide. Some of the higher oxides of metals react as feebly acidic oxides, the salts corresponding with these oxides are generally obtained by fusing the oxides with KOH or NaOH (*v* ANHYDRIDES, vol 1 p 267).

Peroxides This class includes those oxides which react with acids to produce salts that correspond with oxides containing less O than the peroxides. The following reactions exhibit this typical property of peroxides — $\text{BaO}_2 + \text{H}_2\text{SO}_4\text{aq} = \text{BaSO}_4 + \text{H}_2\text{Oaq} + \text{O}$, $2\text{CrO}_3 + 12\text{HClaq} = 2\text{CrCl}_3\text{aq} + 6\text{H}_2\text{O} + 3\text{Cl}_2$. Some oxides which react in this way also dissolve in water to form acids, *eg* $\text{CrO}_3 + \text{H}_2\text{O} + \text{Aq} = \text{H}_2\text{CrO}_4\text{aq}$, others do not form acids with water, but react with strongly basic oxides to produce salts, *eg* $\text{PbO}_2 + \text{K}_2\text{O}$ (molten) — K_2PbO_4 . Peroxides which exhibit acidic functions may be called *acidic peroxides*, *eg* CrO_3 , PbO_2 , peroxides which do not exhibit acidic functions, *ie* which do not form acids with water, nor salts by reacting with basic oxides, may be called *basic peroxides*, *eg* CaO_2 , K_2O_2 . The more important *basic peroxides* are Na_2O_2 , K_2O_2 , CaO_2 , SrO_2 , BaO_2 (?) CaO_2 , CuO_2 , (?) Bi_2O_2 . Among the *acidic peroxides* may be mentioned CrO_3 , PbO_2 , and MnO_2 . Several highest oxides of metals cannot be assigned with certainty to the class of acidic peroxides, or to that of basic peroxides; *eg* Bi_2O_3 reacts with acids as a basic peroxide, and probably forms salts by fusion with a large excess of KOH , but the salts have not been isolated, UO_3 is distinctly acidic, with acids it forms uranyl salts, *eg* UO_2SO_4 ; OsO_3 is slightly acidic, no corresponding salts have been obtained by the re-

action of acids, but these reactions have not been examined sufficiently.

The term *peroxide* is used sometimes to include any oxide of a specified element which contains more O than the highest definitely basic or acidic oxide of that element. This statement does not define peroxide, because no formal definitions of the terms 'definitely basic oxide' and 'definitely acidic oxide' can be given. Such oxides as S_2O_3 , Cr_2O_3 (if it exists), MnO_2 , and Mn_2O_3 , would thus be classed as peroxides (*v* *infra*).

Indifferent or neutral oxides Oxides which do not form acids with water, are not obtained by removing water from acids, and do not form salts by reacting with either basic or acidic oxides, or with acids, are generally called *neutral or indifferent oxides*. Examples of such oxides are H_2O , NO , P_2O , (?) Fe_3O_4 , Pb_2O , Ag_2O , (?) Mn_2O .

None of the qualifying terms applied to oxides can be defined with strictness, such an oxide as MnO_2 is basic, because it forms salts by reacting with acids, MnO_2 is also acidic, because when fused with K_2O , in presence of O, it forms a salt (K_2MnO_4), it is also a peroxide, because the salts which it forms with acids correspond with the lower oxide MnO , and lastly, it may be classed as an indifferent oxide, because it does not form an acid with water, is not obtained by removing water from an acid, and does not form *corresponding* salts by the action of acids, acidic oxides, or basic oxides. When it is remembered that the properties expressed by the terms basic oxide, acidic oxide, and peroxide, are properties which come into play only when the oxides react with other substances, it is evident that these properties must depend to some extent on those of the other substances. Keeping this in mind, one sees how difficult, if not impossible, it must be to define the properties in question.

Oxides have been classified as *indifferent* and *salt forming*, and the salt forming oxides have been subdivided into those which form corresponding salts by reacting with acids or negative oxides, those which form corresponding salts by reacting with basic oxides, and those which form salts but not corresponding salts, by one or other of these reactions. This classification is practically the same as that which has been sketched already, although it is expressed in somewhat different terms.

What is the composition of the basic oxides? Which elements form acidic oxides? Can the composition of peroxides be stated in general terms? Alkali forming oxides are oxides of the most positive metals. The following are usually included in this group — Li_2O , Na_2O , K_2O , Rb_2O , Cs_2O , TiO ; MgO , CaO , SrO , BaO , (?) Ag_2O , (?) PbO . Basic oxides, which are not alkali forming, are oxides of fairly positive metals *eg* BeO , ZnO , CdO , HgO , SeO_2 , I_2O_3 , Al_2O_3 , Ga_2O_3 , Fe_2O_3 , NiO , CoO , TiO_2 , ZrO_2 , SnO , SnO_2 , Bi_2O_3 . Acidic oxides are oxides of negative elements, or they are compounds of elements, which on the whole are positive, with relatively much O, *eg* N_2O , N_2O_3 , P_2O_3 , P_2O_5 , Cl_2O , I_2O_5 , SiO_2 , CO_2 , CrO_3 , UO_3 , Ta_2O_5 . Peroxides, in the sense explained above, are usually the highest oxides of metals; *eg* CrO_3 , PbO_2 , in the widest sense, peroxides

also include some of the highest oxides of non-metals, e.g. S_2O_7 . Two oxides of the same element may exist, and one of these may be a basic, and the other an acidic, oxide, thus Cr_2O_3 is basic, but CrO_3 is acidic. Hence, whether an oxide is basic or acidic seems to depend not only on the general chemical character of the element combined with O, but also on the relative quantities of O and the other element. None of the elements whose lower oxides are alkali-forming forms an acidic oxide, but some of these elements form basic peroxides, in other words, the association of much O with a very distinctly positive element does not produce an acidic oxide, but does produce a basic peroxide. It is impossible to divide the elements into two classes, and say all on one side of the division line generally form basic oxides, but may also form acidic oxides. All that can be said is, the lower oxides of the metallic elements, as a class, are basic, but many of these elements also form higher oxides, some of which are distinctly acidic, and some are acidic peroxides, the oxides of the non-metallic elements as a class, are acidic, but some of these elements also form indifferent oxides, and a few oxides of non-metals are peroxides.

The peroxides have been divided by Mendeleeff into two classes (*J R* 1881, [1] 561, abstract in *B* 15, 242, v also Traube, *B* 19, 1111, 1115, 1117, Richarz, *B* 21, 1675). Mendeleeff distinguishes peroxides belonging to the type H_2O from those which belong to the type H_2O_2 , the latter class he calls *superoxides*, the former *polyoxides*. In the polyoxides, according to M, the O is all in direct union with the other element, and none of the O atoms is directly united with any other, whereas the O atoms of superoxides are regarded as in direct union with each other, as well as with the other element. BaO_2 is a typical superoxide, and is

supposed to have the structure $Ba \begin{smallmatrix} O \\ \diagup \quad \diagdown \\ O \end{smallmatrix} O$, MnO_2

is a typical polyoxide, and is supposed to have the structure $O Mn O$. The views of Traube (*lc*) and Richarz (*lc*) are practically the same as those expressed by Mendeleeff. The superoxides yield H_2O_2 Aq by reacting with dilute acids, and therefore reduce $KMnO_4$ Aq in presence of H_2SO_4 , the polyoxides do not yield H_2O_2 , but H_2O and O. Peroxides formed by the action of alkaline oxidisers, e.g. $KClO_4$ Aq, seem always to belong to the class of polyoxides. The peroxides K_2O_2 , Na_2O_2 , CaO_2 , SiO_2 , BaO_2 , ZnO_2 , CdO_2 , CuO_2 , PbO_2 , and some others, are superoxides, i.e. they give H_2O_2 Aq when acted on by dilute acids. According to Mendeleeff, S_2O_3 is a superoxide, it gives H_2O_2 when dissolved in much water, the constitution is probably $O SO_2 \begin{smallmatrix} O \\ \diagup \quad \diagdown \\ O \end{smallmatrix}$.

Mendeleeff (*lc*) says that the power possessed by any element of forming a characteristic basic or acidic oxide, from which salts are obtained, is connected with the position of the element in the periodic scheme of classification. But besides forming a salt-forming oxide, or more than one such oxide, some elements are also capable of producing superoxides belonging

to the type $HO OH$. Glancing at the groups of elements, as the elements are arranged in the periodic scheme of classification (*v. Classification*, vol. II p 204), and expressing the composition of the highest characteristic oxides of each group in a general formula, we have the result shown in the table on p 661.

The molecular weights of very few oxides have been determined with certainty, on this subject *v. Henry, P M* [5] 20, 81, *cf. Carnelley & Walker, C J* 58, 59. M P M

OXIDO-DI-NAPHTHYL-AMINE *v. Imido-di-NAPHTHYL OXIDE*

OXIMIDO ACETIC ETHER C_6H_5NO , i.e. $OH(NO) CO_2Et$ *Nitroso-acetic ether*. Formed, together with oxalic acid, by the action of fuming HNO_3 on acetoacetic ether (Pröpper, *A* 222, 48). Oil. Yields oxalic acid and hydroxylamine when heated with $HClAq$ at 140° . Cold $KOHAq$ yields KCy , K_2CO_3 , and alcohol. $-(NH_2)C_6H_5NO$, aq silky needles. $-C_6H_5NaNO$, $\frac{1}{2}$ aq. needles (from water).

OXIMIDO-ACETOACETIC ETHER *v. NITROSO-ACETOACETIC ETHER*

OXIMIDO- compounds *v. Nitroso- compounds*

OXIMIDO-ETHER *v. OXALIMIDO-ETHER*

TRI-OXIMIDO-METHYLENE $C_3H_3N_3O_3$, i.e. $CH_2 \begin{smallmatrix} N(OH)CH_2 \\ \diagup \quad \diagdown \\ N(OH)CH_2 \end{smallmatrix} N(OH)$. Formed from formic aldehyde and hydroxylamine (Scholl, *B* 24, 574). White amorphous solid, insol water, alcohol, and ether. At 133° it passes directly into a gas.

OXIMIDO-NAPHTHOL *v. AMIDO NAPHTHO-QUINONE*

OXIMS or OXIMES Compounds containing the group $CNOH$, obtained by the action of hydroxylamine on aldehydes, ketones, and ketonic compounds, by the reaction $R'RCO + H_2N OH = RR'C(NOH) + H_2O$ (*V Meyer, B* 15, 1164, 1324, 1525, 2784, 16, 822, 2992, 19, 1613) (*v. Aldoxims*, vol. I p 111). The reaction is best performed with hydroxylamine hydrochloride (1 mol) and aqueous $NaOH$ (3 mols) in the cold (Auwers, *B* 22, 604). Oxims are split up by boiling $HClAq$ into hydroxylamine and the original aldehyde or ketonic compound. The oxims do not exhibit Liebermann's reaction with phenol and H_2SO_4 . The oxims yield acetyl, alkyl, and sodium derivatives. Ketoxims (i.e. oxims of ketones) are converted by warming with conc H_2SO_4 at 100° into the isomeric amides, thus Ph_2CNOH becomes $Ph CO NHPh$, while $PhMeCNOH$ yields acetanilide (Beckmann, *B* 20, 1507, Wegerhoff, *A* 252, 1, Gunther, *A* 252, 44). PCl_5 acts like H_2SO_4 . Oxims are readily reduced, in alcoholic solution by means of sodium amalgam and acetic acid, to the corresponding amines, thus $XYCNOH + 2H_2 = XYCHNH_2 + H_2O$ (Goldschmidt, *B* 19, 3232). Benzoic aldehyde gives two oxims, two and two sets of alkyl oxims (Beckmann, *B* 22, 1534), and benzil also gives a greater number of oxims and alkyl-oxims than the ordinary formulæ indicate. These isomerisms may perhaps be explained by considering the arrangement in space of the atoms in the molecule (Beckmann, 20, 2766, *B* 23, 1680, Auwers & *V Meyer, B* 21, 784, 22, 1996, 23, 2408, Hantzsch, *B* 24, 81, 1192).

Groups	I	II	III
<i>Elements in group</i>	H, Li, Na, K, Cu, Rb, Ag, Cs, Au	Be, Mg, Ca, Zn, Sr, Cd, Ba, Hg	B, Al, Sc, Ga, Y, In, La, Yb, Tl
<i>Composition of highest basic or acidic oxide characteristic of the group</i>	M ₂ O	MO	M ₂ O ₃
<i>Character of highest basic or acidic oxide characteristic of the group</i>	Basic	Basic	Basic except B ₂ O ₃ , which is a weak anhydride
<i>Superoxides, type of HO OH</i>	H ₂ O ₂ , Na ₂ O ₂ , K ₂ O ₂ , CuO ₂ , Ag ₂ O ₂	CaO ₂ , ZnO ₂ , SrO ₂ , CdO ₂ , BaO ₂	(? TiO ₂)

Groups	IV	V	VI	VII
<i>Elements in group</i>	C, Si, Ti, Ge, Zr, Sn, Ce, Pb, Th	N, P, V, As, Nb, Sb, Bi, Er, Ta, Bi	(O), S, Cr, Se, Mo, Te, W, U	F, Cl, Mn, Br, I (group very incomplete)
<i>Composition of highest basic or acidic oxide characteristic of the group</i>	MO ₂	M ₂ O ₅	MO ₃	M ₂ O ₇ , represented only by Mn ₂ O ₇ , [Cl ₂ O, ClO ₂ , IO ₂ , no oxide of F or Br]
<i>Character of highest basic or acidic oxide characteristic of the group</i>	Acidic when M = C or Si, acidic and basic when M = Ti, Ge, Zr, Sn (? Ce), Pb (becoming more basic as M increases), basic when M = Th	Acidic, except Bi ₂ O ₅ , which acts as a feebly acidic peroxide (polyoxide)	Acidic, becoming less acidic as M increases, UO ₃ shows some basic properties	Acidic
<i>Superoxides, type of HO OH</i>	ThO ₂ , CeO ₂	(? NO ₃ , ? D ₂ O ₅)	S ₂ O ₇ , (? Cr ₂ O ₇ , ? UO ₄)	none

Group VIII

	Family 1 Fe, Ni, Co	Family 2 Ru, Rh, Pd	Family 3 Os, Ir, Pt
<i>Highest characteristic oxide</i>	M ₂ O ₃	MO ₃ , represented only by RuO ₄	MO ₃ , represented only by OsO ₄
<i>Character of oxide</i>	Basic	Feebly acidic (These elements also form MO, M ₂ O ₃ , and MO ₃ , which are feebly basic)	Feebly acidic (These elements also form MO, M ₂ O ₃ , and MO ₃ , which are feebly basic)
<i>Superoxides, type of HO OH</i>	? FeO ₂ , ? NiO ₂ , xO, ? Co ₂ O ₂ , xO	none	none

Oxims of ketones may be changed to phenylhydrazides by heating with phenyl hydrazine (Just, B 19, 1205)

OXINDOLE C₈H₇NO *rac.* C₈H₇ $\left\langle \begin{smallmatrix} \text{CH}_2 \\ \text{NH} \end{smallmatrix} \right\rangle \text{CO}$ or C₈H₇ $\left\langle \begin{smallmatrix} \text{OH} \\ \text{N} \end{smallmatrix} \right\rangle \text{C OH}$ (Baeyer & Comstock, B 16, 1704) *Anhydride of o-amido-phenyl acetic acid* Mol wt 133 [120°] Formed by reducing o-nitro-phenyl-acetic acid with tin and HCl. Obtained also by reducing dioxindole with sodium amalgam (Knop, J pr 97, 65, Baeyer & Knop, A 140, 1, Baeyer, B 11, 588, 12, 457) If the mixture of isomeric nitro-acids got by heating phenyl-acetic acid with fuming HNO₃ on the water bath be reduced with tin and HClAq, and, after removal of tin by H₂S, be boiled with

BaCO₃, only the *m*- and *p*-nitro phenyl acetic acids will form barium salts, and the oxindole may be extracted by ether

Properties — Long colourless needles (from water). Oxidised slowly by moist air, forming dioxindole. Reduces ammoniacal silver nitrate, forming a mirror. Extracted by ether from its alkaline solution. Not affected by boiling baryta water, but at 150° it yields barium amido phenyl-acetate. Nitrous acid forms nitroso-oxindole which is an oxim of isatin. PCl₅ yields C₈H₇ $\left\langle \begin{smallmatrix} \text{CHCl} \\ \text{N} \end{smallmatrix} \right\rangle \text{COCl}$ [104°] reduced by iron-filings and KOH to indole. Salts — AgC₈H₇NO — B/HCl. deliquescent spicules. *Acetyl derivative* [126°]. Long colour-

less needles. Sl sol cold water and ligroin, v sol alcohol (Suida, *B* 12, 1326). Converted by dilute NaOH aq into $\text{C}_6\text{H}_4(\text{NHAc})\text{CH}_2\text{CO}_2\text{H}$ [142°]

Bromo-oxindole $\text{C}_6\text{H}_4\text{BrNO}$ [176°] Formed, together with tri bromo-oxindole $\text{C}_6\text{H}_2\text{Br}_3\text{NO}$ 2aq, by the action of bromine water on oxindole. Both compounds form feathery crystals.

Nitro-oxindole $\text{C}_6\text{H}_4(\text{NO})\text{NO}$ Prepared by nitration. Yellow needles, sol alcohol. Begins to decompose at 175°.

Amido-oxindole $[\text{4 } \frac{1}{2}] \text{C}_6\text{H}_4(\text{NH}_2) \langle \frac{\text{CH}}{\text{NH}} \rangle \text{CO}$ [c 200°] Prepared by reduction of (4,2,1)-di-nitro phenyl acetic acid (Gabriel a Meyer, *B* 14, 832). Long spikes, v sol hot water.

Ezo-amido-oxindole $\text{C}_6\text{H}_4 \langle \frac{\text{CH}(\text{NH}_2)}{\text{NH}} \rangle \text{CO}$
The hydrochloride, formed by reducing isatin-oxim with tin and HCl aq , is decomposed by water, yielding a red resin.

Methyl-oxindole v p 351

Ethyl-oxindole $\text{C}_6\text{H}_4 \langle \frac{\text{CH}_2}{\text{NEt}} \rangle \text{CO}$ Formed by heating oxindole with EtI and NaOEt . Liquid, sl sol water. Very difficult to saponify by acids or alkalis.

Dioxindole $\text{C}_6\text{H}_4\text{NO}_2$ s.e. $\text{C}_6\text{H}_4 \langle \frac{\text{CH}(\text{OH})}{\text{NH}} \rangle \text{CO}$
Isatin dihydride. Anhydride of Hydronic acid. Mol w 149 [180°] S 85 in the cold, 17 at 100° S (alcohol) 7 in the cold, 10 at 78° Formed by reducing an aqueous solution of isatin with zinc dust and HCl aq , and extracting with ether (Baeyer, *B* 12, 1309). Yellowish monoclinic prisms (from water) or colourless crystals (from alcohol). Forms a violet liquid on melting. Its aqueous solution becomes red on exposure to the air, forming isatyde and isatin. In acid solution it may be reduced to oxindole. Chlorine forms chloro dioxindole $\text{C}_6\text{H}_3\text{ClNO}_2$ and dichloro dioxindole $\text{C}_6\text{H}_2\text{Cl}_2\text{NO}_2$. The corresponding bromo-derivatives melt at 165° and 170° respectively. PCl_5 yields $\text{C}_6\text{H}_4\text{Cl}_2\text{N}$. Ammonia colours its solution violet, and on boiling throws down a violet colouring matter, sol HCl aq . Potash and baryta produce a dark-violet colour, changing to red and finally yellow.

Salts — $\text{C}_6\text{H}_4\text{NO}_2\text{HCl}$ nodular crusts — $\text{C}_6\text{H}_4\text{NO}_2\text{H}_2\text{SO}_4$ aq ppd by adding water to its solution in H_2SO_4 — $\text{NaC}_6\text{H}_4\text{NO}_2$ 2aq silvery scales, v sol water, insol alcohol — BaA' 4aq white cubes, sl sol water. Gives off aniline on heating — PbA' 2aq ppd by lead subacetate — AgA' crystalline pp. Gives off benzoic aldehyde at 60°.

Acetyl derivative $\text{C}_6\text{H}_4\text{AcNO}_2$ [127°] Short prisms, sl sol cold water (Suida, *B* 12, 1326). Cold baryta water converts it into acetyl-hydronic acid $\text{C}_6\text{H}_4(\text{NHAc})\text{CH}(\text{OH})\text{CO}_2\text{H}$ [142°] which is also formed by reducing acetyl-isatinic acid with 3 p.c. sodium amalgam in acid solution (Suida, *B* 11, 586).

Nitroso-dioxindole $\text{C}_6\text{H}_4(\text{NO})\text{NO}_2$ [300°–310°] Formed by passing nitrous acid vapour into an alcoholic solution of dioxindole. Yellowish crystals, sl. sol water. May be sublimed. On boiling with FeSO_4 and KOH aq it yields 'azodioxindole' $\text{C}_6\text{H}_4\text{N}_2\text{O}_2$, which is ppd by HCl aq in white needles [300°], yielding with AgNO_3 and NH_3 a white pp of $\text{Ag}_2\text{C}_6\text{H}_4\text{N}_2\text{O}_2$. Sodium-amalgam

and a little water reduce nitroso dioxindole to 'azodioxindole' $\text{C}_6\text{H}_4\text{N}_2\text{O}_2$ ppd by HCl aq as an amorphous powder (containing 3aq) and crystallising from alcohol in cubes. Bromine water converts nitroso dioxindole into $\text{C}_6\text{H}_4\text{Br}_2\text{N}_2\text{O}_2$ 3aq [275°].

Salts — $\text{NH}_4\text{C}_6\text{H}_4(\text{NO})\text{NO}_2$ 1½aq white silky laminae — $\text{BaC}_6\text{H}_4\text{N}_2\text{O}_2$ — $\text{AgC}_6\text{H}_4\text{N}_2\text{O}_2$, yellowish-white pp.

OXOCTENOL $\text{C}_8\text{H}_{14}\text{O}_2$ s.e.

$\text{OMe}, \text{C}(\text{OH}) \langle \frac{\text{CMe}_2}{\text{O}} \rangle$ [49.5°] (178°). V D

48 (at 185°). A product of the oxidation of isodibutylene by KMnO_4 (Butleroff, *J R* 14, 203, *C J* 42, 936, *Bl* [2] 38, 553). Long thin prisms, smelling like camphor, sl sol water, v sol alcohol and ether. Does not react with hydroxylamine (Meyer a Nagel, *B* 16, 1622). Not attacked by hot bromine.

Acetyl derivative $\text{C}_8\text{H}_{12}\text{AcO}_2$ (201°) Formed by heating with Ac_2O at 150°.

OXOCTYLIC ACID v Oxy-octonic acid

OXONIC ACID $\text{C}_8\text{H}_{14}\text{N}_2\text{O}_4$ A salt of this acid is formed by the atmospheric oxidation of an alkaline solution of uric acid (Strecker, *Medicus*, A 175, 230, *B* 10, 546). The free acid splits up at once into glyoxyl urea, CO_2 , and NH_3 — $\text{NH}_4\text{HA}''$ aq — KHA'' — $\text{K}_2\text{A}''$ 1½aq — $\text{Ba}(\text{HA}'')$ ½aq minute needles.

OXYACANTHINE $\text{C}_{16}\text{H}_{11}\text{NO}_2$ Occurs in the root of *Berberis vulgaris*, together with berberine, berbamine, and at least one other alkaloid. Needles (from alcohol or ether), or amorphous solid. In the amorphous state it melts at [138°–150°], in the crystalline state at [208°–214°]. V sol benzene and chloroform, scarcely in petroleum spirit. In chloroform $[\alpha]_D = +131.6$. Conc H_2SO_4 , or conc H_2SO_4 and molybdic acid, gives no colour at first, but on standing or heating, a yellow colour. From iodic acid it separates iodine.

Salts — $\text{B}'\text{HCl}$ 2aq small colourless needles, in aqueous solution $[\alpha]_D = +163.6$ — $\text{B}'\text{HNO}_3$ 2aq, colourless needles — $\text{B}'\text{H}_2\text{SO}_4$ 2aq microscopic plates — $\text{B}'\text{H}_2\text{SO}_4$ 6aq small prisms — $\text{B}'\text{H}_2\text{Cl}_2\text{PtCl}_4$ 5aq yellow pp (Hesse, *B* 19, 3190, cf Poley, *Ar Ph* 6, 265, Wacker, *J* 1861, 545).

DIOXY-ACENAPHTHENE $\text{C}_{10}\text{H}_6 \langle \frac{\text{CH}(\text{OH})}{\text{CH}(\text{OH})} \rangle$

Acenaphthylene-glycol [205°] Formed by saponifying its acetyl derivative, which is obtained from di bromo acenaphthene $\text{C}_{10}\text{H}_6 \langle \frac{\text{CHBr}}{\text{CHBr}} \rangle$ (Ewan a Cohen, *C J* 55, 578). Long colourless needles, sl sol cold MeOH and hot water. On treatment with Na and alcohol it yields the ketone $\text{C}_{10}\text{H}_6 \langle \frac{\text{CH}_2}{\text{CO}} \rangle$ [119°].

Mono-acetyl derivative $\text{C}_{11}\text{H}_7\text{AcO}$, [112°] Long needles, v sol alcohol.

Di-acetyl derivative $\text{C}_{12}\text{H}_8\text{Ac}_2\text{O}_2$ [130°] **Benzoyl derivative** $\text{C}_{11}\text{H}_7\text{BzO}_2$ [190°]

OXY-ACETIC ACID v GLYOXYLIC ACID

Di-oxy-acetic acid v GLYOXYLIC ACID

Tri-oxy-acetic acid v OXALIC ACID.

DI-OXY-ACETOACETIC ETHER

Di-ethyl derivative $\text{C}_{16}\text{H}_{20}\text{O}_4$ s.e. $\text{CH}_3(\text{OEt})\text{COCH}(\text{OEt})\text{CO}_2\text{Et}$ (245°) Formed by the action of Na on a solution of $\text{CH}_3(\text{OEt})\text{CO}_2\text{Et}$ in benzene (Conrad, *B* 11, 58).

Liquid, gives a violet colour with FeCl_3 . Forms Na and Ba derivatives. Split up by alkalis into alcohol and $\text{CH}_3(\text{OEt})\text{CO}_2\text{H}$

OXY-ACETONE v ACETYL CARBINOL

s*-Di oxy acetone** ***Di ethyl derivative $\text{C}_6\text{H}_5\text{O}_2$, vs $\text{CO}(\text{CH}_2\text{OEt})_2$ (195°) SG 12 98 VD 4 95 Formed by allowing the ether $\text{CH}_3(\text{OEt})\text{CO CH}(\text{OEt})\text{CO}_2\text{Et}$ (v the preceding article) to stand for three days with cold dilute (2.5 p.c.) KOH, neutralising with H_2SO_4 , extracting with ether, and distilling (Grimaux & Lefèvre, C R 107, 914) Sweet aromatic liquid, sl sol Aq, sol alcohol, volatile with steam. Combines with NaHSO_3 . Reduces Fehling's solution and yields a mirror with warm ammoniacal AgNO_3 .

***u*-Di oxy acetone v PYRUVIC ALDEHYDE**

***p*-OXY ACETOPHENONE** $\text{C}_6\text{H}_4(\text{OH})\text{CO CH}_3$ [14] [107°] Obtained by diazotising *p* amido acetophenone and boiling the solution (Klingel, B 18, 2691) Formed also by heating phenol with ZnCl_2 and HOAc (Michael & Palmer, Am 7, 277) White needles Sol water, alcohol, and ether FeCl_3 gives a dark brown colouration

Methyl derivative $\text{C}_6\text{H}_4(\text{OMe})\text{CO CH}_3$ [39°] (Gattermann, B 23, 1201) (258°) (G), (221°) (O) Formed by heating anisic aldehyde with Na and MeI in ethereal solution (Oliveri, G 13, 275)

Ethyl derivative [81°] (above 260°)

o*-Oxy acetophenone v BENZOYL CARBINOL** The ***phenyl derivative $\text{Br CH}_2(\text{OPh})$ [72°] and the ***p nitro-phenyl derivative*** $\text{Bz CH}_2(\text{OC}_6\text{H}_4\text{NO}_2)$ [144°] may be obtained from *o* bromo acetophenone (Mohlan, B 15, 2497) The ***phenyl hydrazides*** $\text{C}_6\text{H}_5\text{C}(\text{N}_2\text{HPh})\text{CH}_2\text{OH}$ [112°] and $\text{C}_6\text{H}_5\text{C}(\text{N}_2\text{HPh})\text{CH}_2\text{N}_2\text{HPh}$ [152°] have been prepared by Laubmann (A 243, 247).

(4 2 1) ***Di oxy acetophenone***

[4 2 1] $\text{C}_6\text{H}_3(\text{OH})_2\text{CO CH}_3$ ***Resacetophenone*** [142°] Formed by fusion of β methyl umbelliferon with KOH (Pechmann & Dunsberg, B 16, 2123) Prepared by heating resorcin (1 pt) with HOAc (1½ pts) and ZnCl_2 (1½ pts) at 150° (Nencki & Sieber, J pr [2] 23, 147, 546) Fine white needles Gives a red colouration with FeCl_3 . With HOAc , ZnCl_2 , and POCl_3 it gives $\text{C}_6\text{H}_3(\text{OH})_2\text{Ac}$ [180°] (Crépeux, Bl [3] 6, 152)

Acetyl derivative $\text{C}_6\text{H}_3(\text{OAc})\text{CO CH}_3$ [73°] (303°) White needles

Phenyl hydrazide

$\text{C}_6\text{H}_3(\text{OH})_2\text{C(OMe)N}_2\text{HPh}$ [139°] Tables (from xylene) (Michael & Palmer, Am 7, 276)

Methyl derivative

[2 4 1] $\text{C}_6\text{H}_3(\text{OH})_2(\text{OMe})\text{CO CH}_3$ [47°] Occurs in Japanese peonies (Will, B 19, 1776)

Di ethyl derivative [68°] (G)

(5, 2, 1) ***Di oxy acetophenone*** [202°] Made in like manner from hydroquinone (Nencki)

Tri oxy acetophenone $\text{C}_6\text{H}_2\text{O}_3$, vs

$\text{C}_6\text{H}_3(\text{OH})_2\text{CO CH}_3$ ***Gallacetophenone*** [168°] Formed by heating pyrogallol (1 pt) with HOAc (1½ pts) and ZnCl_2 (1½ pts) at 150° (Nencki & Sieber, J pr [2] 23, 147, 538) Pearly plates Alcoholic KOH ppts $\text{C}_6\text{H}_2\text{O}_3\text{KOH}$

OXY-ACETOPHENONE-CARBOXYLIC ACID

Phenyl derivative $\text{C}_6\text{H}_4\text{O}_2$, vs $\text{C}_6\text{H}_4(\text{CO}_2\text{H})\text{CO CH}_2\text{OPh}$ [110°] Prepared by dissolving $\text{C}_6\text{H}_4\text{C(=O)CH}_2\text{OPh}$ in alkalis and acidifying the solution. Needles (Gabriel, B. 14, 928) —A'Ag white flocculent pp

β -OXY-ACETYL PROPIONIC ACID

$\text{C}_6\text{H}_4(\text{OH})\text{O}_2$ ***β -Oxy-levulic acid*** Formed in chief quantity, together with aceticrylic acid, by the action of aqueous Na_2CO_3 upon β -bromo-levulic acid. Yellowish oil V sol water and alcohol, sl sol other solvents Very prone to enter into reactions Reduces alkaline silver solutions Hydroxylamine gives an oxim [145°] By heating with NH_3 it yields tetra-methyl-pyrazine (di-methyl ketine), with evolution of CO_2 The salts are amorphous and easily decomposable (Wolff, B 20, 426, A 264, 234) On heating it yields two anhydrides [240°] and [263°]

The isomeric α -oxy acetyl propionic acid $\text{CH}_3\text{Ac CH}(\text{OH})\text{CO}_2\text{H}$ [104°] is crystalline

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OXY-ACRYLIC ACID v GLYCIDIC ACID and PYRUVIC ACID

OXY-ADIPIC ACID $\text{C}_6\text{H}_8\text{O}_4$, vs $\text{CO}_2\text{H CHMe CMe}(\text{OH})\text{CO}_2\text{H}$ Formed by treating methyl-acetoacetic ether with HCy and boiling the product with HClAq (König, B 12, 768) Crystalline mass — NaA'' — AgA''

Isomeric v ADIPOMALIC ACID

Di-oxy-adipic acid $\text{C}_6\text{H}_8\text{O}_4$ Formed by the action of Ag_2O on the di bromo adipic acid prepared from hydromuconic acid (v p 443) (Limpicht, A 165, 267) Syrup, sol alcohol and ether — BaA'' 4aq hygroscopic powder

Di-oxy-adipic acid Got from its nitrile, which is formed by warming the anhydride $\text{C}_6\text{H}_4\text{O}_2$ of erythrite with dry HCy at 55° (Przybytek, B 17, 1094) Crystalline — KHA'' — CdA'' 4aq — PbA'' 2aq amorphous pp

Tri-oxy-adipic acid $\text{C}_6\text{H}_6\text{O}_6$ Prepared by boiling tri bromo adipic acid with baryta water (Limpicht) Prisms (from alcohol benzene) — BaA'' 3aq v sol water

Tri-oxy-adipic acid $\text{C}_6\text{H}_6\text{O}_6$ [147°] Formed by oxidising metasaccharin with HNO_3 (SG 12) at 50° (Kiliani, B 18, 644, 1555) Small monoclinic plates, v sol water, v sl sol alcohol and ether Reduced by HI to adipic acid [149°] — CaA'' 4aq — ZnA'' 3aq — CuA'' 4aq — AgA''

Tetra-oxy-adipic acid v MUCIC ACID and SACCHARIC ACID

OXY-ALDEHYDO-BENZOIC ACID v ALDEHYDO OXY-BENZOIC ACID

OXY-ALDEHYDO-CYMENE v CARVACROTIC ALDEHYDE

OXY-ALDEHYDO-PHENOXY-ACETIC ACID

Methyl derivative

$\text{C}_6\text{H}_3(\text{OMe})(\text{CHO})\text{O CH}_2\text{CO}_2\text{H}$ [2 4 1] [188°] Prepared by melting chloro acetic acid with vanillin, adding aqueous KOH (SG 12), and heating on the water bath (Elkan, B 19, 3054) Slender needles (from water), v sol alcohol and ether — AgA'' white pp

OXY-TRI-ALDINE v ALDEHYDE

OXYAMINO-NAPHTHOQUINONE v LAFACIFIC ACID

OXY-AMIDO-ACETIC ACID $\text{C}_6\text{H}_4\text{NO}_2$, vs $\text{CH}(\text{NH}_2)(\text{OH})\text{CO}_2\text{H}$ Formed by the action of alcoholic NH_3 on glyoxylic acid (Böttger, A 198, 217) Syrup — CaA'' insol cold water. Yields pyrrole on distillation

OXY-AMIDO-ACETOPHENONE-OXIM v OX-AMIDO ACETOPHENONE-OXIM

(a) **OXY-AMIDO-ANTHRAQUINONE** $\text{C}_6\text{H}_4\text{C}_2\text{O}_2\text{C}_6\text{H}_3(\text{OH})(\text{NH}_2)$ [1 2 4 3] Formed, together with a smaller quantity of the (b) iso-

meride by heating alizarin with NH_4Aq for 8 hours at 200° (Liebermann & Troschke, *A* 183, 202) Brown needles (from alcohol) with green lustre, insol water Yields alizarin on fusing with potash, and on heating with HClAq at 250° . —BaA⁺.

Acetyl derivative $\text{C}_{14}\text{H}_9\text{O}_2(\text{OH})(\text{NHAc})$. [170°] Formed by heating with Ac_2O for 2 hours at 120° (Von Perger, *J pr* [2] 18, 143) Yellow needles (from HOAc), insol water, sol alkalis
Ethyl ether $\text{C}_{14}\text{H}_9\text{O}_2(\text{OEt})(\text{NH}_2)$ [182°] Red plates (Liebermann & Hagen, *B* 15, 1796)

(8)-Oxy-amido-anthraquinone

$\text{C}_{14}\text{H}_9\text{O}_2\text{C}_2\text{H}_4(\text{OH})(\text{NH}_2)$ [1284] Formed by boiling di-amido anthraquinone with KOHAc (Von Perger, *J pr* [2] 18, 139) Brown needles (from alcohol), insol. water, sol H_2SO_4 Begins to sublime at 150° On boiling with baryta it yields an insoluble Ba salt (difference from the (a)-isomeride) Its solution in KOHAc deposits the K salt on standing Potash fusion forms alizarin Elimination of NH_2 yields erythroxy anthraquinone

Acetyl derivative $\text{C}_{14}\text{H}_9\text{O}_2(\text{OH})(\text{NHAc})$ [242°] Small brownish needles (from alcohol)

Oxy-amido-anthraquinone

$\text{C}_{14}\text{H}_9\text{O}_2(\text{OH})(\text{NH}_2)$ [301°] Prepared by heating sodium anthraquinone sulphate with NH_4Aq at 180° (Bourchart, *B* 12, 1418) Red needles Yields the tri acetyl derivative $\text{C}_{16}\text{H}_6\text{Ac}_3\text{NO}_6$, [257°]

Oxy-amido-anthraquinone

$\text{C}_{14}\text{H}_9\text{O}_2(\text{NH}_2)(\text{OH})$ Formed by heating purpur-oxanthin with NH_4Aq (Liebermann, *A* 183, 217) Brown needles with green lustre

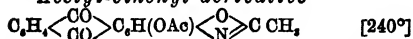
Di-oxy-amido-anthraquinone $\text{C}_{14}\text{H}_8(\text{OH})_2\text{NH}_2$

(a) **Amido alizarin** Formed by reducing (a)-nitro alizarin with sodium-amalgam (Perkin, *C J* 30, 578) Crystallises from alcohol in black needles with green lustre Its alcoholic and alkaline solutions are crimson

Di-oxy-amido-anthraquinone $\text{C}_{14}\text{H}_8(\text{OH})_2\text{NH}_2$

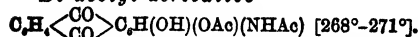
(8) **Amido alizarin** [above 800°] Formed by reducing (8) nitro alizarin with zinc dust and KOH , or with glucose and H_2SO_4 (Schunck & Roemer, *B* 12, 588, Brunner & Chuard, *B* 18, 445) Lustrous red prisms, sl sol alcohol, forming a reddish-yellow solution Its alkaline solution is blue Dyes iron mordants grey, and alumina red

Acetyl-ethenyl derivative



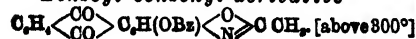
Formed by heating amidoalizarin with acetic anhydride Yellowish brown crystals (from benzene or acetic anhydride), sublimes in small yellow plates By boiling with HCl it is reconverted into amidoalizarin

Di-acetyl derivative



Formed by boiling the preceding body with dilute acetic acid till dissolved Red-brown crystals, soluble in alcohol with a yellow colour Alcoholic $\text{Pb}(\text{OAc})_2$ gives a violet pp, alcoholic $\text{Cu}(\text{OAc})_2$ a red solution It dissolves in aqueous Na_2CO_3 with a violet, in KOH with a blue, colour Alumina-mordants are dyed a deep red

Benzoyl-benzoyl derivative



Formed by heating amidoalizarin with benzoyl chloride Small glistening needles, sublimes in yellow needles, nearly insol all solvents Alcoholic KOH gives a blue solution on boiling (Roemer, *B* 18, 1666)

Di-oxy-amido-anthraquinone

$\text{C}_{14}\text{H}_9\text{O}_2(\text{OH})_2(\text{NH}_2)$ **Amidoanthraflavic acid** Formed by heating anthrapurpurin with ammonia at $150^\circ\text{--}180^\circ$ (Perkin, *C J* 33, 216) Dark-green crusts (from alcohol), almost insol water Its alkaline solution is purple Nitrous acid converts it into isoanthraflavic acid

Tetra-amido-di-oxy-anthraquinone

$\text{C}_{14}\text{H}_8\text{O}_2(\text{NH}_2)_4(\text{OH})_2$ **Hydrochrysamide** Formed by reducing tetra nitro di oxy anthraquinone (chrysammic acid) (Schunck, *A* 65, 234, Stenhouse & Miller, *A* 142, 91, Liebermann, *A* 183, 182) Bluish black needles with coppery lustre, insol boiling water, sl sol hot alcohol Its alkaline solution is blue

(a) **OXY-AMIDO-ANTHRAQUINONE SULPHONIC ACID** $\text{C}_{14}\text{H}_9\text{O}_2(\text{OH})(\text{NH}_2)\text{SO}_3\text{H}$

Formed from (a) oxy amido anthraquinone and H_2SO_4 at 180° (Von Perger, *J pr* [2] 18, 182) Brick-red crystalline aggregates (from alcohol) V sol water, giving it a reddish brown colour Insol ether Forms a violet solution with NaOH , a violet-blue pp with baryta water, and a reddish brown pp with BaCl_2 Dyes, with iron mordant, a pale brown With N_2O it gives oxy anthraquinone sulphonic acid

(8) Oxy-amido-anthraquinone sulphonic acid

Formed from amido erythro oxy-anthraquinone and H_2SO_4 at 115° Crystals obtained from water exhibit a green metallic lustre Aqueous solutions are red Insol ether Forms a purple solution with NaOH , a reddish violet pp with baryta water and a violet red pp with BaCl_2 Dyes, with iron mordants, yellow With N_2O it gives erythro-oxy anthraquinone sulphonic acid

Oxy-amido-anthraquinone sulphonic acid

Got by heating sodium anthraquinone disulphonate with NH_4Aq at 180° (Bourchart, *B* 12, 1419) Violet pp, forming a red solution in ammonia — $\text{NH}_4\text{A}^+\text{S}^{2-}\text{aq}$

Di oxy-amido-anthraquinone sulphonic acid

$\text{C}_{14}\text{H}_8(\text{OH})_2(\text{NH}_2)\text{SO}_3\text{H}$ Formed by the action of boiling alkali upon the anhydride $\text{C}_{14}\text{H}_8\text{N}_2\text{S}_2\text{O}_6$ or the sulphate, which are got by heating (a) nitro-anthraquinone sulphonic acid with H_2SO_4 at 200° (Claus, *B* 15, 1522, 16, 903, Lifschutz, *B* 17, 902) Red powder with green lustre Decomposed by heat Its alkaline solution is bluish violet

OXY-AMIDO-AZO- v Azo-

OXY-AMIDO-BENZENE v AMIDO PHENOL

Di-oxy-amido-benzene v AMIDO HYDROQUIN

ONE, AMIDO PYROCATHEIN, and AMIDO RESORCIN

Di-oxy-di-amido-benzene v DI AMIDO HYDRO

QUINONE and DI AMIDO RESORCIN

Tri-oxy-tri-amido-benzene Tri-benzoyl derivative $\text{C}_6(\text{OH})_3(\text{NHBz})_3$ [135246] **Tribenzoyl tri-amido phloroglucin** [c 156°] Got by the action of NAOEt on hippuric ether (Rugheimer, *B* 21, 8329) Needles (containing $1\frac{1}{2}\text{aq}$), v sl sol water — $\text{Cu}_2\text{A}^{3+}\text{--Pb}_2\text{A}^{3+}$

Tetra oxy amido-benzene

$\text{C}_6\text{H}_4(\text{OH})_4(\text{NH}_2)_4$ [12456] Formed from nitro di oxy quinone, SnCl_4 , and HCl (Nietzki & Schmidt, *B* 22, 1661) — $\text{B}^+\text{HCl aq}$ needles.

Penta-acetyl derivative. [242°]

Tetra-oxy di amido benzene $C_6(NH_2)_2(OH)_4$. Formed by reduction of mitranilic acid or of nitro-amido tetra oxy benzene with an excess of $SnCl_2$, the yield is 90 p.c. of the theoretical. The base could not be isolated, being readily oxidised to di-amido di oxy quinone $C_6(NH_2)_2(OH)_2O_2$. By HNO_3 it is converted into benzene-tri-quinone C_6O_3 . By boiling with KOH , NH_3 is evolved with separation of a black crystalline substance, if this is boiled with water and evaporated with a little KOH , it yields croconic acid $C_6H_2O_4$. Distillation with zinc dust yields *p* phenylene-diamine *Hydrochloride* $C_6(OH)_4(NH_2)_2H_2Cl_2$, colourless needles.

Di acetyl derivative Needles

Hexa acetyl derivative

$C_6(NHAc)_2(OAc)_4$, [c 240°], small colourless tables (Nietzki a Benckiser, *B* 18, 502, 19, 2727, 21, 1852)

OXY-AMIDO BENZOIC SULPHONIC ACID

v AMIDO PHENOL SULPHONIC ACID

Di oxy amido benzene sulphonic acid

$C_6H_4(OH)_2(NH_2)(SO_3H)$ *Amido resorcin sulphonic acid* Formed by reducing the nitro-acid (Hazura, *M* 4, 613, Brunner a Kramer, *B* 17, 1870) Plates, with greenish lustre, sl sol hot water. Its alkaline solution soon becomes blue, then green, and finally black.

OXY-AMIDO-BENZOIC ACID $C_6H_4NO_2$, *ie* $C_6H_4(OH)(NH_2)CO_2H$ [2 5 1] *Amido-salicylic acid* Mol w 153. Obtained by reducing the nitro-acid (Beilstein, *A* 130, 243, Hubner, *A* 195, 18). Obtained also by reducing $C_6H_5N_2C_6H_4(OH)CO_2H$ (P F Frankland, *C J* 37, 748) and $C_6H_5N_2C_6H_4(OH)CO_2H$ (Limpriht, *B* 22, 2908). Satiny needles, insol alcohol, sl sol hot water. $FeCl_3$ colours its aqueous solution cherry red.

Reactions—1 *Distillation* produces *p* amido-phenol—2 *Nitrous acid* forms diazo salicylic acid $C_6H_4(OH)N_2O$, which is converted by conc HI into iodo salicylic acid $C_6H_4I(OH)CO_2H$ [193 5°] (Frankland)—3 *Urea* forms, on heating, crystalline uramidosalicylic acid $C_6H_4N_2O_3$, which at 200° forms 'carboxamidosalicylic' acid $C_6H_4N_2O_4$, (Griess, *J pr* [2] 1, 235).

Salts— $HA'HCl$ — $HA'HSnCl_4$ (Goldberg, *J pr* [2] 19, 362)— $HA'HI$ (Schmitt, *J* 1864, 383, 423)— $HA'_2H_2SO_4$ aq prisms— $CaA'_2 \cdot 6\frac{1}{2}aq$ — $BaA'_2 \cdot 4aq$ Needles, v sol water— $MgA'_2 \cdot 8aq$ — $ZnA'_2 \cdot 10aq$ needles (Wattenberg, *B* 8, 1221).

Acetyl derivative $C_6H_4(OH)(NHAc)CO_2H$. [218°] Thick needles (containing $\frac{1}{2}aq$), v sol water.

Benzoyl derivative

$C_6H_4(OH)(NH_2)CO_2H$ [252°] Yields the salts $BaA'_2 \cdot 6aq$ and CaA'_2 (Dabney, *Am* 5, 22).

Oxy-amido-benzoic acid

$C_6H_4(OH)(NH_2)(CO_2H)$ [2 3 1] Formed by reducing the nitro acid (Hubner, *A* 195, 17)— $HA'HCl$ aq needles, v sol water.

Benzoyl derivative [189°] Needles

Oxy-amido-benzoic acid

$C_6H_4(OH)(NH_2)CO_2H$ [5 2 1] [235°] Formed from benzene-azo-*m*-oxy-benzoic acid and $SnCl_2$ (Limpriht, *A* 263, 234) Prisms, v. sl sol. water— $HA'HCl$ white needles.

Oxy-amido-benzoic acid

$C_6H_4(OH)(NH_2)CO_2H$ [4 8 1]. Got by reducing

nitro-*p* oxy benzoic acid (Barth, *Z* 1866, 648; Deninger, *J pr* [2] 42, 553) Needles— $H_2A'_2H_2SO_4$ needles, m sol water.

Ethyl ether EtA' Plates

Methyl derivative $C_6H_4(OMe)(NH_2)CO_2H$.

Amido amic acid [181°] S 125 at 100° Formed by reducing $C_6H_4(OMe)(NO_2)CO_2H$ (Zimin, *A* 92, 827, Cahours, *A Ch* [3] 53, 322)— AgA' curdy pp— $HA'HCl$ — $H_2A'_2H_2PtCl_6$ — $HA'HNO_3$ — $H_2A'_2H_2SO_4$ —Ethers MeA' — $(Me_4')_2H_2PtCl_6$ reddish prisms— EtA' — $EtA'HCl$ — $(EtA')_2H_2PtCl_6$ brownish-red prisms.

Oxy-amido-benzoic acid **Methyl derivative**

$C_6H_4(OMe)(NH_2)(CO_2H)$ [4 2 1] Formed from $C_6H_4Br(OMe)(NH_2)CO_2H$ by treatment with zinc and HCl (Balbiano, *G* 14, 247). It melts at 204°.

Oxy-di-amido-benzoic acid

$C_6H_4(OH)(NH_2)CO_2H$ [2 3 5 1] Formed by reducing $C_6H_4(OH)(NO_2)CO_2Me$ with HI and P (Saytzeff, *A* 133, 321). Small needles, sl sol cold water— $HA'HCl_2$ — $HA'_2H_2I_2aq$ — $HA'_2H_2SO_4$ aq dimetric prisms, sl sol water.

Di-oxy-amido benzoic acid **Di-methyl derivative** $C_6H_4(NH_2)(OMe)_2CO_2H$ Formed with evolution of CO_2 by reduction of nitro-hemic acid $C_6H_4(NO_2)(OMe)_2(CO_2H)$, with tin and HCl (Grune, *B* 19, 2305)— $A'HCl$ white needles.

Di-oxy-amido benzoic acid **Di-methyl derivative** [4 3 x 1] $C_6H_4(OMe)_2(NH_2)CO_2H$ *Amido veratric acid* Formed by reducing nitro veratric acid (Tiemann a Matsumoto, *B* 9, 942, 11, 135) Tables *Ethyl ether* EtA' [89°]

Di-oxy-amido-benzoic acid **Di-methyl derivative** $C_6H_4(OMe)_2(NH_2)CO_2H$ [5 3 4 1] [182°] Got by reduction (Meyer, *M* 8, 432). Six sided prisms (from alcohol)— $CuA'_2 \cdot 2aq$ — $HA'HCl$ needles, m sol cold water.

Di oxy-amido-benzoic acid **Acetyl-methylene derivative** of the *Nitris* $C_6H_4(O_2CH_2)(NHAc)CN$ [216°] Formed from the oxim of amido piperonal, Ac_2O , and $NaOAc$ (Haber, *B* 24, 626). Yellow needles (from chloroform).

Reference—BROMO OXY AMIDO BENZOIC ACID

DI-OXY-AMIDO-BENZOIC ALDEHYDE

Oxim of the methylene derivative $C_6H_4(O_2CH_2)(NH_2)CHNOH$ [175 5°] Got by reducing the oxim of *o*-nitro piperonal with ammonium sulphide (Haber, *B* 24, 625). Yellow plates. Yields a di acetyl derivative [188°].

OXY-AMIDO-BENZYLAMINE **Methyl derivative**

$C_6H_4(OMe)(NH_2)CH_2NH_2$ [1 2 4] Formed from $C_6H_4(OMe)(NO_2)CH_2NHAc$, tin, and $HClAq$ (Goldschmidt a Polonowska, *B* 20, 2412)— $B'HCl$ needles— $B'_2H_2PtCl_6$ plates.

Di-acetyl derivative [185°] Needles

α OXY- β AMIDO-BUTYRIC ACID $C_4H_7NO_3$, *ie* $CH_3CH(NH_2)CH(OH)CO_2H$ S 4 at 15°. Formed by heating β -methyl glycidic acid in sealed tubes with NH_3 aq at 100° (Pavloff, *Bl* [2] 43, 115, Melkoff, *J R* 16, 525). Prisms, with sweet taste.

Oxy-amido-isobutyric acid S 55 at 15°. Formed from α -methyl-glycidic acid and NH_3 .

OXY-AMIDO CINNAMIC ACID

Methyl derivative

[5 2 1] $C_6H_4(NH_2)(OMe)CHCHCO_2H$. [189°]. Formed by reduction of nitro-methoxy-cinnamic

acid with FeSO_4 and NH_3 (Schnell, *B* 17, 1884). Colourless needles $\sqrt{}$ sol alcohol and ether, almost insol cold water. An aqueous solution of the ammonium salt gives pps with AgNO_3 , CuSO_4 , $\text{Pb}(\text{OAc})_2$, and ZnSO_4 .

Anhydride $\text{C}_6\text{H}_3(\text{NH}_2) < \begin{smallmatrix} \text{O} & \text{CO} \\ \diagdown & \diagup \\ \text{CH} & \text{CH} \end{smallmatrix} \text{Amido-coumarin}$ [170°] (Frapolli a Chiozza, *A* 95, 253), [161°] (Taege, *B* 20, 2110). Got by reduction of nitro coumarin— $\text{B}_2\text{H}_6\text{PtCl}_6$.

Di-oxy-amido cinnamic acid *Methylene derivative*.

$\text{CH}_2 < \begin{smallmatrix} \text{O} \\ \diagdown \\ \text{O} \end{smallmatrix} > \text{C}_6\text{H}_3(\text{NH}_2) \text{CH CH CO}_2\text{H}$. [207°]

Formed by reducing the nitro-acid with NH_3 and hot aqueous FeSO_4 (F M Perkin, *C. J* 59, 158). Brownish needles, $\sqrt{}$ sol HOAc .

DI OXY-AMIDOETHYL BENZOIC ACID

Methylene derivative $\text{C}_{12}\text{H}_{11}\text{NO}_4$, $\text{v.e. C}_6\text{H}_5(\text{O} \cdot \text{CH}_2)(\text{CH}_2 \text{CH}_2 \text{NH}) \text{CO}_2\text{H}$. *Amido ethylpyrperonylcarboxylic acid* [182°] Formed, together with hemipic acid, by boiling anhydro berberic acid with water (W H Perkin, jun, *C. J* 57, 1055). Needles, sl sol alcohol, m sol cold water— $(\text{HA}')_2\text{H}_2\text{SO}_4$ 2aq [203°] Colourless prisms— $\text{HA}'\text{HCl}$ — $\text{HA}'\text{H}_2\text{C}_2\text{O}_4$ [203°]— $(\text{HA}')_2\text{H}_2\text{PtCl}_6$ [222°]— $\text{HA}'\text{H}_2\text{AuCl}_3$.

Di-benzoyl derivative $\text{C}_{22}\text{H}_{19}\text{NO}_6$ [150°]

Anhydride $\text{CH}_2 < \begin{smallmatrix} \text{O} \\ \diagdown \\ \text{O} \end{smallmatrix} > \text{C}_6\text{H}_4 < \begin{smallmatrix} \text{O} \\ \diagdown \\ \text{O} \end{smallmatrix} \text{CH}_2$

Formed by heating the acid, and, along with its ψ opianate and ψ opianic acid by the action of alkalis on berberal. It is also a product of the oxidation of berberine with KMnO_4 . Yields the compound $\text{C}_{10}\text{H}_7\text{BrNO}_3$ [240°]—Opianate $\text{B}'\text{C}_6\text{H}_5(\text{OMe})_2\text{CO}_2\text{H}_2$ [182°]— ψ Opianate $\text{C}_6\text{H}_5\text{NO}_3$ [149°] Needles, sl sol cold water. Yields berberal when heated.

OXY AMIDO GLUTAMIC ETHER $\text{C}_8\text{H}_{11}\text{N}_2\text{O}_4$

$\text{v.e. CO}(\text{NH}_2) \text{CH}_2 \text{C}(\text{OH})(\text{NH}_2) \text{CH}_2 \text{CO}_2\text{Et}$ [86°] Formed from acetone dicarboxylic ether and conc NH_3Aq (Stokes a Von Pechmann, *B* 19, 2694). Needles, sl sol cold water, insol alkalis. FeCl_3 gives a red colour. Dilute HClAq yields a body melting at 61° turned purple by FeCl_3 . Nitrous acid yields a nitroso derivative [178°]. Boiling alkaline carbonates yield glutazone (*v. Di OXY AMIDO PYRIDINE*).

OXY AMIDO-HEPTOIC ACID $\text{C}_8\text{H}_{11}\text{NO}_4$, $\text{v.e. CMe}_2(\text{NH}_2) \text{CH}_2 \text{CMe}(\text{OH}) \text{CO}_2\text{H}$

Amido trimethylbutylactic acid [210°] Formed by the action of boiling baryta water on its anhydride which is obtained by the action of boiling conc HClAq upon the product of the union of HCy with diacetoneamine (Heintz, *A* 189, 231, 192, 329, Weil, *A* 232, 208). Prisms (from water), insol alcohol and ether. Neutral to litmus. Yields its anhydride when heated—Salts $\text{CuC}_6\text{H}_7\text{NO}_4$ 2aq minute greenish blue prisms— $(\text{HA}')_2\text{H}_2\text{SO}_4$ — $\text{HA}'\text{HCl}$ crystalline.

Anhydride $\text{CMe}(\text{OH}) < \begin{smallmatrix} \text{CO} & \text{NH} \\ \diagdown & \diagup \\ \text{CH}_2 & \text{CMe}_2 \end{smallmatrix} \text{Di oxy- tri-methyl-pyrrole}$ Prisms (by sublimation) Melts above 180° .

OXY-AMIDO HYDRO ISATIN *v. ISATIN*

OXY-AMIDO-IMIDO-DI PHENYL SUL.

PHIDE $\text{C}_{12}\text{H}_9\text{N}_2\text{SO}$ $\text{v.e. N} < \begin{smallmatrix} \text{C}_6\text{H}_4(\text{NH}_2) \\ \diagdown \\ \text{O}-\text{C}_6\text{H}_4 \end{smallmatrix} \text{S}$.

Thionolins Formed by adding H_2S and FeCl_3 successively to an aqueous solution of *p*-

amido phenol hydrochloride (Bernthsen, *A* 280, 202). Yellowish brown plates or needles with green lustre (from alcohol).

OXY-AMIDO-DI-ISATIN DIAMIDE ISATIN DI-OXY-DI AMIDO-TETRAMETHENYL

Di-benzoyl derivative $\text{C}(\text{NHBz}) \text{C}(\text{OH}) \text{C}(\text{OH}) = \text{C}(\text{NHBz})$

[138°] Formed by heating hippuric ether with NaOEt , or with Na , at 160° (Rugheimer, *B* 21, 3325, 22, 114). Needles, m sol alcohol, $\sqrt{}$ e sol benzene. FeCl_3 colours its alcoholic solution violet. Decomposed by acids into benzoic acid and di amido acetone. Hydr xylamine does not affect it. Methyl alcohol and gaseous HCl form a body $(\text{C}_{11}\text{H}_{10}\text{N}_2\text{O}_2?)$ crystallising in plates [200°].

Oxy-tetra amido-pentamethenyl hydride

$\text{CH}(\text{OH}) < \begin{smallmatrix} \text{C}(\text{NH}_2) & \text{C}(\text{NH}_2) \\ \diagdown & \diagup \\ \text{C}(\text{NH}_2) & \text{C}(\text{NH}_2) \end{smallmatrix} \text{Tetra amido pentol}$

Formed from the tetra oxim of leuconic acid SnCl_4 , and HClAq (Nietzki a Rosemann, *B* 22, 923)— B^{14}HCl octahedra— $\text{B}^{14}(\text{H}_2\text{SO}_4)_2$ aq.

Tetra oxy-tetra-amido octomethenyl

Tetra benzoyl derivative $\text{C}_{24}\text{H}_{18}\text{N}_8\text{O}_8$, $\text{v.e. C}(\text{OH}) < \begin{smallmatrix} \text{C}(\text{NHBz}) & \text{C}(\text{OH}) & \text{C}(\text{NHBz}) \\ \diagdown & \diagup & \diagdown \\ \text{C}(\text{OH}) & \text{C}(\text{OH}) & \text{C}(\text{NHBz}) \end{smallmatrix} > \text{C}(\text{OH})$

A product of the action NaOEt on hippuric ether (Rugheimer, *B* 22, 1962). Small yellow needles. Does not melt below 270° — $\text{BaC}_2\text{H}_3\text{N}_2\text{O}_4$ pp.

Anhydride $\text{C}_8\text{H}_8\text{N}_4\text{O}_4$

OXY-AMIDO-METHYL-ANTHRAQUINONE

$\text{C}_{12}\text{H}_{11}\text{NO}_4$, $\text{v.e. C}_6\text{H}_4\text{Me}(\text{OH})(\text{NH})\text{O}$. Formed by heating chrysophanic acid with NH_3Aq at 200° (Liebermann, *A* 183, 218). Brown plates.

Oxy-di-amido-methyl-anthraquinone

$\text{C}_{12}\text{H}_{11}\text{N}_2\text{O}_5$, *Di amido chrysophanic acid* Formed by heating chrysophanic acid with excess of NH_3Aq at 150° . With AcO it yields $\text{C}_{12}\text{H}_{11}\text{AcN}_2\text{O}_5$ crystallising from chloroform in lustrous violet needles.

OXY DIAMIDO METHYL-DIPHENYL

Ethyl derivative

$\text{C}_6\text{H}_4(\text{NH}_2) \text{C}_6\text{H}_4\text{Me}(\text{OEt})(\text{NH}_2)$ [1 2 5 4] [107°] Formed from the hydrazo derivative $\text{C}_6\text{H}_4\text{NH NH C}_6\text{H}_4\text{Me}(\text{OEt})$ by dissolving in cold HClAq (Noelting a Werner, *B* 23, 3263). Needles, $\sqrt{}$ sl sol water.

Oxy di amido methyl diphenyl [3 4 1]

$\text{C}_6\text{H}_4\text{Me}(\text{NH}_2) \text{C}_6\text{H}_4(\text{OH})(\text{NH}_2)$ [1 3 4] [177°] Formed by the action of water at 180° on the sulphonic acid $\text{C}_6\text{H}_4\text{Me}(\text{NH}_2) \text{C}_6\text{H}_4(\text{OH})(\text{NH})\text{SO}_3\text{H}$ [1 3 4 6], got by reducing the dye from diazotised *o* toluidine and *p* phenol sulphonic acid (Weinberg, *B* 20, 3174). Plates, $\sqrt{}$ sl sol water.

Ethyl derivative

$\text{C}_6\text{H}_4\text{Me}(\text{NH}_2) \text{C}_6\text{H}_4(\text{OEt})(\text{NH}_2)$ [117 5°] Formed in like manner from $\text{C}_6\text{H}_4\text{Me}(\text{OEt})(\text{NH})\text{SO}_3\text{H}$, which yields $\text{HA}'\text{HCl}$ 4aq and BaA' 8aq.

OXY-AMIDO-METHYL QUINOLINE

$\text{C}_8\text{H}_7(\text{OH})(\text{NH}_2)\text{MeN}$ Formed by reducing nitro (*Py* 1)-oxy (*Py* 3) methyl quinoline (Conrad a Limpach, *B* 20, 950). Prisms, decomposing at 225° without melting— $\text{B}'\text{HCl}$ aq.

OXY AMIDO NAPHTHALENE v. AMIDO-NAPHTHOL

(1,2,3)-Di oxy amido naphthalene

$\text{C}_6\text{H}_4 < \begin{smallmatrix} \text{C}(\text{OH}) & \text{C}(\text{OH}) \\ \diagdown & \diagup \\ \text{N} & \text{C}(\text{NH}_2) \end{smallmatrix} \text{Di-oxy naphthylamine}$ Obtained by reduction of nitro (β) naphtho-

quinone (Korn, *B* 17, 906) Reduces cold AgNO_3 . Fe_2Cl_6 gives a blue black pp.

(2,2',1)-di oxy amido naphthalene

$\text{C}(\text{OH})\text{CH}=\text{C}(\text{NH}_2)\text{C}(\text{OH})$. Formed by reducing the (a) oxim of (2') oxy (β) naphthoquinone with SnCl_4 (Clausius, *B* 23, 521) — B^{HCl} needles, turning blue in air

Di-oxy-amido-naphthalene Got by reducing amido (a)-naphthoquinone (Graebe & Ludwig, *A.* 154, 320) Tables, v. e sol water Blackens in moist air

Tri-oxy amido naphthalene

$\text{C}_6\text{H}_4\text{C}(\text{OH})\text{C}(\text{NH}_2)\text{C}(\text{OH})$ Formed by reducing nitro oxy (a) naphthoquinone with SnCl_4 (Kehrmann, *B* 21, 1780, *J pr* [2] 40, 179) Monoclinic crystals Cold conc HNO_3 appears to form $\text{C}_6\text{H}_4\text{C}(\text{CO})\text{C}(\text{NO})\text{CO}$ crystallising in red dish yellow needles Yields a tetra acetyl derivative [145°]

OXY AMIDO NAPHTHALENE SULPHONIC ACID v AMIDO NAPHTHOL SULPHONIC ACID

Oxy di amido naphthalene sulphonic acid $\text{C}_6\text{H}_4(\text{NH}_2)_2(\text{OH})(\text{SO}_3\text{H})$ [α 4' 14] The hydrochloride B^{HCl} is formed by reducing di-nitro naphthol sulphonic acid with SnCl_4 and HCl (Nietzki & Zubelen, *B* 22, 455) It crystallises in colourless needles

OXY m AMIDO NAPHTHOIC ACID $\text{C}_{10}\text{H}_7(\text{OH})(\text{NH})\text{COH}$ Formed by reducing the nitro acid (Schmitt & Burkard, *B* 20, 2700) Decomposes above 200°

Acetyl derivative [185°]

Oxy p amido naphthoic acid Got by reducing $\text{C}_6\text{H}_4(\text{SO}_3\text{H})\text{N}^-\text{C}_{10}\text{H}_7(\text{OH})\text{COH}$ with tin and HCl (S & B) Decomposes above 200°

Acetyl derivative [195°].

Oxy amido naphthoic acid

$\text{C}_{10}\text{H}_7(\text{OH})(\text{NH}_2)(\text{CO}_2\text{H})$ [1 4 2 ?] [above 200°] Formed from (a) oxy naphthoic acid by combining it with diazobenzene chloride and warming the product with SnCl_4 and HCl (Nietzki & Gutermann, *B* 20, 1275) Crystalline powder, insol water Decomposes above 230° into CO_2 and (a) amido (a) naphthol

OXY AMIDO (a)-NAPHTHOQUINONE

$\text{C}_6\text{H}_4\text{C}(\text{CO})\text{C}(\text{NH}_2)\text{C}(\text{OH})$ Amido-naphthalic acid

Formed by reduction from nitro oxy naphthoquinone (Diehl & Merz, *B* 11, 1319) and from di nitroso di oxy naphthalene (Kostanecki, *B* 22, 1346) Formed also by oxidising tri-oxy amido-naphthalene with aqueous FeCl_3 (Kehrmann, *B* 21, 1781) Red prisms, sl sol water Its alkaline solutions are blue — BaA_2 : violet pp — AgA' : grey pp

Acetyl derivative $\text{C}_{10}\text{H}_7\text{O}_2(\text{OH})(\text{NHAc})$ [220°] Formed from $\text{C}_{10}\text{H}_7(\text{OAc})_2(\text{NHAc})$ and KOHAq (Kehrmann & Weichardt, *J pr* [2] 40, 182) Yellow needles, insol water — KA' : blue-black needles

Oxym $\text{C}_6\text{H}_4\text{C}(\text{CO})\text{C}(\text{NH}_2)\text{C}(\text{OH})$. Greenish flocculent pp — B^{HCl} yellow needles.

Acetyl derivative of the oxym

$\text{C}_6\text{H}_4\text{C}(\text{CO})\text{C}(\text{NHAc})\text{C}(\text{OH})$ Golden needles, sl sol alcohol Decomposes at 190°–200°.

OXY DIAMIDO DIPHENYL $\text{C}_{12}\text{H}_{10}\text{N}_2\text{O}$ i.e.

[4 1] $\text{C}_6\text{H}_4(\text{NH}_2)_2\text{C}_6\text{H}_4(\text{NH}_2)(\text{OH})$ [1 4 2] [185°] Formed by the action of water at 180° on the hydrochloride of its sulphonic acid, which is got by reducing $\text{C}_6\text{H}_4\text{N}_2\text{C}_6\text{H}_4(\text{OH})\text{SO}_3\text{H}$ with SnCl_4 at 80° (Weinberg, *B* 20, 3173, Noeltling & Werner, *B* 23, 3256) Plates, sl sol water — $\text{HA}'\text{H}_2\text{SO}_4$ insol water — $\text{HA}'\text{HCl}$ — $(\text{HA}')_2\text{H}_2\text{PtCl}_6\text{aq}$ sl sol water

Ethyl derivative

$\text{C}_6\text{H}_4(\text{NH}_2)_2\text{C}_6\text{H}_4(\text{NH}_2)(\text{OEt})$ [135°]. Needles, sl sol water — $\text{HA}'\text{H}_2\text{SO}_4$ prisms

Di-oxy-di-amido-diphenyl Di-ethyl-derivative

[4 3 1] $\text{C}_6\text{H}_4(\text{NH}_2)(\text{OEt})\text{C}_6\text{H}_4(\text{NH}_2)(\text{OEt})$ [1 4 3] Di-ethoxy benzidine [117°] Formed from [2 1] $\text{C}_6\text{H}_4(\text{OEt})\text{NH NH C}_6\text{H}_4(\text{OEt})$ [1 2] and conc HClAq (Mohlau, *J pr* [2] 19, 381) Needles or plates, sol hot water — B^{HCl} — $\text{B}^{\text{H}_2\text{SO}_4}$ — $\text{B}^{\text{H}}\text{PtCl}_6$ — $\text{B}^{\text{H}}\text{H}_2\text{SO}_4$ needles, sol hot water

Di oxy di amido diphenyl

[3 4 1] $\text{C}_6\text{H}_4(\text{NH}_2)(\text{OH})\text{C}_6\text{H}_4(\text{NH}_2)(\text{OH})$ [1 3 4] [above 300°] Formed by reducing di nitro di-oxy diphenyl (Kunze, *B* 21, 3332, Schutz, *B* 21, 3531) Needles or plates AcO yields the di-acetyl derivative AcCl forms the di ethenyl derivative [195°] — B^{HCl} needles — $\text{B}^{\text{H}_2\text{SO}_4}$

Di-acetyl derivative

$\text{C}_{12}\text{H}_8(\text{OH})(\text{NHAc})_2$ [210°] Needles, sol alkalis, insol acids

Tetra-acetyl derivative $\text{C}_{12}\text{H}_6\text{Ac}_4\text{N}_2\text{O}_2$ [225°] White needles (K)

Di oxy-tetra amido diphenyl

$\text{C}_6\text{H}_4(\text{NH}_2)_2(\text{OH})\text{C}_6\text{H}_4(\text{NH}_2)(\text{OH})$ Formed from tetra nitro di oxy diphenyl (Kunze) Colourless needles or silvery plates, turning blue in air AcCl converts it into a compound $\text{C}_{12}\text{H}_8\text{N}_2\text{O}_2$ — B^{HCl} 4aq — $\text{B}^{\text{H}}\text{H}_2\text{SO}_4$ white needles

Tetra acetyl derivative

$\text{C}_{12}\text{H}_8(\text{NHAc})_2(\text{OH})_2$ [280°] Got by boiling the hexa acetyl derivative with NaOHaq

Hexa acetyl derivative

$\text{C}_{12}\text{H}_8(\text{NHAc})_2(\text{OAc})_2$ [300°] Formed from the base and boiling AcO

Tetra oxy di amido diphenyl

$\text{C}_6\text{H}_4(\text{OH})_2(\text{NH})\text{C}_6\text{H}_4(\text{OH})\text{NH}_2$

Tetra-methyl derivative

$\text{C}_{12}\text{H}_8(\text{OMe})_2(\text{NH}_2)_2$ [210°] Formed by molecular change from tetra methoxy hydrazobenzene under the influence of acids (Baessler, *B* 17, 2123) Silky white needles, v sol chloroform, CS_2 , hot alcohol, and hot benzene, sl sol water and ligroin — B^{HCl} small white, easily soluble needles — $\text{B}^{\text{H}}\text{Cl}_2\text{PtCl}_6$ yellow pp The di-acetyl derivative forms white needles [251°], sol hot alcohol, benzene, chloroform, and CS_2

Tetra-ethyl derivative

$\text{C}_{12}\text{H}_8(\text{NH}_2)_2(\text{OEt})_2$ [129°] Formed, in like manner, from the hydrazo-compound got by reducing the di ethyl ether of nitro hydroquinone with powdered zinc and alcoholic potash (Nietzki, *B* 12, 39) Leaflets — $\text{B}^{\text{H}}\text{Cl}_2$ — $\text{B}^{\text{H}}\text{H}_2\text{PtCl}_6$ yellow crystalline pp

Tetra-oxy tetra-amido-diphenyl Tetra-acetyl derivative $\text{C}_{12}\text{H}_8(\text{NHAc})_4(\text{OH})_4$ White needles, formed by reduction of the corresponding quinone $\text{C}_6\text{H}_4(\text{NHAc})_2(\text{OH})_2\text{O}_2$ [268°] which is formed by oxidation of tri-acetyl tri-amido phenol (Bamberger, *B* 16, 2403)

• OXY-*o*-AMIDO-PHENYL-ACETIC ACID*Hydrazide acid**Acetyl derivative*

$C_6H_4(NHAc)CH(OH)CO_2H$ [142°] Got by reducing acetyl isatic acid with sodium amalgam in presence of acetic acid (Suida, *B* 11, 586) Needles, v sol water

Anhydride v Di oxindole

p-Oxy-*o*-amido-phenyl acetic acid *Methyl derivative* $C_6H_4(OMe)CH(NH)CO_2H$ [225°] Formed from anisic aldehyde by successive treatment with HCy, alcoholic NH_3 , and HCl (The mann & Köhler, *B* 14, 1979). Needles (from dilute alcohol) — CuA' , blue amorphous pp

Di-oxy o amido-phenyl-acetic acid *Methyl-ene derivative* $CH_2O_2C_6H_4CH(NH_2)CO_2H$ [210°] Formed from piperonal by successive treatment with HCy, alcoholic NH_3 , and HCl (Lorenz, *B* 14, 794) Needles, sl sol water

OXY-AMIDO-PHENYL-ACRIDINE v CHERYSPHENOL**OXY-AMIDO DI PHENYL-AMINE**

$C_6H_5NH C_6H_4(OH)(NH_2)$ [134°] [135°] Formed by reducing nitroso-oxy di phenyl amine (Köhler, *B* 21, 910) Plates, v sol chloroform

OXY-AMIDO-PHENYL CARBAMICETHERS

Formed from the nitro compounds (Köhler, *J pr* [2] 29, 263)

[431] $C_6H_4(OEt)(NH_2)NHCO_2Et$ [88°] Small needles — $B'HCl$ [156°] Needles

[432] $C_6H_4(OEt)(NH_2)NHCO_2Et$ Needles — $B'HCl$ [238°] Six sided plates, sol water

$C_6H_4(OEt)(NH_2)NHCO_2Et$ Small needles, rapidly oxidising in air — $B'HCl$ [233°] Needles

OXY-AMIDO-DI-PHENYL-METHANE v. AMIDO BENZYL-PHENOL

Oxy di amido-tri-phenyl-methane Methyl derivative $C_6H_4(OMe)CH(C_6H_5NH_2)_2$ Formed by heating anisic aldehyde with aniline and $HClAq$ (Mazzara & Possetto, *G* 15, 57) Crystalline crust (containing C_6H_5) melting at 65°

Di-oxy-tri-amido-tri-phenyl-methane

Di-methyl derivative $C_6H_4(NH_2)CH(C_6H_5NH_2)OMe_2$ [183°] Obtained by reduction of the di methyl derivative of nitro di-amido-di-oxy tri-phenyl methane (Fischer, *B* 15, 681) Plates or tables Yields on oxidation 'rosaniline,' a bluish red colouring matter with blue fluorescence

OXY-AMIDO-PHENYL-METHYL-PYRAZOLE

$C_6H_4N_2O$ *ie* $C_6H_4N_2COCH(NH_2)Me$ Formed by reducing the nitro compound (Knorr, *A* 238, 189) Rapidly oxidised by air — $B'HCl$

OXY DI-AMIDO PHENYL-METHYL-PYRIMIDINE

$C_{11}H_8N_4O$ [232°–240°] Formed from $C_6H_5CH=N C(OH)Me$ by nitration and reduction (Pinner, *B* 20, 2364) Golden needles, sol $NaOHaq$ — $B'H_2PtCl_4$ — $B'H_2I_2$ prisms

OXY-DI-AMIDO-PHENYL NAPHTHALENE

Ethyl derivative [214] $C_{10}H_6(OEt)(NH_2)C_6H_4NH_2$ [14] [72°] Formed by reducing $C_6H_5N_2C_6H_4OEt$ in alcoholic solution with $SnCl_2$ and HCl (Wernberg, *B* 20, 3177) White flakes — $B'HCl$ — $B'H_2SO_4$, sl sol water.

OXY-AMIDO-PHENYL PROPIONIC ACID

$C_6H_4CH(NH_2)CH(OH)CO_2H$ [190°] (P) Formed from sodium phenyl glycidate and NH_3 , (Plöchl, *B* 16, 2822, Erlenmeyer, jun, *B* 22, 1482) Decomposes at 221° (E)

o Oxy-*o*-amido phenyl propionic acid.

Anhydride $C_6H_4CH_2CH(OH)NHCO$. *Oxyhydro-*

carbostyryl [198°] Got by reducing the product of the nitration of *o*-oxy phenyl propionic acid (Erlenmeyer & Lipp, *A* 219, 229) White plates (from alcohol) Sl sol ether, v² sol hot water

o Oxy-*p*-amido-phenyl-propionic acid

$C_6H_4(NH_2)CH_2CH(OH)CO_2H$ [189°] Got by reducing the nitrate of nitro oxy phenyl-propionic acid (Erlenmeyer & Lipp, *A* 219, 227) Slender needles (containing $\frac{1}{2}$ aq) from dilute (93 p c) alcohol More sol alcohol or ether than tyrosine Readily soluble in alkalis or acids Unlike tyrosine, its solution is acid and it does not give Piria's reaction Boiled with $Hg(NO_3)_2$ it forms a yellow flocculent pp, which, on adding HNO_3 , turns red $HA'HCl$ V sol. water or alcohol, without decomposition

p Oxy-*p* amido phenyl propionic acid v.**TYROSINE***o*-Di-oxy-*o*-amido phenyl propionic acid

$C_6H_4(NH_2)CH(OH)CH(OH)CO_2H$ [218°] Produced in the reduction of *o* nitro cinnamic acid by tin and $HClAq$ (Morgan, *J* 1877, 788) Yellow needles (from alcohol) Its solutions fluoresce green

Di oxy amido propionic acid Anhydride of the methylene derivative

$CH_2<O>C_6H_4<NHCOCH_2CH_2$ 'Ethylamidopiperonyl *o* carboxylic anhydride' [235°] Formed by reducing $(CH_3O)_2C_6H_4(NO_2)CHCHCO_2Et$ in alcoholic solution with tin and gaseous HCl (F M Perkin, *C J* 59, 159) Crystalline pp

OXY-AMIDO PHENYL PYRAZOLE CARB- OXYLIC ACID

Benzoyl derivative $NPh<COCHNHBz>N=C O_2H$ [185°–190°] Formed from its ether and $NaOH$ (Wilschusen, *B* 24, 1261) Yellowish needles, sl sol water

Ethyl ether EtA' [195°] Formed by boiling the phenyl hydrazide of benzoyl amido-oxalacetic ether with $HOAc$ Yellow crystals

OXY m-AMIDO PHENYL-PYROTARTARIC ACID

Anhydride $C_{11}H_8N_4O_{12}$ *ie* $C_6H_4(NH_2)CH<CH(CO_2H)>CO_2CH_2$ *Amidophenyl-*

paraconic acid Formed by reducing the corresponding nitro compound (Salomonson, *R T C* 6, 18) — $B'HCl$ prisms — $B'H_2PtCl_4$. The isomeric oxy *p* amido phenyl pyrotartaric anhydride yields a similar hydrochloride

(B 2) **OXY-*p*-AMIDO-(Py 3)-PHENYL-**

QUINOLINE $C_6H_4CH(OH)CH_2CH_2CH_2NH_2$ $CH=CHON C_6H_4NH_2$

[294°] Formed by passing oxygen over a mixture of (B 2)-oxy-quinoline hydrochloride, aniline hydrochloride, aniline, and platinised asbestos at 220° (Weidel & Georgievitch, *M* 9, 146) Needles (from amyl alcohol), sl sol. alcohol and ether — $B'HCl$ $\frac{1}{2}$ aq — $B'H_2SO_4$ $\frac{1}{2}$ aq

Acetyl derivative $C_{11}H_{10}Ac_2N_2O$. Scales.

(Py 412) Oxy amido phenyl-isoquinoline

$C_{17}H_{12}N_2O$ i.e. C_6H_5 $\begin{matrix} \text{C(NH)} \text{CPh} \\ | \\ \text{C(OH)N} \end{matrix}$. Amido iso-
benzal pthalimidine [c 190°] Formed by
redn of the nitro-derivative with P and HI
(Gabriel, B 19, 833) Fine yellow needles
v sol acetic acid and hot alcohol

DI-OXY-DI AMIDO-DI-PHENYL SUL-

PHONE $C_{17}H_{12}N_2SO_4$ i.e. $SO_2(C_6H_4(NH_2)OH)_2$
Formed by reducing the nitro-compound (Anna-
heim, B 7, 436, 8, 1063) Crystals (from
water) Alcoholic isomyl nitrite forms golden
plates of $C_{17}H_{12}N_2SO_4$ Salts— $B''H_2Cl_2$ 2aq—
 $B''H_2L_2$ 2aq— $B''H_2SO_4$ 2aq thick prisms—
 $C_2H_5Me_2N_2SO_4H_2$ Forms long needles—
 $C_{17}H_{12}Et_2N_2SO_4H_2$ needles

OXY-DI-AMIDO-DIPHENYL SULPHONIC

ACID [41] $C_{12}H_8(NH_2)_2C_6H_4(OH)SO_3H$
[1436] Prepared by reducing the azo-compound
 $C_6H_5N=N_2C_6H_4(OH)SO_3Na$ with aqueous
 $SnCl_2$ (Weinberg, B 20, 3172, Feer & Muller,
C C 1888, 1358) Needles, v sol water— $B'HCl$
transparent crystals

Ethyl derivative

$C_6H_4(NH_2)_2C_6H_4(NH_2)(OEt)SO_3H$ Got in like
manner Needles— $B'HCl$ 2aq needles, v sol
water

OXY-AMIDO-DIPHENYL DISULPHONIC

ACID $C_6H_4(NH_2)(SO_3H)C_6H_4(OH)(SO_3H)$
Formed from benzidine disulphonic acid by the
diaz reaction (Limpriht, A 261, 315) Light-
yellow crystalline mass, v e sol water, insol
ether— BaA'' 8aq reddish-yellow crystals

OXY-AMIDO PHENYL TOLYL KETONE

$C_{17}H_{12}NO_2$ i.e. $C_6H_4(NH)CO C_6H_4Me(OH)$ or
 $C_6H_4(OH)CO C_6H_4Me(NH_2)$ A product of the
action of water at 270° on commercial rosaniline
(Liebemann, B 16, 1927) Small colourless
needles, sol acids and alkalis

Di-benzoyl derivative [193°] Needles

DI OXY-AMIDO-PHTHALIC ACID Methyl
derivative of the anhydride

$C_6H_4(OMe)(OH)(CO_2H) \begin{matrix} \text{CO} \\ | \\ \text{NH} \end{matrix}$ [4321] Nor-
methylazoopropionic acid [175°] Formed by

reducing $C_6H_4(NO_2)(OMe)(OH)(CHO)CO_2H$ with
conc $SnCl_2$ and HCl (Elbel, B 19, 2307)
Colourless needles, nearly insol ether Yields
an acetyl derivative $C_6H_4NO_2(OMe)(OAc)$ [198°],
and a di acetyl derivative [105°]

Di-methyl derivative of the anhy-
dride v Amido hemipic acid, vol 11 p 672

DI-OXY-AMIDO-PHTHALIDE Di-

methyl derivative v Amido MECONIN (p
198) and Amido ψ MECONIN (p 199)

 α -OXY β AMIDO PROPIONIC ACID

$CH(NH)CH(OH)CO_2H$ Amido lactic acid
b 15 at 20° Formed from $CHClCH(OH)CO_2H$
and conc NH_4Aq (Melikoff, C C 1881, 354, B
13, 958, 1266, Erlenmeyer, B 13, 1077) Mono-
clinic prisms— $B'HCl$ needles

OXY AMIDO-ISOPROPYL BENZOIC ACID

$OMe_2(OH)C_6H_4(NH_2)CO_2H$ [421] [158°] Got
from the nitro-acid (Widman, B 19, 271)
Prisms Yields amido-propenyl benzoic acid
on boiling with $HClAq$

Acetyl derivative [174°] Tables

Exo oxy amido isopropyl benzoic acid
 $OMe_2(OH)C_6H_4(NH_2)(CO_2H)$ [431] Does not

melt below 270° (Widman, B 16, 2571) Its
acetyl derivative is not melted at 280°

OXY-AMIDO PYRIDINE $C_6H_4N(OH)(NH_2)$
[214°] Got by reducing oxy comazine (Krippen-
dorff, J pr [2] 32, 162) Pyramids (containing
aq), v e sol hot water— $B'HCl$ — $B''H_2PtCl_4$

Di-oxy-amido-pyridine $NH \begin{matrix} \text{CO} CH_2 \\ | \\ \text{CO} CH_2 \end{matrix} C NH$

or $N \begin{matrix} \text{C(OH)CH} \\ | \\ \text{C(OH)CH} \end{matrix} C NH_2$ Glutazime [c 300°].

Formed by boiling β -oxy- β amido glutamic ether
 $CO_2EtCH_2C(OH)(NH_2)CH_2CONH_2$ with aque-
ous Na_2CO_3 (Stokes & Von Pechmann, B 19,
2694, 20, 2655, Am 8, 375) Rectangular
plates, m sol hot water, almost insol hot alcoh-
ol Gives a deep red colour with $FeCl_3$, turning
dark green on warming

Reactions—1 Hot $HClAq$ yields tri-oxy-
pyridine—2 Bromine in excess forms
 CBR_3 , $CO CBR_2$, $CONH_2$ [c 148°]—3 By heating
with PCl_5 (6 to 7 pts) it is converted into a
mixture of tri chloro oxy-amido pyridine [282°]
and tetra chloro amido-pyridine [212°], together
with small quantities of di chloro di oxy amido-
pyridine [242°] and tri chloro amido pyridine
[158°]—4 Yields nitro glutazime [170°–180°]
and crystalline di nitro glutazime on treatment
with nitrous acid gas With $NaNO_2$ and acetic
acid there is formed the nitro nitrosamine
 $C_6H_4N_2O_2(NO_2)(NO)$ whence $NaC_6H_4N_2O_2$ crys-
tallising in yellow needles, which on warming
with $NaNO_2$ and $HOAc$ yields the di nitro-
nitrosamine salt $NaC_6H_4N_2O_2$, a scarlet powder
Salts— $B'HCl$ aq prisms, v sol alcohol—
 AgA' plates

Acetyl derivative $C_6H_4N_2H_2AcO_2$ [285°–
290°] Tables, not attacked by hydroxylamine.
— NH_4A' aq

Di benzoyl derivative $C_6H_4N_2Bz_2O_2$
[216°] Brownish plates, insol water

DI OXY AMIDO PYRIMIDINE CARBOXY-

LIC ACID $NH \begin{matrix} \text{CO NH} \\ | \\ \text{CO C(NH}_2) \end{matrix} C CO_2H$ Formed
by reducing the nitro acid (Behrend, A 240, 21)
Not affected by boiling potassium cyanate—
 KA' aq

Ethyl ether EtA' [260°] (Kohler, A
236, 32)

OXY AMIDO-PYROTARTARIC ACID

$CH_2C(OH)(CO_2H)CH(NH_2)CO_2H$ or
 $CH_2C(NH_2)(CO_2H)CH(OH)(CO_2H)$ Amido-
citramalic acid S 31 at 18° Formed from
oxoacetic acid and NH_4Aq at 100° (Melikoff
& Feldmann, A 253, 92) Monoclinic prisms—
 $B'HCl$ prisms, v sol water

DI OXY AMIDO-PYRROLE Benzoyl de-
rivative $N \begin{matrix} \text{CH} \\ | \\ \text{C(OH)CH} \end{matrix} NHbz$ [2005°].

Got by heating at 140° the product of the action
of $NaOEt$ on hippuric ether (Rugheimer, B 22,
1957) Plates (from $MeOH$), v sol warm water

OXY-AMIDO-QUINIZINE-CARBOXYLIC

ACID ? $C_{19}H_{12}N_2O_2$ Anhydride of amido oxal-
acetic acid phenyl hydrazide Formed by warm-
ing the phenyl hydrazide of amido-oxalacetic
acid (vol 1 p 169) with conc H_2SO_4 White
glistering plates Insol hot water and ether,
v sl sol hot alcohol Dissolves in alkalis and
in conc H_2SO_4 Reduces Fehling's solution on
gently warming Mercuric salts are reduced in

Erythro - oxyanthraquinone $C_{14}H_8O_4$, *s.s.*
 $C_6H_4 \begin{array}{c} \text{CO C C(OH) CH} \\ \text{CO C CH=CH} \end{array}$ Mol. w 224 [191°]

Formation—1 By fusing *o*-bromo-anthraquinone with KOH at 150° (Pechmann, B. 12, 2127)—2 Together with its isomeride by heating phenol with phthalic anhydride and H_2SO_4 (Baeyer & Caro, B. 7, 968)—3 By the action of nitrous acid on *o*-amido anthraquinone (Roemer, B. 15, 1798)—4 By passing nitrous acid gas into an alcoholic solution of di-amido anthraquinone or (β) oxy amido anthraquinone (Von Perger, J pr [2] 18, 148)—5 Together with three isomeric di-oxy anthraquinones, by heating a mixture of benzoic acid (120 grms) and *m*-oxy-benzoic acid (60 grms) with 1200 grms of H_2SO_4 and 120 grms of water for 10 hours at 180°–200° (Liebemann & Kostanecki, B. 19, 829)—6 By oxidation of oxy-anthranol dihydride (Liebemann, A. 212, 20)

Properties—Orange feathery needles (from alcohol), insol water, sol ether and benzene. Yields alizarin on fusion with KOH. Insol cold baryta-water, but boiling baryta yields a violet salt. Insol NH_4Aq , sol $HOAc$. Its ethereal solution shaken with baryta water forms a violet pp, while the isomeric oxyanthraquinone gives a red solution. KOHAq dissolves it with difficulty, forming a brown solution. May be sublimed at 140° in a current of gas.

Acetyl derivative $C_{14}H_8AcO_4$ [176°–179°]

Oxy-anthraquinone $C_6H_4 \begin{array}{c} \text{CO C CH C(OH)} \\ \text{CO C CH CH} \end{array}$

[302°]

Formation—1 A by product in the preparation of alizarin by fusing anthraquinone sulphonic acid with KOH or NaOH (Liebemann, B. 4, 108, 5, 868, A. 160, 141, Simon, B. 14, 464). Occurs also in the product of the dry distillation of sodium anthraquinone sulphonate (A. G. & W. H. Perkin, C. J. 47, 680)—2 Obtained also from *m*-bromo anthraquinone by careful potash fusion (Graebe & Liebemann, A. 212, 25, Suppl. 7, 290)—3 By the action of nitrous acid on *m*-amido anthraquinone.

Properties—Yellow needles (by sublimation). Readily soluble in excess of baryta water. Its alkaline solution is reddish brown. Potash-fusion yields alizarin. HIAq reduces it to anthrol and anthracene hexahydride. Sol NH_4Aq , forming a reddish yellow liquid— $Ba(C_{14}H_8O_4)_2 \cdot Ba(C_4H_9O_2)_2 \cdot aq$.

Acetyl derivative [158°]. Small needles. Formed by acetylation and also by oxidation of the acetyl derivative of anthrol.

Ethyl ether $C_{14}H_8O_4(OEt)$ [135°]. Sol alcohol. Very difficult to saponify (Liebemann & Hagen, B. 15, 1798).

Di-oxy-anthraquinone

$C_6H_4 \begin{array}{c} \text{CO C C(OH) C(OH)} \\ \text{CO C CH=CH} \end{array}$ Alizarin Mol. w.

240 [290°]. S 034 at 100° (Plessy & Schutzenberger, C. R. 43, 167). Occurs in madder, the root of *Rubra tinctoria*, which contains its glucoside $C_{22}H_{20}O_{14}$ (ruberythric acid), and the glucosides of purpurin, purpurin carboxylic acid, purpuroxanthin, and purpuroxanthin carboxylic acid (Rohbquet & Colin, A. Ch. [2] 34, 225, Runge, J pr 5, 362, Schunck, A. 66, 174, 201, 81, 336, 87, 344, P. M. [4] 5, 410, 495, 12, 200, 270, J pr. 59, 465, Roehleider, A. 80, 321, 82,

205, Debus, A. 66, 351; Wolff & Strecker, A. 75, 1, Wartha, B. 3, 545, 673, Willigk, A. 82, 339, Rosenstiehl, A. Ch. [5] 18, 235, C. R. 88, 1194, Wurtz, C. R. 96, 465, Liebemann, B. 20, 2241, Bergami, B. 20, 2247).

Formation—1 By fusing di-bromo-anthraquinone with potash (Graebe & Liebemann, Z. [2] 11, 516, A. Suppl. 7, 300)—2 By fusing anthraquinone sulphonic acid with potash (Perkin, C. J. 23, 133, B. 9, 281). The yield may be improved by addition of $KClO_4$ —3 By heating pyrocatechin with phthalic anhydride and H_2SO_4 at 140° (Baeyer & Caro, B. 7, 972).

Purification—1 By repeated solution in NaOHAq and ppn by CO_2 , the pp being decomposed by HCl (Liebemann & Troschke, B. 8, 379)—2 Crude commercial alizarin, a mixture of alizarin, flavo purpurin, and isopurpurin, is fractionally sublimed when the alizarin comes over from 100° to 160°, and a mixture of flavo purpurin and isopurpurin from 160° to 170°, the crystals of these two bodies can be distinguished in the sublimate with a microscope and can be separated by means of benzene, in which isopurpurin is only slightly soluble, but the flavo purpurin very soluble (Schunck & Römer, B. 13, 41).

Properties—Red trimetric needles, ∇ sol alcohol and ether. Sometimes crystallises in golden scales (containing 2aq (Schunck). Its solutions in alkalis and alkaline carbonates are purple. $BaCl_2$ and $CaCl_2$ give purple pps in these solutions. Alumina decolourises its alcoholic solution forming a red lake. Salts of Mg, Fe, Cu, and Ag give purple pps in the ammoniacal solution.

Reactions—1 Yields phthalic acid on oxidation with HNO_3 —2 Distillation with zinc dust yields anthracene (G. & L.)—3 Zinc dust and ammonia reduce it to di-oxy anthranol—4 Aqueous ammonia at 200° gives oxy amido anthraquinone, di-amido anthraquinone, and other bodies (Von Perger, J pr [2] 18, 129).

Salts— CaA^+aq — BaA^+aq (dried at 100°)

— PhA^+ violet brown pp

Di-acetyl derivative [160°] (P.), [179°–183°] (Baeyer, B. 9, 1232). Pale yellow crystals (Perkin, C. J. 26, 21).

Methyl ether $C_{14}H_8O_4(OH)(OMe)$. From alizarin, KOH, and MeI (Schunck, C. N. 27, 171).

Di-ethyl ether $C_{14}H_8O_4(OEt)$. Got by heating alizarin with K_2SO_4 and KOH at 170° (Habermann, M. 5, 238). Needles.

Bromo alizarin *v* vol 1 p 599

Chloro alizarin *v* vol 11 p 102

Nitro alizarin *v* NITRO DI OXY ANTHRAQUINONE

Amido alizarin ∇ DI OXY AMIDO ANTHRAQUINONE

Alizarin blue $C_{17}H_8NO_4$, *s.s.*

$C_6H_4 \begin{array}{c} \text{CO C} \\ \text{CO C C(OH) C(OH) C N=CH} \end{array}$ [270°]

Formed by heating (β) nitro alizarin (1 pt) with glycerin (1½ pts) and H_2SO_4 (5 pts) at 107°–200° (Prudhomme, Bl. [2] 28, 62, Auerbach, C. J. 35, 799, Graebe, A. 201, 333, B. 12, 1416). Brown needles. Alkalis form a blue solution, but excess of alkali throws it down again. Dyes chromium acetate mordant blue. Yields $C_{17}H_8NO_4(OH)NH_2$ [255°]. Gives anthraquinol-

ine when distilled with zinc dust —B'HCl—
B'HOAc— $C_6H_5(NO_2)_2OH$ [245°]—

$Ba_2OC_6H_4NO_2$, $\frac{1}{2}$ eq greenish-blue pp

Di-acetyl derivative $C_{11}H_8O_4$, [225°]

Di-benzoyl derivative [244°]

Alizarin blue S $C_{17}H_{11}NO_5(SO_2Na)_2$ Prepared by dissolving alizarin-blue in a cold concentrated solution of $NaHSO_3$ (Brunck & Graebe, *B* 15, 1783) Reddish brown crystalline powder V sol water, sl sol 95 p c of alcohol At 60° it begins to decompose into its constituents

Isolizarin occurs in madder that has been heated with HClAq or dilute H_2SO_4 (Rochleder, *B* 3, 292) Its alkaline solutions are blood red, and it forms a red solution with baryta water It does not dye mordants

p **Di-oxy-anthraquinone**

$C_6H_4 \begin{matrix} \diagup CO C C(OH) CH \\ \diagdown CO C C(OH) CH \end{matrix}$ **Quinizarin** [195°]

Formed by heating hydroquinone or *p* chloro phenol with H_2SO_4 and phthalic anhydride at 140° (Grinna, *B* 6, 506) Red needles (from alcohol) Its alkaline solutions are blue Yields anthracene on distillation with zinc dust Its ethereal solution shows greenish yellow fluorescence Baryta gives a bluish violet lake Oxidised by MnO_2 and H_2SO_4 at 140° to purpurin (Bayer & Caro, *B* 8, 152)

Reactions —1 HI (S G 17) and a little red

P give a dihydride, $C_6H_4 \begin{matrix} \diagup C(OH) \\ \diagdown C(OH) \end{matrix} C_6H_4(OH)_2$,

which forms golden plates or needles (from alcohol) It forms with NaOH a yellow solution, turned blue by air $SnCl_2$ and HCl also reduce quinizarin to this dihydride (Liebermann, *A* 212, 11) —2 A longer treatment with HI forms hydroquinizarol, $C_{11}H_{12}O_2$ It forms orange needles (from alcohol) This body is probably $C_6H_4 \begin{matrix} \diagup CH(OH) \\ \diagdown CH_2 \end{matrix} C_6H_4(OH)_2$ —3 Conc HI (S G 19) and red P after an hour's boiling reduce quinizarin to oxy anthranol dihydride (*q* v)

Di-acetyl derivative [200°] Prisms

Ethyl ether $C_{11}H_8O_4(OH)(OEt)$ [151°]

Di-ethyl ether $C_{11}H_8O_4(OEt)_2$ [177°]

Yellow needles (Liebermann, *B* 21, 1168)

m **Di-oxy-anthraquinone**

$C_6H_4 \begin{matrix} \diagup CO C C(OH) CH \\ \diagdown CO C CH = C(OH) \end{matrix}$ **Purpurazanthin**

Xanthopurpurin [265°] Occurs, together with its carboxylic acid, in madder (Schützenberger, *Bl* 4, 12) May be got by reducing purpurin with HIAq or with $SnCl_2$ and NaOH Formed also by dissolving P in a hot alkaline solution of purpurin (Rosenstiehl, *C R* 79, 764), by the action of nitrous acid on purpuramide (di-oxy amido anthraquinone) (Liebermann & Fischer, *B* 8, 974), and, together with anthrachrysone, by heating a mixture of *s*-di-oxy benzoic acid (1 pt), benzoic acid (5 pts), and H_2SO_4 at 105°–110° (Noah, *B* 19, 332)

Properties —Yellow needles, sol alcohol, HOAc, and benzene, insol water May be sublimed Its alkaline solution is red Its barium salt is easily soluble Gives purpurin on fusion with KOH Nitric acid oxidises it to phthalic acid Reduced by HI and P to a dihydride, which dyes alumina mordants like quercitron (Rosenstiehl, *A Ch* [5] 18, 224, *C R* 79, 764) Yields anthracene on distillation with zinc-dust

Vol. III

NH_3Aq at 150° yields brown needles of $C_{11}H_8O_4(OH)(NH_2)$ (Liebermann, *A* 183, 217)

Acetyl derivative $C_{11}H_8O_4(OAc)_2$ [184°]

Di-methyl ether [180°] (Plath, *B* 9, 1204)

Di-ethyl ether $C_{11}H_8O_4(OEt)_2$ [170°].

(*B* 1, 4') **Di-oxy-anthraquinone**

$CH CH = C CO C C(OH) CH$ **Anthraflavin**
 $CH C(OH) C CO C CH = CH$

[280°] Formed in small quantity (2 g), together with anthraflavic acid (30 g) and *m*-'benzdioxyanthraquinone' (5 g), by the action of H_2SO_4 on *m*-oxy benzoic acid (100 g) (Schunck & Römer, *B* 11, 1176, 1616) Formed also by potash fusion from anthraquinone (*p*) disulphonic acid (Liebermann & Dehnat, *B* 12, 1287), and by the action of nitrous acid on the di amido anthraquinone Obtained by reduction of di nitro anthraquinone [above 300°] (Römer, *B* 16, 369)

Properties —Yellow tables, insol water, sl sol alcohol, HOAc, ether, and CS_2 , v sol benzene Forms oxyanthraflavin on fusion with KOH Almost insol baryta water, Na_2CO_3 , and NH_3Aq , sol KOHAq Its solution in H_2SO_4 is cherry red with scarlet fluorescence Its Ca and Ba salts are crimson and insoluble

Acetyl derivative $C_{11}H_8O_4(OAc)_2$ [244°]

Got by oxidising (*p*) di oxy anthracene with CrO_3 in HOAc Yellow needles

Di-oxy-anthraquinone $C_{11}H_8O_4$, **Anthraflavic acid** [above 330°] S (95 p c alcohol) 118 at 10° (Rosenstiehl, *Bl* [2] 29, 401, 434) A by product in the preparation of alizarin, being formed by fusing anthraquinone (*a*)-disulphonic acid with KOH (Schunck, *B* 4, 360, 8, 1628, 9, 379, 679, Perkin, *C J* 24, 1109, 26, 19, 29, 851) Formed also from *m*-oxy benzoic acid and H_2SO_4 (*v supra*) Radiating yellow needles (from alcohol), insol ether and benzene, sl sol HOAc Its solution in H_2SO_4 is yellow Its solution in alkalis is yellowish red Its Ba and Ca salts are sl sol water It does not dye mordanted cotton Potash fusion forms flavopurpurin Distillation with zinc dust forms anthracene Boiling with zinc-dust and NH_3Aq forms $C_{11}H_8O_2$, whence $C_{11}H_7(OAc)$, [165°] (Liebermann, *B* 21, 445)

Salts — Na_2 5aq sl sol water — BaA 7aq reddish brown needles

Di-acetyl derivative $C_{11}H_8O_4(OAc)_2$ [229°]

Di-benzoyl derivative $C_{11}H_8O_4(OBz)_2$ [275°]

Di-methyl ether $C_{11}H_8O_4(OMe)_2$ [248°]

Di-ethyl ether [232°] Yellow needles

Di-oxy anthraquinone $C_{11}H_8O_4$ [above 330°]

Isoanthraflavic acid Occurs in crude commercial alizarin, being formed by fusing anthraquinone (*p*) disulphonic acid with potash (Schunck & Römer, *B* 9, 379, 15, 1041, Perkin, *C J* 29, 851) Yellow needles (containing aq), almost insol benzene and ether, sl sol HOAc and alcohol It dissolves in cold baryta-water and in KOHAq with dark red colour Does not possess tinctorial properties Its solution in H_2SO_4 is red Yields anthrapurpurin on fusion with KOH — BaA 7aq red needles, v sol aq

Di-acetyl derivative [c 195°] Crystals.

Di-ethyl ether [194°] Yellow needles

X X

(B. 1, 1' or 3')-Di-oxy-anthraquinone $C_{14}H_8(OH)_2$, $C_{14}H_8O_2$, *Chrysazin* [192°]. Formed by eliminating amidogen from di oxy-tetra-amido-anthraquinone (hydrochrysazimide) (Liebemann & Giesel, *B* 8, 1643, 9, 329, *A* 183, 184) Formed also by fusing anthraquinone χ disulphonic acid with potash (Liebemann & Dehnst, *B* 12, 1287) Golden laminae or reddish-brown needles, m sol alcohol and HOAc. Insol cold NH_4Aq and Na_2CO_3Aq Its solution in KOHAq is yellowish red, and that in H_2SO_4 is red Potash fusion yields oxychrysazin, and *o*- and *m*-oxy-benzoic acid Yields anthracene when distilled with red hot zinc dust Baryta- and lime water give red pps Does not dye mordanted cotton

Di-acetyl derivative [232°] Prepared by oxidising di-acetyl-chrysazol $C_{14}H_8(OAc)_2$ with CrO_3 (Liebemann, *B* 12, 186)

Di-oxy-anthraquinone $C_{14}H_8O_2$, *Isochrysazin* [175°-180°] Formed, together with *m*-benzodioxanthraquinone, by the action of nitrous acid and alcohol on the dye $C_{22}H_{14}N_2O_6$, got from *o* di nitro anthraquinone and H_2SO_4 (Lütschütz, *B* 17, 897) Deep red needles (from alcohol), readily sublimed Its solutions in NH_4Aq and KOHAq are violet red, that in H_2SO_4 is reddish-yellow Its Ba salt is insoluble It does not dye mordants

Di-acetyl derivative [160°-165°] Needles

Di oxy-anthraquinone $C_{14}H_8O_2$, '*m*-Benzodioxanthraquinone' [293°] A product of the action of H_2SO_4 on *m*-oxy benzoic acid (Schunck & Römer, *B* 11, 969) Formed also as above Yellow needles, insol water and CS_2 , sol alcohol, HOAc, ether, and benzene Its alkaline solution is yellow Gives purpurn by potash-fusion Its solution in H_2SO_4 is brownish-yellow When freshly ppt it dissolves in hot baryta-water, and on cooling the Ba salt separates in red needles, which, after drying, are insoluble Has no tinctorial power

Di-acetyl derivative [199°] Needles

Di-oxy-anthraquinone $C_{14}H_8O_2$, *Franguluc acid* [264°] Obtained, together with glucose, by hydrolysing frangulin which occurs in the bark of *Rhamnus Frangula* (Faust, *A* 165, 229) Orange needles (containing 14aq), sl sol hot water, m sol alcohol Its solution in KOHAq is cherry red Its ammoniacal solution gives a red pp with $BaCl_2$ Distillation over zinc dust gives anthracene

Di-acetyl derivative [184°] Prisms

Di oxy anthraquinone

$C_{14}H_8 \begin{matrix} \diagup CO \ C \ CH \ C(OH) \\ \diagdown CO \ C \ CH \ C(OH) \end{matrix}$. *Hystazarin* [282°]

Formed by heating pyrocatechin (*5g*) with phthalic anhydride (*68g*) and H_2SO_4 (*75g*) at 145° for 5 hours (Liebemann & Scholler, *B* 21, 2508, 22, 683) Orange yellow needles (from acetone), almost insol benzene, v sol alcohol, ether, and HOAc Its solution in KOHAq is blue, that in NH_4Aq is violet, and that in H_2SO_4 blood-red The Ba salt is blue, the Ca salt violet and insol water It dyes mordants slightly Yields anthracene on distillation with zinc-dust

Di-acetyl derivative [207°]. Needles

Ethyl ether $C_{14}H_8O_2(OH)(OEt)$. [284°-

240°] Yellow needles, forming a crimson solution in alkalis

Di-ethyl ether $C_{14}H_{10}O_2(OEt)_2$, [160°-163°]. From hystazarin, KOH, and EtI Yellow needles

Eleven di oxy anthraquinones have been described, but ten only are indicated by theory

Tri-oxy-anthraquinone

$C_{14}H_5 \begin{matrix} \diagup CO \ C \ C(OH) \ C(OH) \\ \diagdown CO \ C \ C(OH) \ CH \end{matrix}$ *Purpurn* [253°]

Occurs in madder root, probably as glucoside (Colin & Robiquet, *A Ch* [2] 84, 244, Gauthier de Claubry & Persoz, *A Ch* [2] 48, 69, 51, 110, Runge, *A Ch* [2] 63, 282, Schiel, *A* 60, 74, Debus, *A* 66, 351, 86, 117, Wolff & Strecker, *A* 75, 1, Rochleder, *A* 80, 321, 82, 205, Stenhouse, *Pr* 12, 633, 13, 145, Schützenberger, *J* 1864, 542, *Bl* [2] 4, 12) Formed by the action of MnO_2 and H_2SO_4 at 150° on alizarin (De La lande, *C R* 79, 669) and on quinzarin (Baeyer & Caro, *B* 8, 152), and also by heating tri bromo-anthraquinone with H_2SO_4 at 200° (Diehl, *B* 11, 184) It may be separated from alizarin by repeatedly dissolving in boiling alum solution and ppg by acid Orange prisms containing aq (from dilute alcohol) or dark-red anhydrous needles (from absolute alcohol) May be sublimed Sl sol boiling water, forming a yellow liquid Its ethereal solution is yellow and shows two absorption bands (Stokes, *C J* 12, 220, Vogel, *B* 9, 1641) Its solution in H_2SO_4 is rose red and shows three absorption bands Aqueous KOH, Na_2CO_3 , and NH_3 yield purple red solutions Almost insol alcoholic potash Baryta-water forms an insoluble purple lake Its alkaline solution is oxidised in daylight by the air becoming yellow (unlike alizarin) the product containing phthalic acid (Schunck & Römer, *C J* 31, 665, Dralle, *B* 17, 376) Boiling alum forms a pink solution with yellow fluorescence Leadacetate gives in an alcoholic solution a dark-crimson pp, soluble in excess, forming a crimson liquid with three absorption bands (the lead compound of alizarin is insol alcoholic lead acetate) Dyes cotton, mordanted with alumina, red

Reactions —1 Yields anthracene on heating with zinc dust —2 Forms some quinzarin when heated in sealed tubes at 300° —3 Phosphorus and NaOHAq reduce it to purpuroxanthin —4 Nitric acid oxidises it to phthalic acid —5 Aqueous ammonia forms brown purpuramide $C_{14}H_5O_4(NH_2)(OH)_2$

Acetyl derivative $C_{14}H_5O_2(OAc)_2$, [198°] (*L*), [200°] (*S* & *R*) Yellow needles

Ethyl ether $C_{14}H_5O_2(OEt)_2$, Red crystals

Tri oxy anthraquinone $C_{14}H_5O_3(OH)_2$, *Anthrapurpurn* *Isoapurpurn* [above 330°] A by product in the preparation of artificial alizarin, being formed by fusing anthraquinone (*8*) disulphonic acid with potash (Perkin, *C J* 25, 659, 26, 425, 29, 851, Caro, *B* 9, 682) Formed also by potash-fusion from isanthraflavic acid, from *m* benzodioxanthraquinone (Schunck & Römer, *B* 9, 679, 11, 972), and from (a) di bromo anthraquinone (Perkin, *C J* 37, 557) Orange needles (from alcohol), v sol hot alcohol, sl sol hot water and ether Conc H_2SO_4 forms a red solution, potash, NH_4Aq , and Na_2CO_3Aq give a violet solution. Sl sol hot baryta-water, forming

a violet solution Alcoholic lead acetate gives a purple pp, sol excess It colours mordants

Reactions—1 *Nitric acid* gives no phthalic acid—2 Aqueous ammonia at 100° forms an unstable blue dye, decomposed by HCl or KOH with regeneration of anthrapurpurin—3 Aqueous ammonia at 170° forms anthrapurpuramide $C_{12}H_8O_4(OH)_2(NH_2)$ which does not dye mordants (Perkin, *C J* 83, 216)

Tri-acetyl derivative [222°] Yellow scales

Tri-benzoyl derivative [185°] Crystals

Mono ethyl ether $C_{12}H_8O_4(OH)(OEt)$ [265°] Orange red needles (Liebermann & Jellinek, *B* 21, 1170)

Di-ethyl ethers $C_{12}H_8O_4(OH)(OEt)_2$ [162°] and [170°] Yellow needles (L a J)

Tri-oxy-anthraquinone $C_{12}H_6O_6$ *Flavopurpurin* [above 330°] Formed by potash fusion from anthraflavic acid and from anthraquinone (a) disulphonic acid Purified by means of its lead salt (S a R, C, Liebermann, *B* 21, 441, 2524) Golden needles (from alcohol), v sol. cold alcohol Its solution in conc H_2SO_4 is red, that in KOHAq is purple, becoming red on dilution Sl sol baryta water, forming a red solution Its solution in NH_4Aq and Na_2CO_3Aq is yellowish red Alcoholic lead acetate forms a reddish brown pp, v sl sol excess On heating with phenyl cyanate at about 160° it forms $C_{12}H_8O_4(O CO NHPh)_2$ crystallising in yellowish plates (Tesmer, *B* 18, 2610)

Di-acetyl derivative [238°] Golden plates

Tri acetyl derivative [196°] Yellow needles

Di-benzoyl derivative [210°] Needles

Ethyl ether $C_{12}H_8O_4(OH)(OEt)$ V sol ether

Di-ethyl ether [209°] Needles

Tri oxy anthraquinone

$C_{12}H_6O_6 \begin{cases} CO C(OH) C(OH) \\ CO C CH= C(OH) \end{cases}$ *Anthragallol* [310°] (Cahn, *B* 19, 2335) Formed by heating a mixture of benzoic acid, gallic acid, and H_2SO_4 at 70° (Seuberlich, *B* 10, 38) Formed also from pyrogallol, phthalic anhydride, and H_2SO_4 . Orange needles (by sublimation), nearly insol water Its alkaline solution is green Dilute HNO_3 forms phthalic acid Distillation over zinc dust gives anthracene Dyes alumina mordants brown Alcoholic lead acetate ppts violet brown $C_{12}H_6O_4Pb_2O_4$ Boiling alcoholic NH_3 forms anthragallolamide $C_{12}H_6O_4(NH_2)(OH)_2$ crystallising in black needles with green reflex HCl and HOAc form $C_{12}H_6O_4$, whence $C_{12}H_6O_4$, [205°] (L)

Tri-acetyl derivative [173°] Needles

Ethyl ether $C_{12}H_8O_4(OH)(OEt)$ [175°]

From the K salt and EtI at 80° The Pb salt and EtI at 220° yields an isomeride [245°]

Di-ethyl ether [134°] Formed from the K salt (L a J) The Pb salt yields an isomeride [198°]

Tri-oxy anthraquinone $C_{12}H_6O_6$ *Oxychrysaem* Formed by potash-fusion from chrysaem and from anthraquinone ρ and χ disulphonic acids (Liebermann, *A* 183, 191, 12, 1889). Probably identical with oxyanthrarufin Red needles (from alcohol). Its alkaline solutions are blue.

Baryta-water gives a blue insoluble pp Dyes mordants

Tri-acetyl derivative [193°] Yellow needles

Tri-oxy-anthraquinone

$C_{12}H_6(OH)C_6O_4C_6H_2(OH)_2$ *Oxyanthrarufin*. Prepared by fusing anthrarufin with KOH (Liebermann & Boeck, *B* 11, 1617) Red needles (by sublimation) Its alkaline solutions are blue. Dyes mordants like alizarin

Tetra-oxy anthraquinone $C_{12}H_4O_8$ *t.e.*

$C(OH) CH C(O) C(O) CH= C(OH) CH C(OH) C(O) C(O) C(OH) CH$ *Anthrachry-one* Mol w 272 Formed by heating *s* di-oxybenzoic acid alone or with H_2SO_4 (Barth & Senhofer, *A* 164, 109, Noah, *B* 19, 754) Silky needles (containing 2aq), not melted at 360° V. sol alcohol, v sl sol water and ether Yields anthracene on distilling with zinc-dust Its solution in KOHAq is reddish-yellow—Ba($C_{12}H_4O_8$), 11aq red needles

Tetra-acetyl derivative [253°] Needles.

Tetra-oxy-anthraquinone $C_{12}H_4O_8$ *Oxyppurpurin* Formed by heating purpurin with KOH at 240° (Diehl, *B* 11, 185) Brown nodules, not melted at 290° Almost insol alcohol Its alkaline solution is brownish red. Its acetyl derivative melts above 240°

Tetra-oxy-anthraquinone

$CH C(OH) C(O) C(O) C(OH) C(OH) CH C(OH) C(O) C(O) CH= CH$ *Quinalizarin*. [above 275°] Obtained by heating its di-methyl ether with HOAc and HCl at 200° Formed also by heating alizarin with H_2SO_4 at 210° (Graebe, *B* 23, 3739) Long red needles, sol HOAc, sl sol alcohol Dyes mordants well Its solution in H_2SO_4 is blue The Ba and Ca salts are insol water Yields anthracene on distillation with zinc dust

Tetra-acetyl derivative [201°] Needles.

Di-methyl ethyl [225°–230°] Obtained by heating hempic acid $C_6H_3(OMe)_2(CO_2H)_2$ [6 5 2 1] with hydroquinone and H_2SO_4 at 130° (Liebermann & Wense, *B* 20, 864, *A* 240, 298) Minute brownish red plates Forms a bluish-violet solution in alkalis and a blue solution in H_2SO_4 . Does not dye. Yields $C_{12}H_4O_4(OMe)_2(OAc)_2$ [211°]

Tetra-oxy-anthraquinone $C_{12}H_4O_8$ *Rufin*

opin Formed by heating opianic acid with H_2SO_4 at 180° (Liebermann & Chojnacki, *B* 4, 194, *A* 162, 322) Yellowish red needles, sl sol boiling water, m sol hot alcohol Its solution in KOHAq is violet red, that in NH_4Aq is reddish-brown, and is ppd by $BaCl_2$ and $CaCl_2$. Yields anthracene on distilling with zinc dust. Conc H_2SO_4 gives a violet red solution Dyes alumina mordants brownish red—Ba($C_{12}H_4O_8$), aq.

Tetra-oxy-anthraquinone $C_{12}H_4O_8$ (a) *Oxyanthragallol* [above 350°] Formed, together

with the (β)-isomeride and rufigallol acid, by heating pyrogallol with *m*-oxy-benzoic acid and H_2SO_4 at 150° (Noah, Liebermann & Kostanecki, *A* 240, 270) Golden needles (from alcohol) or red needles (from benzene) Conc. H_2SO_4 forms a violet solution KOHAq gives a green solution Insol baryta-water Dyes mordants

Tetra-acetyl derivative [209°].

Tetra-oxy-anthraquinone $C_{12}H_4O_8$ (β) *Oxyanthragallol*. [above 380°] Formed as above.

Red needles (from alcohol), insol benzene. Conc H_2SO_4 gives a brownish yellow solution KOHAq forms a green solution Dyes mordants

Tetra-acetyl derivative [189°]

Of the two oxyanthraquinones one should have the hydroxyls in the position 1,2,3,2', and the other in 1,2,3,4'

Penta-oxy-anthraquinone

$\text{C}(\text{OH})\text{CH}(\text{C})\text{CO}\text{C}(\text{OH}) = \text{C}(\text{OH})\text{CH}(\text{C}(\text{OH}))\text{CO}\text{C}(\text{OH})\text{C}(\text{OH})$ Formed by heating gallic acid with *s*-di oxy benzoic acid H_2SO_4 for 15 minutes at 165° (Liebemann & Noah, *B* 19, 751, *A* 240, 273) Small yellowish-red plates (by sublimation), not melted at 360° V sol hot alcohol, sl sol ether, nearly insol benzene and hot water Its solution in KOHAq is green, that in H_2SO_4 is brownish red Dyes mordant fabrics

Penta-acetyl derivative [229°]

Needles

Hexa oxy-anthraquinone

$\text{C}(\text{OH})\text{C}(\text{OH})\text{C}(\text{CO})\text{C}(\text{CH}) = \text{C}(\text{OH})\text{C}(\text{OH}) = \text{CH}(\text{C})\text{CO}\text{C}(\text{OH})\text{C}(\text{OH})$ *Rufgallisch acid* Formed by heating gallic acid with H_2SO_4 (Robiquet, *A* 19, 204, Wagner, *C* 1861, 47, Löwe, *J* pr 107, 296, Jaffé, *B* 3, 694, Widman, *B* 9, 856, Klobulowski & Nöltung, *B* 8, 819, 9, 1256, 10, 880) Red crystals (containing 2aq), v sl sol alcohol and ether Its solution in conc KOHAq is blue, becoming violet red on dilution Conc H_2SO_4 forms a red solution Baryta-water forms a blue insoluble salt Dyes fabrics, mordanted with alumina, brown, with iron salts, black

Reactions —1 Yields anthracene on heating with *conc*-*acid* —2 Nitric acid gives no phthalic acid —3 Boiling HIAq and P form minute needles of $\text{C}_{14}\text{H}_{10}\text{O}_4$ —4 *Potash-fusion* yields *m*-oxy benzoic acid, γ -oxy-isophthalic acid, and an anhydride of hexa-oxy-diphenyl $\text{C}_{20}\text{H}_{10}\text{O}_{11}$ 4sq, crystallising in colourless needles (Mahn, *A* 141, 346, Schreder, *M* 1, 432)

Hexa-acetyl derivative Needles

Chloro-acetyl derivative $\text{C}_{14}\text{H}_9\text{ClO}_6$

Needles

Tri-ethyl ether [195°] Orange needles

Tetra-methyl ether [c 220°] Needles

Tetra-ethyl ether [above 180°] Red needles

Hexa-ethyl ether $\text{C}_{14}\text{H}_{20}(\text{OEt})_6$ [c 140°] Orange needles, sol hot alcohol

References —Bromo-, Chloro-, and Nitro-OXYANTHRAQUINONE and OXY AMIDO ANTHRAQUINONE

OXY-ANTHRAQUINONE CARBOXYLIC

ACID $\text{C}_6\text{H}_4\text{C}(\text{CO})\text{C}_6\text{H}_4(\text{OH})\text{CO}_2\text{H}$ [260°]. Prepared by fusing anthraquinone carboxylic acid with caustic soda (Hammerschlag, *B* 11, 83) Orange needles (by sublimation) Yields phthalic acid on oxidation with HNO_3 Its Ba salt is a blue pp Its alkaline solutions are purple

(*erythro*)-Oxy-anthraquinone-carboxylic acid [238°] Formed by heating (*erythro*)-oxy-methyl-anthraquinone with H_2SO_4 , a portion of the quinone probably oxidising the methyl of the rest (Birukoff, *B* 20, 2488) Long yellow needles V sol boiling water Heated to about 270° it loses CO_2 and gives (*erythro*)-oxy-anthra-

quinone. The Ca and Ba salts are sparingly soluble

Di oxy-anthraquinone carboxylic acid

$\text{C}_6\text{H}_4(\text{OH})\text{C}_6\text{H}_3(\text{CO}_2\text{H})\text{CO}_2\text{H}$ *Alizarin carboxylic acid* [305°] Formed by soda-fusion from the sulphonic acid got by heating anthraquinone carboxylic acid with H_2SO_4 (Hammerschlag, *B* 11, 86) Dull-red powder or red needles (by sublimation), v sol aqueous NaOAc Decomposed by heat into CO_2 and alizarin Its alkaline solutions are purple Gives a red lake with alumina Nitric acid oxidises it to trimellitic acid — $\text{Ba}_2(\text{C}_{12}\text{H}_6\text{O}_6)_2$ blue pp

Di oxy-anthraquinone carboxylic acid

$\text{C}_{12}\text{H}_6\text{O}_6$ *Munjistin Purpuranthric acid* [231°] Occurs in madder (Schunck & Römer, *B* 10, 172, *C* J 81, 666, 83, 422) and in munjeet or East Indian madder (Stenhouse, *Pr* 12, 633, 13, 86, 145) Golden scales (from HOAc), split up at 233° into CO_2 and purpuranthrin M sol boiling water and alcohol Its alkaline solutions are red Conc H_2SO_4 forms an orange solution Dilute nitric acid oxidises it to phthalic acid Fuming HNO_3 gives a di nitro-derivative Dyes fabrics, mordanted with alumina, orange, with iron mordants, brownish red Forms a red insoluble Ba salt Yields purpurin when boiled for a long time with conc KOHAq Ammonia at 100° forms purpurin amide Br in HOAc forms di bromo-purpuranthrin [231°]

Tri-oxy-anthraquinone carboxylic acid

$\text{C}_{12}\text{H}_6\text{O}_7$ *Purpurin carboxylic acid Pseudopurpurin* [220°] Occurs in madder (Schutzenberger & Schiffrt, *Bl* 4, 13, Rosenstiehl, *C* R 84, 561, Liebemann, *B* 10, 1618) Red plates, almost insol water and alcohol Split up by heat, or by boiling with water, potash, or alcohol into CO_2 and purpurin Its alkaline solution is orange-red Does not dye mordanted fabrics unless the water is free from CaCO_3 (difference from purpurin) Bromine water yields bromo-purpurin [275]

OXY-ANTHRAQUINONE SULPHONIC

ACID $\text{C}_6\text{H}_4\text{C}_6\text{H}_4(\text{OH})(\text{SO}_3\text{H})$ [1 6 2 3 or 4] *Erythro oxy anthraquinone sulphonic acid* Formed from *o* amido anthraquinone sulphonic acid by the action of nitrous acid (Litschütz, *B* 17, 900) Yellow plates, v sol water, alcohol, and ether Its alkaline solution is red On fusion with potash it gives alizarin — AgA' yellow needles

Anhydride $\text{C}_{12}\text{H}_6\text{O}_2\text{SO}_2$ Grey needles, insol water, alcohol, and ether, sl sol HOAc Requires to be heated with alkalis before yielding the acid

Oxy anthraquinone sulphonic acid Formed by sulphonating erythro oxy anthraquinone at 130° Sol water and alcohol, insol ether Its alkaline solution is reddish yellow Baryta gives a blood red pp, crystallising from hot water BaCl_2 gives a similar yellow pp Lead acetate gives a yellowish-brown pp, sol hot water Gives neither alizarin nor purpurin on potash fusion

m-Oxy-anthraquinone sulphonic acid

$\text{C}_{12}\text{H}_6\text{O}_7(\text{SO}_3\text{H})$ Formed by heating *m*-oxy-anthraquinone with H_2SO_4 at 120° (Von Perger, *J* pr [2] 18, 176) Crystals (from alcohol), m. sol cold water, insol ether Its alkaline solu-

tion is reddish brown Lead acetate gives a brownish yellow, and lime-water a brown, pp Yields alizarin sulphonic acid on fusion with potash at 180° — $\text{BaCl}_2 \cdot \text{H}_2\text{SO}_4$ orange crystals

When (a)- or (β)-anthraquinone disulphonic acid is fused with moist potash, a mixture of an anthraquinone disulphonic acid, oxyanthraquinone sulphonic acid, and tri oxy-anthraquinone is got The mono sulphonic acid is obtained by stopping the fusion as soon as the blue colour begins to turn violet (Graebe & Liebermann, *A* 160, 139) According to Von Perger (*J pr* [2] 18, 168) it is doubtful whether the product is not a di oxy-anthraquinone sulphonic acid The acid prepared from (α)-anthraquinone disulphonic acid gives on fusion with potash anthraflavic acid and flavo-purpurin, and is therefore isomeric with that from (β) anthraquinone disulphonic acid, which gives isanthraflavic acid and anthra purpurin on potash fusion

Di-oxy anthraquinone sulphonic acids

$\text{C}_{14}\text{H}_2\text{O}_2(\text{OH})_2(\text{SO}_3\text{H})$ Pure alizarin heated with strong H_2SO_4 at 120° forms at least three sulphonic acids On adding water two sulphonic acids dissolve and another remains on the filter as a brown mass The latter dissolves in alkalis, forming a cherry red liquid It is slightly soluble in water, but is slowly decomposed by boiling water, regenerating alizarin Of the two sulphonic acids which are readily soluble, the one that is formed in greatest quantity is the most soluble, and it is decomposed by potash fusion at 200° without forming either alizarin or purpurin The other sulphonic acid is formed in very small quantity, but by potash fusion it is converted at 140° into purpurin, the mass becoming crimson (Von Perger, *J pr* [2] 18, 173)

Di-oxy-anthraquinone sulphonic acid

$\text{C}_{14}\text{H}_2\text{O}_2(\text{OH})_2\text{SO}_3\text{H}$ *Quinizarin sulphonic acid* Formed in small quantity in the preparation of quinizarin from hydroquinone, phthalic anhydride, and H_2SO_4 (Liebermann, *A* 212, 11) Its Na salt forms a deep orange solution, turned deep blue by alkalis It does not dye mordants

OXY-AZELAIC ACID $\text{C}_8\text{H}_8(\text{OH})(\text{CO}_2\text{H})_2$

[91°] Formed by the action of NaOHAq upon the product of the action of Br and a little red P upon azelaic acid (Bujard & Hell, *B* 22, 68) Nodules (from water or ether)— $\text{BaA}'' \frac{1}{2}\text{aq}$ S 256 at 18° — $\text{CaA}'' \frac{1}{2}\text{aq}$ S 65 at 20° — $\text{ZnA}'' \frac{1}{2}\text{aq}$ S 623 at 20° — $\text{SrA}'' \frac{1}{2}\text{aq}$ — $\text{MgA}'' \frac{1}{2}\text{aq}$ — $\text{CdA}'' \frac{1}{2}\text{aq}$ — $\text{CuA}'' \frac{1}{2}\text{aq}$ — $\text{PbA}'' \frac{1}{2}\text{aq}$ — AgA''

OXY-AZO- compounds v Azo-compounds

OXY-AZOPHENINE $\text{C}_{10}\text{H}_7\text{N}_3\text{O}$ [197°]

Formed by heating nitroso *m*-oxy-diphenylamine with aniline and aniline hydrochloride on the water bath (O Fischer & Hepp, *B* 20, 2481, Kohler, *B* 21, 910) Needles (from toluene), sol alcoholic NaOH, insol NaOHAq Conc. H_2SO_4 forms a reddish-brown solution.

DI-OXY-BEHENIC ACID $\text{C}_{22}\text{H}_{40}(\text{OH})_2\text{O}_2$

[127°] (H), [138°] (H & G) Formed by boiling oxy-erucic acid with potash (Haussknecht, *A* 143, 53), or by oxidising erucic acid with alkaline KMnO_4 (Irwantsoff, *J R* 21, 13, *J pr* 39, 884; Hazura & Grüssner, *M* 9, 947) Plates (from alcohol), insol water and ether With Pb , it gives iodobehenic acid reduced by zinc and

HCl to behenic acid NaA' grains, sl. sol water— BaA'_2 insoluble pp

ISO-DI OXY-BEHENIC ACID $\text{C}_{22}\text{H}_{40}(\text{OH})_2\text{O}_2$ [99°] Formed by oxidising brassic acid with alkaline KMnO_4 (Grüssner & Hazura, *M* 10, 197) Minute plates, insol water and ligroin, v sol hot alcohol

DI-OXY-BEHENIC ACID $\text{C}_{22}\text{H}_{40}\text{O}_2$ [91°]

Formed by oxidising behenic (benolic) acid with fuming HNO_3 (Haussknecht, *A* 143, 48) Yellowish scales (from alcohol)— AgA' white pp

OXYBENZALDEHYDE v OXYBENZOIC ALDEHYDE

HYDE

OXYBENZAMIDE v Amide of OXYBENZOIC ACID

O-OXY-BENZAMIDINE Ethyl ether $\text{C}_8\text{H}_7(\text{OEt})\text{C}(\text{NH})\text{NH}_2$ A product of the action of alcoholic HCl, followed by alcoholic NH_3 , on $\text{C}_8\text{H}_7(\text{OEt})\text{CN}$ (Pinner, *B* 23, 2952) The hydrochloride $\text{B} \cdot \text{HCl}$ [218°] crystallises in short hexagonal columns, v sol water

p Oxy benzamidine Ethyl ether The hydrochloride $\text{C}_8\text{H}_7(\text{OEt})\text{C}(\text{NH})\text{NH}_2\text{Cl}$ [260°] is formed by the action of alcoholic NH_3 on the hydrochloride of ethyl p oxy benzimidio-ether It is converted by acetoacetic ether and NaOHAq into ethylated dioxy phenyl methyl pyrimidine $\text{C}_8\text{H}_7(\text{OEt})\text{C} \begin{smallmatrix} \text{N}^{\text{CM}} \\ \text{C}(\text{OH}) \end{smallmatrix} \text{CH}$

OXY-BENZENE v PHENOL

Di-oxy-benzene v HYDROQUINONE, PYROCAT-CHIN, and RESORCIN

u-Tri-oxy-benzene $\text{C}_6\text{H}_3(\text{OH})_3$ [124°] *Oxy hydroquinone* [140.5°] Formed, together with hexa oxy diphenyl, by fusing hydroquinone (1 pt) with moist NaOH (9 pts) (Barth & Schreder, *M* 4, 176, 5, 589) Monoclinic plates (from ether), $a b c = 75.1101$, $\beta = 91^\circ 46'$ v e sol water, alcohol, ether, and HOAc, almost insol chloroform and benzene Its alkaline solution rapidly becomes brown through absorption of oxygen FeCl_3 gives a transient bluish green colour H_2SO_4 forms a green solution becoming cherry-red on warming Bromine forms $\text{C}_6\text{Br}_3(\text{OH})_3$ [206°] Nitric acid yields greyish blue crystals of oxyquinhydrone

Tri-acetyl derivative $\text{C}_6\text{H}_3(\text{OAc})_3$ [96°]

Methyl ether $\text{C}_6\text{H}_3(\text{OMe})(\text{OH})_2$ [24.1°] [84°] Got by reducing the methyl ether of oxyquinone with aqueous SO_2 (Will, *B* 21, 606) Colourless plates Turned yellow by FeCl_3 , being reconverted into $\text{C}_6\text{H}_3(\text{OMe})_3$

Tri-methyl ether $\text{C}_6\text{H}_3(\text{OMe})_3$ (247°) Formed from the preceding ether, methyl iodide, and KOH (W) Obtained also from asarone $\text{C}_8\text{H}_7(\text{OMe})_2\text{CH}_2\text{CHMe}$ by oxidation to $\text{C}_6\text{H}_3(\text{OMe})_3\text{CO}_2\text{H}$ and distillation of this acid with lime (Rizsa & Butlerow, *J R* 1887, 1). Oil, volatile with steam

Mono-ethylether $\text{C}_6\text{H}_3(\text{OH})(\text{OEt})$ [148°] [112°] Formed by reduction of the ethyl ether of oxyquinone $\text{C}_6\text{H}_3(\text{OEt})_2\text{O}_2$ with SO_2 Colourless prisms Sublimes in fine plates v sol water, alcohol, and ether, m sol benzene Fe_2Cl_6 gives a dark brown colouration.

Tri-ethyl ether $\text{C}_6\text{H}_3(\text{OEt})_3$ [84°] Formed by ethylation of the preceding body. Long white glistening needles, v e sol. alcohol and ether, insol water, volatile with steam. This body is identical with that obtained from esculetin, which is therefore a derivative of

***m*-tri-oxy-benzene** (Will & Fukall, *B*, 20, 1133, Herzog & Zeisel, *M*, 10, 150)

Isomerides are described as PHLOROGLUCIN and PYROGALLOL.

***s*-Tetra-oxy-benzene** $C_6H_2(OH)_4$ [1 2 4 5] [α 218°]. Formed by reducing di-oxy-quinone with $SnCl_2$ (Nietzki, *B*, 21, 2377). Colourless plates, v sol water, alcohol, and ether. Its aqueous solution rapidly turns brown, and its alkaline solution is oxidised by air or by $FeCl_3$ to di-oxy-quinone.

Acetyl derivative $C_6H_2(OAc)_4$ [217°]. Colourless plates.

***Di*-methyl ether** $C_6H_2(OH)_2(OMe)_2$ [1 4 2 5] [166°]. Formed by reducing $C_6H_2O_2(OMe)_2$ with stannous chloride (Nietzki & Reichberg, *B*, 23, 1217). Colourless plates.

***Di*-ethyl ether** $C_6H_2(OH)_2(OEt)_2$ [1 4 2 5] [188°]. Got in like manner. Colourless needles (from hot water). Yields $C_6H_2O_2(OEt)_2$ on oxidation. Acetic anhydride forms the compound $C_6H_2(OAc)_2(OEt)_2$ [148°].

***Tetra*-ethyl ether** $C_6H_2(OEt)_4$ [143°]. Formed from $C_6H_2(OH)_2(OEt)_2$, alcoholic $NaOEt$ and $EtBr$ at 100° (Nietzki, *B*, 23, 1214). Colourless plates, smelling like anise. May be sublimed.

***u*-Tetra-oxy-benzene** ***Di*-methyl ether** $C_6H_2(OH)_2(OMe)_2$ [1 4 3 5] [158°]. Formed by reducing the corresponding $C_6H_2O_2(OMe)_2$ with Zn and $HClAq$, with $SnCl_2$, or with SO_2 (Hofmann, *B*, 8, 67, 11, 332, Will, *B*, 21, 609). Needles, reconverted into the quinone by $FeCl_3$. $HOAc$ forms an acetyl derivative [133°] which forms a green solution in H_2SO_4 . Ac_2O forms an acetyl derivative [128°] which does not give a green solution in H_2SO_4 . (H) Benzoyl chloride yields $C_6H_2(OMe)_2(OBz)_2$ [245°], v sl. sol alcohol.

***Tre*-methyl ether** $C_6H_2(OH)(OMe)_3$ [146°]. Formed, together with the *tetra*-methyl ether by the action of methyl iodide and KOH on the *di*-methyl ether. Needles (from alcohol).

***Tetra*-methyl ether** $C_6H_2(OMe)_4$ [47°]. [271°]. Plates (from ether). Yields $C_6Br_4(OMe)_4$ [76°].

***o*-Tetra-oxy-benzene** ***Di*-methyl ether** $C_6H_2(OH)_2(OMe)_2$. ***Di*-methyl apionol** [106°] [296°]. Formed by heating apionol acid $C_6H_2O_2(OMe)_2CO_2H$ with alcoholic potash, at 180° (Ciamician & Silber, *B*, 22, 119, 2482, 23, 2391). Insol water, sol alcohol, ether, and benzene. Its solution in KOH becomes brownish-red. Ferric chloride gives a violet-black colour. Lead acetate gives a gelatinous pp. $AgNO_3$ forms minute needles, quickly turning black. Yields a crystalline acetyl derivative [144°].

***Tetra*-methyl ether** $C_6H_2(OMe)_4$ [1 2 3 4] [31°]. White needles, v sol ether.

Hexa-oxy-benzene $C_6(OH)_6$ (so-called '*trihydrocarboxylic acid*' of Lerch). Long nearly colourless needles. Soluble in hot water, slightly soluble in cold water, alcohol, ether, and benzene.

Formation—1 By the action of dilute HCl upon freshly prepared carbonic-oxide potassium. 2 By reduction of tri-quinone C_6O_3 with stannous chloride.

Properties—Reduces $AgNO_3$ in the cold. Is oxidised by HNO_3 to benzené-tri-quinone. Gives a violet colouration with $FeCl_3$. The solution in aqueous Na_2CO_3 is readily oxidised by the air to tetra-oxy-quinone $C_6(OH)_4O_2$. By distillation with zinc dust it gives benzene and diphenyl. By evaporation in an open dish with dilute KOH it yields orcinic acid $C_6H_2O_4$.

Salt $C_6(OK)_6$. Formed by combination of CO with K at 80°, occurring as a by-product in the preparation of K (Laebig, *A*, 11, 182, Brodie, *A*, 113, 358, Lerch, *A*, 124, 20). Grey mass, becoming explosive on exposure to air.

Hexa-acetyl derivative $C_6(OAc)_6$ [203°], small colourless prisms, sl sol hot acetic acid, nearly insol alcohol, ether, and benzene (Nietzki & Benckiser, *B*, 18, 505, 1833).

OXY-BENZENE CARBOXYLIC ACID v Oxy BENZOIC ACID.

Oxy-benzene dicarboxylic acid v Oxy PHTHALIC, OXY-ISOPHTHALIC, and OXY TEREPHTHALIC ACIDS.

Tri-oxy-benzene carboxylic acid v GALLIC ACID.

Tri-oxy-benzene tri-carboxylic acid v PHLOROGLUCIN TRICARBOXYLIC ACID.

Di-oxy benzene tetra-carboxylic acid v HYDROQUINONE TETRA-CARBOXYLIC ACID.

DI OXY-BENZENE-DIQUINONE v Di OXY DIQUINONE.

OXY-BENZENE SULPHONIC ACID v PHENOL SULPHONIC ACID.

Di-oxy-benzene sulphonic acid $C_6H_2(OH)_2SO_3H$ [280°]. Formed by heating phenol β' -disulphonic acid with potash at 240° (Senhofer, *J*, 1879, 749). Crystallises from water in needles (containing aq). Coloured violet by $FeCl_3$. — BaA' , 7aq — ZnA' , 27aq — PbA' , 8aq long thin plates.

Isomerides HYDROQUINONE, PYROCATECHIN, and RESORCIN SULPHONIC ACIDS.

OXY-BENZENYL-AMIDO-PHENYL MER CAPTAN $C_{12}H_9NSO$ \approx $C_6H_4 \begin{smallmatrix} N \\ \diagup \diagdown \end{smallmatrix} C_6H_4OH$.

[129°]. Formed by heating salicylic aldehyde with *o*-amido phenyl mercaptan (Hofmann, *B*, 13, 1237). Needles — $B'HCl$ — Platinochloride B'_2H_2PtCl , pp.

***o*-OXY-BENZENYL-AMIDOXIM**

$C_6H_4(OH)C(NO)NH_2$. ***Salcenyl amidoxim*** [99°]. Formed from $C_6H_4(OH)CSNH_2$, hydroxylamine hydrochloride, and Na_2CO_3Aq (Spilker, *B*, 22, 2774). Colourless needles, v sol alcohol and ether, m sol hot water. Gives a greenish pp with $CuSO_4$, and a mirror with $AgNO_3$.

Reactions—1 Ac_2O forms the acetyl derivative and, on further digestion, the azoxim $C_6H_4(OH)C \begin{smallmatrix} N \\ \diagup \diagdown \end{smallmatrix} OMe$ [77°] which yields an acetyl derivative [74°].—2 Concentrated sulphuric acid at 150° yields the sulphonic acid $C_6H_4(SO_3H)(OH)C(NO)NH_2$, v sl sol cold water, insol alcohol.—3 $ClCO_2Et$ yields $C_6H_4(OH)C(NH_2)NOCO_2Et$ [96°] (Miller, *B*, 22, 2799).—4 *Succinic anhydride* forms, on melting, the azoxim $C_6H_4(OH)C(N_2O)CO_2H$, CO_2H [117°].—5. **Potassium cyanate** and HCl yield $C_6H_4(OH)C(NO)NHCO_2NH_2$ [148°].—6,

Phenyl cyanate forms by direct combination $C_6H_5(OH)C(NO_2)NHCO NHPh$ [119°]

Salts— $BHCl$ [175°] Ve sol water — $B_2H_4PtCl_6$ — $Na_2C_6H_4N_2O_2$ — $NaOCH_2N_2O_2$

Acetyl derivative
 $C_6H_5(OH)C(NO_2)NH_2$ [117°] White plates, sl sol water

Di acetyl derivative Formed from the Na salt and $AcCl$ Crystalline

Benzoyl derivative
 $C_6H_5(OH)C(NO_2)NH_2$ [173°] Needles, v. sol ether Yields $C_6H_5(OH)C \begin{smallmatrix} NO \\ N \end{smallmatrix} > CPh$ [128°], which forms a benzoyl derivative [120°]

Di-benzoyl derivative
 $C_6H_5(OBz)C(NO_2)NH_2$ [127°] Formed from the amidoxim, $NaOEt$, and $BzCl$ in ether Insolent crystals

Ethyl ether $C_6H_5(OH)C(NO_2)NH_2$ (278°) Formed from the amidoxim, alcoholic $NaOEt$, and EtI Oil By HCl and $NaNO_2$ it is converted into $C_6H_5(OH)C(NO_2)Cl$ (234°)

Methyl derivative
 $C_6H_5(OMe)C(NO_2)NH_2$ [123°] Formed from $C_6H_5(OMe)CN$ and alcoholic hydroxylamine (Miller, *B* 22, 2791) Needles (from hot water) Yields $BHCl$ [168°] With Ac_2O it yields

$C_6H_5(OMe)C \begin{smallmatrix} NO \\ N \end{smallmatrix} > CMe$ [68°] Aldehyde forms

$C_6H_5(OMe)C \begin{smallmatrix} NO \\ NH \end{smallmatrix} > CHMe$ [127 5°] $ClCO_2Et$ forms $C_6H_5(OMe)C(NH_2)NOCO_2Et$ [120°], which on heating becomes $C_6H_5(OMe)C \begin{smallmatrix} NO \\ NH \end{smallmatrix} > CO$ [208°]

Methyl-acetyl derivative
 $C_6H_5(OMe)C(NO_2)NH_2$ [106°] Formed from the oxim and $AcCl$ in chloroform Prisms

Methyl-benzoyl derivative
 $C_6H_5(OMe)C(NO_2)NH_2$ [148°] White granules, v sol alcohol

Methyl-ethyl derivative
 $C_6H_5(OMe)C(NO_2)NH_2$ [52°] Prisms, v sol. alcohol

Di ethyl derivative
 $C_6H_5(OEt)C(NO_2)NH_2$ (195° at 160 mm) Formed from the amidoxim, $NaOEt$, EtI , and alcohol Oil, miscible with alcohol and ether

m Oxy benzenyl amidoxim
 $C_6H_5(OH)C(NH_2)(NOH)$ [71°] Made by heating *m* oxy-benzonitrile with hydroxylamine hydrochloride and Na_2CO_3 (Clemm, *B* 24, 829) Groups of needles, v sol water

Di-benzoyl derivative
 $C_6H_5(OBz)C(NH_2)NOBz$ [162 5°] Crystals

Acetyl derivative
 $C_6H_5(OH)C(NH_2)NOAc$ [90°] Plates

Di-ethyl ether $C_6H_5(OEt)C(NH_2)NOEt$ [109°] Needles *p*-Oxy benzenyl-amidoxim [158°] Made in like manner (Krone, *B* 24, 834— $BHCl$ [179°])

Acetyl derivative [122 5°]

Benzoyl derivative [166°]

Di-benzoyl derivative [186°]

Di ethyl ether [84°]

***o*-OXY-BENZENYL-*o*-PHENYLENE-DI-**

AMINE $C_6H_5 \begin{smallmatrix} NH \\ N \end{smallmatrix} > COC_6H_5OH$ [223°] Formed by reducing the *o* nitro-anilide of salicylic acid with tin and HCl (Hübner a Mensching, *B* 18, 488; *A* 210, 345) Needles, v. sol. alcohol and

ether — $BHCl aq$ — $B_2H_4SO_4 aq$ sparingly soluble needles

***o*-OXY-BENZENYL-DI-UREA** $C_6H_5N_2O_2$ $1e$ $C_6H_5(OH)CH(NHCO NH_2)_2$ Formed from salicylic aldehyde and aqueous urea (Scheff, *A* 151, 199) Nodular groups of needles (containing aq) V sl sol water, sl sol alcohol, insol ether — $Cu(C_6H_5N_2O_2)_2$ Green pp

Ethyl ether $C_6H_5(OEt)CH(NH_2CO)_2 aq$ Crystals

By fusing salicylic aldehyde with urea there is formed $(C_6H_5(OH)CH)_2(N_2H_2CO)_2$

***p*-Oxy-benzenyl-di-urea** *Methyl ether* $C_6H_5(OMe)CH(NHCO NH_2)_2$ Formed from anisic aldehyde, an aqueous solution of urea, and a little $HOAc$ Yellow plates

DI-OXY-BENZYL *Di-methyl derivative* $C_6H_5(OMe)CO CO C_6H_5(OMe)$ *Anisil* [138°] Formed by oxidising anisoin with alkaline copper solution (Boesler, *B* 14, 327) Golden needles (from alcohol)

Hexa-oxy-benzil *Hexa-methyl derivative* $C_6H_5(OMe)_6 CO CO C_6H_5(OMe)_6$ [189°]

Formed by the action of sodium amalgam on an alcoholic solution of the tri methyl derivative of the amide of gallic acid (Marx, *A* 263, 253) Satiny needles, coloured bluish green by H_2SO_4

***p*-OXY-BENZIMIDO-ETHER** *Ethyl derivative* $C_6H_5(OEt)C(NH)(OEt)$ The hydrochloride is crystalline and is formed from $C_6H_5(OEt)CN$ and alcoholic HCl (Pinner, *B* 23, 2953)

***o*-OXY-BENZOIC ACID** $C_6H_5O_2$ $1e$ $C_6H_5(OH)CO_2H$ *Salicylic acid* Mol w 138 [156°] (Hübner, *A* 162, 74), [159° cor] (Beisert, *B* 23, 2244), [157° cor] (Dunstan a Bloch, *Ph* [3] 21, 429) S 09 at 0° (Ost, *J pr* [2] 17, 230), 15 at 0°, 225 at 15°, 7 925 at 100° (Bourgoin, *J Pharm Chim* [4] 30, 488, *C R* 87, 62), *H C* 734,990 (Berthelot a Recoura, *A Ch* [6] 13, 320), 729,500 (Stohmann, *J pr* [2] 40, 129) *H F* 106,000 (Von Rechenberg), 135,500 (S) Occurs in the blossoms of the meadow sweet (*Spiraea ulmaria*) (Löwig a Weidmann, *P* 46, 83), and in the leaves and stems of *Tulipa*, *Yucca*, and *Hyacinthus* (Griffiths, *C J Proc* 5, 122) Occurs as methyl ether in the oil of wintergreen (from *Gaultheria procumbens*) (Cahours, *A* 48, 60, Hartley, *C J* 53, 664) Methyl salicylate constitutes the essential oils of *Gaultheria punctata* and *G leuco carpa* (Köhler, *B* 12, 246)

Formation—1 By oxidation of *o*-oxy-benzyl alcohol (saligenin), and of *o*-oxy-benzoic aldehyde (salicylic aldehyde)—2 By potash-fusion from salicin, coumarin, indigo, *o*-oresol (Barth, *A* 154, 360), toluene *o*-sulphonic acid, *p*-chlorotoluene sulphonic acid (Vogt, *Z* [2] 5, 577), and other bodies—3 By heating cupric benzoate with water in sealed tubes for 3 hours at 180° (Smith, *Am* 2, 338), *cf* Ettinger, *A* 53, 83)—4 From *o*-amido-benzoic acid by the diazo-reaction (Hübner a Petermann, *A* 149, 129, *cf* Gerland, *A* 86, 147)—5 Together with *p*-oxy benzoic acid by heating phenol with $COCl_2$ and alcoholic potash (Tiemann a Reimer, *B* 9, 1235)—6 By oxidising *o*-tolyl sulphuric acid $C_6H_5MeO SO_3H$ with alkaline $KMnO_4$ (Heymann a Königs, *B* 19, 706)—7 By heating C_6H_5ONa with sodium carbonate in a current of carbonic oxide at 200° $PhONa + Na_2CO_3 + CO$

= $\text{C}_6\text{H}_5(\text{ONa})\text{CO}_2\text{Na} + \text{HCO}_2\text{Na}$ (Schroeder, *A* 221, 41) — 8 By oxidising toluene *o*-phosphonic acid with alkaline KMnO_4 (Heymann a Königs, *B* 19, 8806; — 9 By distilling sodium phenyl carbonate with NaOEt in a current of CO_2 (Hentschel, *J pr* [2] 27, 39) — 10 By heating phenyl ethyl carbonate with NaOPh at 200° in a current of hydrogen, PhOEt being also formed 11. Together with PhOEt by heating Ph_2CO , with NaOEt in a current of hydrogen (*H*)

Preparation — By passing CO_2 over sodium-phenol heated at 180° $\text{C}_6\text{H}_5(\text{ONa})\text{CO}_2\text{Na}$ being formed and phenol distilling over (Kolbe, *A* 115, 201, *J pr* [2] 10, 93) CO_2 is first absorbed, and this can also take place at a lower temperature, and the resulting sodium phenyl carbonate $\text{C}_6\text{H}_5\text{O CO}_2\text{Na}$ changes into the isomeric sodium salicylate $\text{C}_6\text{H}_4(\text{OH})\text{CO}_2\text{Na}$ This change can take place at 130° The sodium salicylate reacts upon excess of sodium phenol at 180° , setting free phenol, which distils over, leaving basic sodium salicylate behind $\text{C}_6\text{H}_4(\text{OH})\text{CO}_2\text{Na} + \text{PhONa} = \text{PhOH} + \text{C}_6\text{H}_5(\text{ONa})\text{CO}_2\text{Na}$ (R Schmitt, *J pr* [2] 31, 404, cf Baumann, *B* 11, 1910) Salicylic acid is set free by acidifying the basic sodium salicylate By using a measured quantity of CO_2 , starting the operation at a low temperature and finishing it at 130° , the formation of basic sodium salicylate may be avoided, so that half the phenol may be saved

Properties — Colourless needles (from hot water) or monoclinic prisms (from alcohol) Sublimes at 200° Volatile with steam FeCl_3 colours its aqueous solution violet, the colour not being removed by acetic acid Prevents pptn of copper sulphate ($\frac{1}{2}$ mol) by alkalis With albuminoids it forms compounds containing about 14 pc of the acid (Farsky, *C C* 1877, 148) Bromine-water yields a pp of $\text{C}_6\text{H}_2\text{Br}_2\text{O}$ in dilute aqueous solutions Iodine and potash give a red powder $\text{C}_6\text{H}_2\text{I}(\text{OI})\text{CO}_2\text{K}$ (Messinger a Vortmann, *B* 22, 2321) Antiseptic Antirheumatic

Reactions — 1 Split up into CO_2 and phenol when rapidly heated to 220° , at 250° diphenylene-ketone oxide $[\text{174}^\circ]$ is formed (Klepl, *J pr* [2] 28, 217) Conc HClAq at 150° decomposes it in like manner (Graebe, *A* 139, 143) Potash-fusion also gives phenol — 2 KMnO_4 oxidises it to formic acid and CO_2 — 3 *Chlorous acid mixture* acts in like manner — 3 *Chlorous* forms chloro-oxy-benzoic and di-chloro-oxybenzoic acids — 4 *Iodine* and HIO_3 give mono, di-, and tri-, iodo-oxy benzoic acids and tri-iodo-phenol — 5 *Nitrous acid* passed into its ethereal solution forms nitro- and diazo oxy-benzoic acids (Goldberg, *J pr* [2] 19, 368) — 6 PCl_5 forms $\text{C}_6\text{H}_4(\text{COCl})\text{O.PCl}_2$ (168° at 11 mm) This compound is partially decomposed on distillation, yielding *o*-chloro-benzoyl chloride Moist air converts it into $\text{C}_6\text{H}_4(\text{CO}_2\text{H})\text{O.P}(\text{OH})_2$ Further treatment with PCl_5 at 170° forms $\text{C}_6\text{H}_4(\text{COCl})\text{OPCl}_2$ (179° at 11 mm) converted by water into $\text{C}_6\text{H}_4(\text{CO}_2\text{H})\text{OP}(\text{OH})_2$ Excess of PCl_5 forms $\text{C}_6\text{H}_4\text{Cl CCl}_2$ (Couper, *A* 109, 370, Anschütz, *A* 228, 314, 239, 314, Chasanovitch, *B* 20, 1166) — 7 PCl_5 forms $\text{C}_6\text{H}_4\text{ClPO}_2$, which may be $\text{C}_6\text{H}_4\text{ClPO}_2$ It forms crystals $[\text{87}^\circ]$ (127° at 11 mm), and is sol ether, benzene, and CHCl_3 , but decomposed by water into salicylic and phosphorous acids (Anschutz a Emery, *A* 239, 301) It is converted by PCl_5 or free chlorine into $\text{C}_6\text{H}_4\text{Cl}_2\text{PO}_2$ (167° at 11 mm), *SG* $\frac{1}{2}$ 1557, whence water forms $\text{C}_6\text{H}_4(\text{OH})_2\text{PO}_2$ [142°] The chloride $\text{C}_6\text{H}_4\text{ClPO}_2$ takes up bromine forming $\text{C}_6\text{H}_4\text{OIBr}_2\text{PO}_2$ (*c* 187° at 12 mm) 8 *Aniline* at 210° produces aniline, phenol, and $\text{C}_6\text{H}_4(\text{OH})\text{CONPh}$ (Limppricht, *B* 22, 2906) — 9 *Cyanamide* and alcohol at 100° form urea and *o* oxy-benzoic ether — 10 *Benzamidine* forms a compound $\text{C}_{11}\text{H}_{11}\text{N}_3\text{O}$ (Pinner, *B* 23, 3824) — 11 *Glycerin* and gaseous HCl forms $\text{C}_6\text{H}_4(\text{OH})\text{CO OC}_2\text{H}_5\text{Cl}_2$ (44°), *SG* 1 331 (Gottig, *B* 24, 508), whence sodic salicylate at 180° – 200° forms crystalline $\text{C}_6\text{H}_4(\text{O CO C}_2\text{H}_5\text{OH})$, and $\text{CH}(\text{OBz})(\text{CH}_2\text{O CO C}_2\text{H}_5\text{OH})$, [95°] (Fritsch, *B* 24, 779) — 12 *Aceto-chlorhydratose* forms $\text{C}_6\text{H}_4\text{O}_3$, [185°], whence $\text{C}_2\text{H}_5\text{O}(\text{OAc})_2$, [111°] (Michael, *B* 15, 1922) — 13 *Phenol* and SnCl_4 at 120° yield *o* di oxy benzophenone [144°] (Michael, *Am* 5, 83) — 14 *Resorcin* at 200° forms tri oxy-benzophenone — 15 *Orcin* gives ' β ' oxy-methyl xanthone' $\text{C}_6\text{H}_4\text{C}(\text{O})\text{C}_6\text{H}_4\text{Me}(\text{OH})$ [285°] (Von Kostanecki a Nessler, *B* 24, 1895) 16 *Phloroglucin* forms iso euxanthone $\text{C}_6\text{H}_4\text{C}(\text{O})\text{C}_6\text{H}_4(\text{OH})_2$ [247°] — 17 When taken internally it is excreted as salicyluric acid or salicyl glycecoll $\text{C}_6\text{H}_5\text{NO}_2$ [*c* 160°] (Bertagnini, *Il Nuovo Cimento*, 1 363) — 18 With *camphor* it forms a compound $\text{C}_{14}\text{H}_{10}\text{O}_2$ [60°] ($[\alpha]_D = +27.3$ (in dilute alcohol) (Léger, *C R* 111, 110) — 19 $\text{K}_2\text{S}_2\text{O}_8$ (17 pts) added to a solution of salicylic acid (10 pts) and KOH (8 pts) in water (25 pts) forms the crystalline compound $\text{C}_6\text{H}_4(\text{CO}_2\text{K})\text{O}8\text{O}_2\text{K}$ (Baumann, *B* 11, 1914) — 20 A solution of salicylic acid and borax in water deposits crystals of $\text{C}_6\text{H}_4\text{NaBO}_3$, whence the corresponding salts $\text{NH}_4\text{A}'$, KA' , MgA' . 10aq and CaA' . 10aq (Jahns, *Ar Ph* [3] 12, 212) — 21 *Chloral* at 140° forms $\text{C}_6\text{H}_4\text{C}(\text{O})\text{CO CH Cl}$, [124°] (Wallach, *A* 193, 1)

Salts — $\text{C}_6\text{H}_5(\text{ONa})\text{CO}_2\text{Na}$ With POCl_3 it gives di phenylene ketone oxide $\text{C}_{10}\text{H}_6\text{O}_2$ [174°] The normal salt $\text{C}_6\text{H}_4(\text{OH})\text{CO}_2\text{Na}$ is converted by POCl_3 into an isomeric body [91°] (R Richer, *J pr* [2] 23, 349, 28, 303) — NaHA' . Large crystals, sol alcohol (E Hoffmann, *Ar Ph* [3] 12, 226) — KA' . 2aq Decomposed at 210° into CO_2 , phenol, and disodium *p* oxy benzoate (Ost, *J pr* [2] 11, 392) In presence of excess of KOH the change does not take place The Na salt does not give *p* oxy benzoic acid when heated alone or with NaOH , but when heated in a current of CO_2 , at 300° it yields oxy-isophthalic and oxytrimelic acids — $\text{NH}_4\text{A}'$ — $\text{NH}_4\text{A}'$. 2aq monoclinic crystals — BaA' . 2aq Small needles, sl sol cold water — $\text{BaC}_6\text{H}_4\text{O}_2$. 2aq Needles, sl sol hot water — SrA' . 2aq — CaA' . 2aq Octahedra, v sol water, sol. alcohol — $\text{CaC}_6\text{H}_4\text{O}_2$. 2aq Crystalline powder, nearly insol water — MgA' . 4aq — ZnA' . 3aq *S* 5 at 20° Needles, sol alcohol (Vulpinus, *Ar Ph* [3] 14, 239) — ZnA' . 2aq (Maignac, *J* 1855, 485) — CdA' . 2aq — $\text{HgC}_6\text{H}_4\text{O}_2$ (Lajoux a Grandval, *J Ph* [5] 20, 5) — HgA' . — PbA' . 2aq — $\text{PbC}_6\text{H}_4\text{O}_2$. — *Oxy-salt*. $\text{Ph}_2\text{O}_4(\text{C}_6\text{H}_4\text{O}_2)_2$ — TlA' — $\text{Ti}_2\text{C}_6\text{H}_4\text{O}_2$ — $\text{BiO}_2\text{A}'$. Insoluble powder got by adding sodium salicylate to a solution of bismuth nitrate in glycerine

(Wolff, *Ph* [3] 14, 508) — AlA'_3 (Van der Velden, *J pr* [2] 15, 151) — FeA'_3 , brown pp., quickly becoming violet — $\text{MnA}'_3 \cdot 2\text{aq}$ — $\text{CuA}'_3 \cdot 4\text{aq}$ bluish-green needles, ν sol water and alcohol. — $\text{CuC}_2\text{H}_3\text{O}_3\text{aq}$ Insoluble — $\text{CuK}_2(\text{C}_2\text{H}_3\text{O}_3)_4\text{aq}$ green tables, ν sol water — $\text{CuBa}(\text{C}_2\text{H}_3\text{O}_3)_4\text{aq}$ (Pellizzari, *G* 14, 365) — AgA' monoclinic needles

Methyl ether $\text{C}_6\text{H}_5(\text{OH})\text{CO}_2\text{Me}$ Mol w 152 (224° cor) SG $\frac{1}{2}$ 1.197 SV 155.9 (Ramsay), 156.7 (Lossen, *A* 254, 64) HF 129.224 (Stohmann, *J pr* [2] 36, 353) Constitutes oil of wintergreen, oil of *Gaultheria punctata* and *leucocarpa*, and oil of birch (Kohler, *B* 12, 246, Pettigrew, *Ph* [3] 14, 167) PCl_5 yields $\text{C}_6\text{H}_5(\text{COCl})\text{OPOCl}_2$ (*u supra*) Phenyl cyanate at 160° forms $\text{C}_6\text{H}_5(\text{CO}_2\text{Me})\text{OCONHPh}$ [238°] (Snape, *C J* 47, 775) Hydroxylamine hydrochloride yields $\text{C}_6\text{H}_5(\text{OH})\text{CO}_2\text{NH}(\text{OH})$ [169°] (Jeaurenaud, *B* 22, 1273) Benzamide forms, on heating, phenyl benzoate and a compound $\text{C}_{12}\text{H}_{12}\text{N}_2\text{O}$ [256° cor] crystallising from chloroform in yellow needles (Guareschi, *A* 171, 143) Forms the crystalline salts $\text{C}_6\text{H}_5(\text{OK})\text{CO}_2\text{Me} \frac{1}{2}\text{aq}$ and $\text{BaA}'_3\text{aq}$

Ethyl ether EtA' (227°) HF 139.252 Oil (Gottig, *B* 9, 1473) With benzamidine hydrochloride it gives $\text{C}_6\text{H}_5\text{O}_3\text{C}_2\text{H}_4\text{NO}$ [120°], benzamidine salicylate $\text{C}_{11}\text{H}_{11}\text{N}_2\text{O}_3$ [202°], and a compound $\text{C}_{21}\text{H}_{21}\text{N}_3\text{O}$ [246°], whence $\text{C}_{21}\text{H}_{21}\text{AcN}_3\text{O}$ [141°] (Pinner, *B* 23, 2935)

Ethylene ether $\text{C}_2\text{H}_4\text{A}'_2$ [83°]

Propyl ether PrA' (239°) SG $\frac{1}{2}$ 1.021 HF 147.880

Isoamyl ether $\text{C}_5\text{H}_{11}\text{A}'$ (270°)

Phenyl ether PhA' *Salol* [42°] Formed by slowly adding POCl_3 (28 g) to a mixture of salicylic acid (60 g) with phenol (48 g) at 135° (Seiffert, *J pr* [2] 31, 472) Anti rheumatic The yield is good (99 g) Trimetric tablets (from alcohol, $a b c = 963.1 \ 697$ (Léger), insol water Its alcoholic solution is coloured violet by FeCl_3 , Conc NaOH forms solid $\text{C}_6\text{H}_5(\text{ONa})\text{CO}_2\text{Ph}$, but boiling NaOHaq saponifies it When heated for a long time to boiling it gives CO_2 , phenol, and diphenylene ketone oxide Dissolved in HOAc it is nitrated by HNO_3 to $\text{C}_6\text{H}_3(\text{NO}_2)_3(\text{OH})\text{CO}_2\text{Ph}$ [150°] and a di nitro-compound [183°] HNO_3 (SG 1.53) forms also $\text{C}_6\text{H}_3(\text{NO}_2)_3(\text{OH})\text{CO}_2\text{Ph}$ [100°] (Knebel, *J pr* [2] 42, 158)

Nitro-phenoxy ethyl ether

$\text{C}_6\text{H}_3(\text{NO}_2)_3\text{OC}_2\text{H}_4\text{A}'$ The *o* isomeride [106°] yields an acetyl derivative [80°] The *p* isomeride crystallises from alcohol in needles [131°] (Wagner, *J pr* [2] 27, 215)

Tolyl ethers $\text{C}_6\text{H}_4\text{MeA}'$ The *o*, *m*, and *p* compounds [35°], [74°], and [39°] are insol water, sl sol alcohol (Nencki, *C R* 108, 254)

Methyl derivatives $\text{C}_6\text{H}_5(\text{OMe})\text{CO}_2\text{H}$ [98.5°] SG $\frac{1}{2}$ 1.1801 μ_d 1.5521 R_{20} 64.59 (Brühl) S 5 at 20° Formed by saponifying its methyl ether, which is formed from oil of Wintergreen, KOme , and MeI (Cahours, *A* 92, 315) Monoclinic tables (from water) Its aqueous solution is not coloured by FeCl_3 . Yields the salts BaA'_3 , $\text{PbA}'_3\text{aq}$, and AgA'_3 , and the ethers MeA' (228°), EtA' (235°) (Fölsing, *B* 17, 486), and PhA' [59°] (Seiffert, *J pr* [2] 31, 474)

Ethyl derivatives $\text{C}_6\text{H}_5(\text{OEt})\text{CO}_2\text{H}$ [19.4°] Forms the salts CaA'_3 , BaA'_3 , $\text{PbA}'_3 \cdot 2\text{aq}$,

$\text{CuA}'_3(\text{OH})$, and AgA' (Kraut, *A* 150.1) and the ethers MeA' (245°) and EtA' (251°)

Isopropyl derivative $\text{C}_6\text{H}_5(\text{OPr})\text{CO}_2\text{H}$ Oil, forming the salts $\text{CaA}'_3 \cdot 2\text{aq}$, $\text{BaA}'_3\text{aq}$, and $\text{AgA}'_3 \cdot \frac{1}{2}\text{aq}$ and the ether MeA' (250°)

Allyl derivative $\text{C}_6\text{H}_5(\text{OC}_2\text{H}_5)\text{CO}_2\text{H}$ [113°] Forms the salt AgA' and the ether MeA' (245°) (Scichilone, *G* 12, 449).

Ethylene derivative $\text{C}_6\text{H}_5(\text{O C}_2\text{H}_4\text{CO}_2\text{H})_2$ [152°] Formed by saponification of its ethyl ether EtA' [97°], which is itself got by heating $\text{C}_6\text{H}_5(\text{ONa})\text{CO}_2\text{Et}$ with ethylene bromide at 130° Silky needles (from water) (Weddige, *J pr* [2] 21, 128)

Phenyl derivative $\text{C}_6\text{H}_5(\text{OPh})\text{CO}_2\text{H}$ [113°] (355°) Formed by adding Na to $\text{C}_6\text{H}_5(\text{OH})\text{CO}_2\text{Ph}$ at 300° (Graebe, *B* 21, 502), and also by the action of phenol on *o*-diazobenzoic acid (Gress, *B* 21, 982) Plates, ν sl sol hot water Heated with conc H_2SO_4 it yields diphenylene ketone oxide Heating with baryta forms diphenyl oxide [25°] Bromine water at 150° forms $\text{C}_6\text{H}_5\text{Br}_2\text{O C}_6\text{H}_5\text{CO}_2\text{H}$ [176°], whence AgA' and EtA' [57°] (Arbenz, *A* 257, 86) HNO_3 at 0° forms a di nitro-derivative [153°], which gives the salts $\text{BaA}'_3 \cdot 4\text{aq}$, $\text{CaA}'_3 \cdot 4\text{aq}$, and AgA'_3 , and the ethers MeA' [126°], EtA' [76°], and an amide [166°] The phenyl derivative of salicylic acid forms the salts $\text{NH}_4\text{A}'$, KA' , $\text{CaA}'_3 \cdot 2\text{aq}$, $\text{BaA}'_3\text{aq}$, and AgA'_3 , and the ether, MeA' and EtA' , boiling above 360°, and PhA' [109°]

Nitro-phenyl ethylene derivative $\text{C}_6\text{H}_3(\text{NO}_2)_3\text{OC}_2\text{H}_4\text{O C}_6\text{H}_5\text{CO}_2\text{H}$ The *o*-acid [142°–148°] forms a crystalline ether EtA' [c 100°], and the *p* acid [132°] forms a similar ether EtA' [81°] (Wagner, *J pr* [2] 27, 214) The *o*-acid yields on reduction an amido acid [110°], forming $\text{C}_6\text{H}_3\text{NO}_2\text{HCl}$ [177°]

Benzyl derivative $\text{C}_6\text{H}_5\text{CH}_2\text{O C}_6\text{H}_5\text{CO}_2\text{H}$ [75°] Tables (Perkin, *A* 148, 27) — AgA'

Tolyl derivatives Occurs in natural salicylic acid (J Williams, *Ph* [3] 8, 785)

Acetyl derivative $\text{C}_6\text{H}_5(\text{OAc})\text{CO}_2\text{H}$ [118°] Formed from the acid and AcCl (Kraut, *A* 150, 9) Needles (from water), ν sl sol cold water Hydrolysed by alkalis, not by boiling water

Benzoyl derivative $\text{C}_6\text{H}_5(\text{OBz})\text{CO}_2\text{H}$ The crystalline methyl ether MeA' is formed from methyl salicylate and BzCl (Gerhardt, *A Ch* [3] 45, 104)

Amide $\text{C}_6\text{H}_5(\text{OH})\text{CONH}_2$ [139°] From the ethers and NH_3aq (Lumprecht, *A* 98, 258) Formed by the action of AcCl on *o*-oxy benzal oxim (Claisen, *A Stock*, *B* 24, 138) Plates and tables Gives saligenin on reduction with sodium amalgam (Hutchinson, *C J* 57, 957) With bromine water it yields $\text{C}_6\text{H}_3\text{Br}_2(\text{OH})\text{CONH}_2$ [183°] (Spilker, *B* 22, 2769) When heated in a current of HCl it forms $(\text{C}_6\text{H}_5(\text{OH})\text{CO})_2\text{NH}$ [199°], which yields $(\text{C}_6\text{H}_5\text{O})_2\text{Nag}$ and $(\text{C}_6\text{H}_5\text{NO}_2)_3\text{HCl}$ — AgA' flocculent pp

Benzoyl derivative of the amide $\text{C}_6\text{H}_5(\text{OBz})\text{CONH}_2$ [200°] Needles, sl sol alcohol

Cumyl derivative of amide [200°] *Methyl derivatives of the amide* $\text{C}_6\text{H}_5(\text{OMe})\text{CONH}_2$ [129°] Prisms (from ether) (Grimaux, *Bl* [2] 18, 26) The homologous ethyl derivative [110°] (Lumprecht, *A*

98, 264) and isopropyl derivative are crystalline

Anilide $C_6H_5(OH)CONHPh$ [184°] Prisms (from dilute alcohol) (Wanstrat, *B* 6, 336, Kupferberg, *J pr* [2] 18, 442, Hübner a Meusching, *A* 210, 541) $FeCl_3$ colours its alcoholic solution violet. On heating with sulphur

it forms $C_6H_5\langle\begin{smallmatrix} N \\ S \end{smallmatrix}\rangle C_6H_5OH$ [129°] (Hofmann, *B* 18, 1237) Yields on nitration $C_6H_5(NO_2)(OH)CONHPh$ [224°] Forms the salts $KC_6H_4NO_2$, $2\frac{1}{2}aq$ and $TlO_2C_6H_4NO_2$.

Nitro-anilide $C_6H_5(OH)CO NHOC_6H_4NO_2$. The *o*-, *m*-, and *p*-varieties melt at 154°, 218°, and 230° respectively

***p*-Toluide** $C_6H_4(OH)NHC_6H_5$ [156°]

Piperidine $C_6H_5(OH)CONHC_6H_5$ [142°] Yellowish tables (Schotten, *B* 21, 2252)

Hydroxylamide $C_6H_5(OH)CONH(OH)$. [169°] Needles Yields $Pb(C_6H_5NO_2)_2 \cdot 4aq$ (Jeanrenaud, *B* 22, 1270)

Anhydride $O(C_6H_4CO_2H)_2$, **Disalicylic acid** Formed, together with salicylide, by the action of $POCl_3$ on dry sodium salicylate (Gerhardt, *A. Ch* [3] 87, 322) Amorphous mass, *v* sol alcohol and ether Gives no colour with $FeCl_3$. Dissolves unchanged in aqueous Na_2CO_3 . Boiling KOH aq converts it into salicylic acid An anhydride $C_{12}H_8O_5$, is formed by heating sodium salicylate (3 pts) with $POCl_3$ (1 pt) at 150° (Kraut, *A* 150, 13) It is insol cold alcohol A third anhydride $C_{12}H_8O_5$, is got by heating $C_6H_5(OAc)CO_2H$ at 200°–240° (Kraut) It is sol alcohol, and softens at 70°

Internal anhydride $C_6H_4O_2$ or $C_6H_4O_2 \cdot s$. $C_6H_5\langle\begin{smallmatrix} O \\ CO \end{smallmatrix}\rangle C_6H_5$, **Salicylide** [195°–200°]

Formed as above Nodular groups of plates (from alcohol) (Schiff, *A* 163, 220) Insol water, *sl* sol alcohol Not attacked by $AcCl$ Reconverted by potash into salicylic acid A resinous anhydride $C_{12}H_8O_5$, accompanying salicylide is still less soluble in alcohol

Nitrile $C_6H_5(OH)CN$. ***o*-Cyano-phenol** [98°] Formed by the action of P_2O_5 or P_2S_5 on the amide, and also from $C_6H_5(OH)CHNOH$ and Ac_2O (Miller, *B* 22, 2771, 2797, Tiemann, *B* 20, 3082, Meyer, *B* 20, 3289, cf Grimaux, *Bl* [2] 13, 26, Ahrens, *B* 20, 2953) Formed also from diazophenol chloride by Sandmeyer's reaction In most of these preparations it is accompanied by a small quantity of a substance melting at 195° Needles, *v* sol alcohol, *m* sol cold water Coloured violet by $FeCl_3$ Gives a white crystalline pp. with bromine water Gives a **methyl derivative** $C_6H_5(OMe)CN$ (256°), and an **ethyl derivative** $C_6H_5(OEt)CN$ (258°) which may be got from $C_6H_5(OEt)NH_2$ (Pinner, *B* 23, 2952) The **acetyl derivative** $C_6H_5(OAc)CN$ is an oil (258°) (Lach, *B* 17, 1572) while the **benzoyl derivative** $C_6H_5(OBz)CN$ [149°] is crystalline (Lamprecht, *A* 99, 250, Henry, *B* 2, 491)

Polynitrile $(C_6H_5(OH)CN)_n$ [296°–299°] Obtained by heating the amide to 270° (*L*) and as a by-product in preparing the nitrile (Miller, *B* 22, 2798) Yellow needles, *msol* alcohol, *sl* sol ether. Decomposed by HCl at 200° into CO_2 , phenol, and NH_3 .

***m*-Oxy-benzoic acid** $C_6H_5(OH)CO_2H$ [200°]. **S**. 37 at 0° (Ost), 9 at 18° (Fittica, *B* 11, 1208).

H F 111,000 (Von Rechenberg); 136,000 (Stohmann, *J pr* [2] 40, 129) **F C** 729,000

Formation —1 By the action of nitrous acid on amido benzoic acid (Gerland, *A* 91, 185, Graebe a. Schultzen, *A* 142, 350) —2 By fusing sulpho-benzoic acid with potash (Barth, *A* 148, 80) —3 By potash fusion from *m* chloro benzoic acid (Dembey, *A* 148, 222), *m* cresol, and even benzoic acid (Barth, *A* 154, 361, *M* 3, 802)

Properties —Nodular groups of needles (from water) May be distilled *V* sol boiling water and alcohol Volatile with steam Blackens when quickly heated to 300°, but gives no phenol When the acid (2 mols) is heated with baryta (3 mols) at 350° it forms no phenol unless the baryta is used in large excess (7 mols) (Klepl, *J pr* [2] 27, 159) It tastes sweet Its solutions are not coloured by $FeCl_3$. Iodine and KOH aq, followed by an acid, give a coffee-brown pp (Messinger, *B* 22, 2321) It is not acted upon by hydroxylamine

Reactions —1 **Bromine** (3 mols) forms tribromo-*m*-oxy benzoic acid, soluble in water (Werner, *Bl* [2] 46, 276) —2 **Sodium amalgam** reduces it, in acid solution, to *m* oxy benzoic alcohol —3 Conc H_2SO_4 forms, on heating, several di oxy anthraquinones (*q v*) When benzoic acid is also present, the two oxy anthraquinones are also formed —4 Heated with cinnamic acid and H_2SO_4 , it yields anthracoumarin $C_{12}H_8O_2$ [260°] (*v* CINNAMIC ACID, Reaction 14) —5 By nitration with dilute nitric acid, which takes place extremely readily, it yields a mixture of three nitro oxy-benzoic acids $C_6H_5(NO_2)(OH)CO_2H$ [4 3 1], [2 3 1], and [5 3 1] (Griess, *B* 20, 403) —6 PCl_5 forms the chloride $C_6H_5(COCl)OPOCl_2$ of *m*-carboxy phenyl phosphoric acid (*q v*) —7 The *K* salt heated with $K_2S_2O_8$ forms $SO_3K O C_6H_4 CO_2K$ [220°–225°] (Baumann, *B* 11, 1915) —8 When taken internally, it appears in the urine as oxybenzuroic acid $C_6H_5(OH)CO NH CH_3 CO_2H$, crystallising in needles (Baumann a. Herter, *H* 1, 260)

Salts. — NH_4A' needles, *v* sol cold water — CaA' , $3aq$ *m* sol water — BaA' , gummy — TlA' — $Tl_2C_6H_4O_2$ — PbA'_2 — CdA'_2 — CuA'_2 , *aq* green needles

Acetyl derivative $C_6H_5(OAc)CO_2H$ [127°]

Ethyl ether EtA' [72°] (282°) Tables (from water), nearly insol cold water Conc $NaOH aq$ forms crystalline $C_6H_5(ONa)CO_2Et$

Methyl derivative $C_6H_5(OMe)CO_2H$ [107°] Formed from its methyl ether, and also by oxidising OH , $C_6H_5(OMe)$ with $KMnO_4$ (Oppenheim a. Pfaff, *B* 8, 887) Formed also by the action of CO_2 and Na on $C_6H_5Br(OMe)$ (Körner, *Bull Acad Belg* [2] 24, 155) and by boiling the sulphate of *m*-diazobenzoic acid with $MeOH$ (Griess, *B* 21, 979) White needles, *v* *sl* sol cold water It forms the salts CaA'_2 , *aq* and AgA'_2 and the ether $C_6H_5(OMe)CO_2Me$ which is formed by heating *m*-oxybenzoic acid with KOH and MeI at 140°

Ethyl derivative $C_6H_5(OEt)CO_2H$ [137°] Formed from its ethyl ether $C_6H_5(OEt)CO_2Et$ (263°) (Hantz, *A* 153, 331) or by boiling the sulphate of diazo-benzoic acid with alcohol (Fittica, *B* 11, 1209, Griess, *B* 21, 979) Needles. Yields CaA'_2 , $2aq$, BaA'_2 , $2aq$, and AgA'_2 .

Allyl derivative $C_6H_4(OC_2H_5)CO_2H$ [148°] Colourless laminae Its ethyl ether is a thick pungent oil (284°) (Seichilone, *G* 12, 449)

Phenyl derivative $C_6H_4(OPh)CO_2H$ [145°] Formed by the action of phenol on the sulphate of *m*-diazobenzoic acid (Griess, *B* 21, 980) Needles, almost insol hot water Yields BaA' , 3:5aq

Amide $C_6H_4(OH)CONH_2$ [167°] Formed from the ether and conc NH_3 Aq (Schulerud, *J pr* [2] 22, 290) Thin plates (from water), sol alcohol and ether, insol chloroform

Anilide $C_6H_4(OH)CONHPh$ [155°] Needles, insol water (Kupferberg, *J pr* [2] 16, 442)

Nitrile $C_6H_4(OH)CN$ [82°] Formed by the diazo-reaction from $C_6H_4(NH_2)CN$ by displacing NH_2 by OH (Griess, *B* 8, 859) or from $C_6H_4(NH_2)OH$ by displacing amidogen by Cy (Ahrens, *B* 20, 2953) Plates (from water) with intensely sweet taste Its acetyl derivative $C_6H_4(OAc)CN$ [60°] is got by boiling *m*-oxybenzaloxim with Ac_2O (Clemm, *B* 24, 827)

Anhydride $C_6H_4O_2$ Formed, together with an anhydride $C_6H_4O_2$, [165°] by the action of $POCl_3$ at 45° on *m*-oxy benzoic acid (Schiff, *B* 15, 2588) Minute crystals, sol boiling alcohol

Tetra-hydrate $CH_2<\begin{smallmatrix} CO & CH_2 \\ CH & CH_2 \end{smallmatrix}>CHCO_2H$

Formed by warming the tetrahydrate of oxyterephthalic acid with water (Baeyer & Lutein, *B* 22, 2183) Mixes with water Forms the hygroscopic salt NaA' crystallising in slender needles Yields an oxim $C_6H_4NO_2$ [170°] and a phenyl hydrazide $C_6H_4N_2O_2$ [125°]

***p* Oxy benzoic acid** $C_6H_4(OH)CO_2H$ [213°] *S* 173 at 0° (Ost, *J pr* [2] 17, 230), *S* at 15° (Saytzeff) *H F* 113,000 (Von Rechenberg), 139,100 (Stohmann, *J pr* [2] 40, 130) *H C* 725,900

Formation—1 By heating anisic acid with conc $HIAq$ at 130° for 12 hours (Saytzeff, *A* 127, 129)—2 By the action of nitrous acid on *p* amido benzoic acid (*G Fischer*, *A* 127, 145) 3 By potash fusion from *p* sulpho benzoic acid (Remsen, *Z* [2] 7, 81, *A* 178, 281), anethol (Ladenburg, *A Suppl* 8, 87), anisic acid (Barth, *Sitz W* 54 [2] 633), gum benzoin, acaroid resin (Hlasiwetz & Barth, *A* 134, 265, 138, 61), tyrosine (Barth, *A* 136, 110, Ost, *J pr* [2] 12, 159), carthamin (Malin, *A* 136, 115), phloretic acid, *p* cresol, and even benzoic acid (Barth, *A* 152, 96, 154, 359, 164, 141, *M* 3, 802)—3 By passing CO_2 through C_6H_4OK dissolved in boiling phenol, or, better, by heating C_6H_4OK in a current of CO_2 at 170°–210° (Kolbe, *J pr* [2] 8, 336, 10, 89, 451, 11, 24, Ost, *J pr* [2] 11, 385, Hartmann, *J pr* [2] 16, 35) At 130°–150° the product is salicylic acid—4 The basic salt $C_6H_4(OK)CO_2K$ is formed, together with CO_2 and phenol, by heating potassium (but not sodium) salicylate at 220° *A* mixture of salicylic acid (1 mol) and excess of KOH (8 mols.) is not affected at 250°, but at 300° yields only K_2CO_3 and C_6H_4OK —5 Together with a smaller quantity of salicylic acid by heating phenol with alcoholic potash (or soda) and CCl_4 The reaction takes place less quickly in aqueous solution (Tiemann & Reimer, *B* 9, 1285; Hasse, *B* 10, 2186)

Properties—Monoclinic crystals (containing aq), $a b c = 1.370 : 1.102$, $\beta = 105^\circ 26'$ *V* sol. hot water, alcohol, and ether, sl sol chloroform (unlike salicylic acid) and CS_2 (unlike benzoic acid) Split up at 220° into CO_2 and phenol With iodine and potash it gives a pale-red pp., which becomes yellow on acidifying (Messinger & Vortmann, *B* 22, 2321) Not attacked by hydroxylamine With $FeCl_3$ it gives a yellow amorphous pp Excess of bromine water gives CO_2 and tri-bromo phenol

Reactions—1 *p*-Oxy benzoic acid (1 mol) is converted by PCl_5 (1 mol) into $C_6H_4Cl_2PO_2$, *z* $C_6H_4(COCl)OPOCl_2$ (176° at 14 mm) *S G* 1542 This chloride shows the following reactions (a) Water forms *p* carboxy-phenyl phosphoric acid $C_6H_4(CO_2H)OPO(OH)_2$ which forms plates [200°], *v* sol water, alcohol, and ether It is not decomposed by boiling aqueous KOH , but water at 160° forms phosphoric acid and *p*-oxy benzoic acid (b) Distillation under atmospheric pressure somewhat decomposes it (c) The chloride (1 mol) heated with PCl_5 (1 mol.) at 160° forms *p* chloro benzoyl chloride, $C_6H_4ClCOCl$ (Anschutz & Moore, *A* 239, 342) 2 On distillation half of it splits up into phenol and CO_2 , the rest yields several anhydrides—3 The *K* salt when distilled yields diphenylene oxide, di-phenylene-ketone oxide, and phenol (Goldschmidt, *M* 4, 127)—4 Yields 50 or 60 pc of the theoretical amount of phenol on fusion with $NaOH$ (Barth & Schreder, *B* 12, 1257)—5 The *Na* salt heated in a current of CO_2 at 290° yields salicylic acid (Kupferberg, *J pr* [2] 16, 424)—6 The *Ca* salt on dry distillation yields phenol, CO_2 , salicylic acid, oxyisophthalic acid, diphenylene oxide, and diphenylene ketone oxide—7 H_2SO_4 at 100° forms $C_6H_4(OH)(SO_3H)CO_2H$ (Klepl, *J pr* [2] 23, 196)—8 The *K* salt heated with $K_2S_2O_8$ forms $C_6H_4(COO.K)O SO_3K$ (Baumann, *B* 11, 1916)—9 Taken internally it appears in the urine as *p* oxy benzoic acid $C_6H_4NO_2$ [c 228°]

Salts— NaA' , 5aq very soluble efflorescent plates— $Na_2C_6H_4O_2$ — KA' , 3aq— NH_4A' aq long efflorescent prisms— CaA' , 4aq slender needles— BaA' , aq flat needles— BaA' , 2aq rhombohedra— $BaC_6H_4O_2$, sandy powder— TiA' — CdA' , 4aq— CdA' , 6aq— ZnA' , 8aq— PbA' , 2aq CuA' , 6aq— AgA' , 2aq

Acetyl derivative $C_6H_4(OAc)CO_2H$ [185°] By heating the acid with Ac_2O Silvery plates (from $CHCl_3$)

Methyl ether MeA' [117°] (283°). *H F* 138,800 Formed from the acid, KOH , and MeI (Ladenburg & Fitz, *A* 141, 250) Large tables (from ether), *v* sl sol hot water

Ethyl ether EtA' [112.5°] (*G*), [116°] (Hartmann) (298°) *H F* 147,690 Formed from the acid, alcohol, and HCl (Graebe, *A* 139, 134). Crystalline Yields solid $C_6H_4(ONa)CO_2Et$

Phenyl ether PhA' [176°] Present in the volatile product of the destructive distillation of *p* oxy-benzoic acid (Klepl, *J pr* [2] 28, 214) Trimetric tablets (from chloroform) Saponified by cold $NaOHAq$ With alcohol and HCl it yields phenol and *p* oxy benzoic acid The acetyl derivative $C_6H_4(OAc)CO_2Ph$ crystallises in long plates [84°]

Methyl derivative $C_6H_4(OMe)CO_2H$. *Anisic acid* Mol *w* 152 [184°]. (275°–280°).

S 04 at 18° H C p 895,200 H F 132,800 (Stohmann, *J pr* [2] 40, 181) Formed by oxidation of anise-camphor, and of oils of anise, fennel, and tarragon, being derived from the anethol contained therein (Cahours, *A Ch* [3] 2, 287, 14, 483, 23, 351, 25, 21, 27, 439, Laurent, *Rev scient* 10, 6, 362, Gerhardt, *A Ch* [8] 7, 292, Ladenburg, *A* 141, 241) Obtained also from its methyl ether, which is formed by heating *p* oxy benzoic acid (1 mol) with KOH (2 mols) and MeI (2 mols) at 120° (Ladenburg) It is a product of the oxidation of chics (Erdmann, *J pr* 71, 198) It is also produced by oxidising $C_6H_5Me(OMe)$ (Körner, *Bl* [2] 10, 468) and by boiling the sulphate of *p* diazobenzoic acid with MeOH (Griess, *B* 21, 979) Prepared by mixing basic potassic *p* oxybenzoate (got by heating potassic salicylate at 220°, or by adding KOH to a solution of *p* oxybenzoic acid) with a solution of $KMeSO_3$ and evaporating to dryness The residue is treated with HCl, and the anisic acid separated from undecomposed *p* oxybenzoic acid by solution in chloroform (E v Meyer a P Richter, *J pr* [2] 32, 429) Monoclinic prisms, m sol hot water Yields, on nitration, $C_6H_3(NO_2)_3OMe$, $C_6H_3(NO_2)_2OMe$, and $C_6H_2(NO_2)_2OMe$ HIAq converts it into MeI and *p*-oxy-benzoic acid (Graebe, *A* 139, 148) When taken internally it passes into the urine as anisuric acid (*q v*) $POCl_3$ forms the anhydride $C_6H_4O_2$ [99°] (Pisani, *A* 102, 284) PCl_5 forms crystalline $C_6H_3(OMe)COCl$ Forms the salts NH_4A' , KA' , NaA' , AgA' , BaA' , SrA' , aq , CaA' , aq , MgA' , aq , PbA' , aq , ZnA' , $3aq$, CdA' , $3aq$, $Pb(OH)A'$, Cr_2A' , $(OH)_2$, MnA' , $3aq$, CoA' , $3aq$, NiA' , $3aq$, CuA' , $3aq$, $CuA'(OH)$, and AgA' (Borrella, *G* 15, 303) Its ethers are MeA' [47°], (255°) and EtA' (c 253°) Its amide $C_6H_4(OMe)CONH_2$ [163°] is formed by the action of NH_3 on $C_6H_4(OMe)COCl$ It is also formed by passing cyanic acid vapour and dry HCl through $C_6H_4(OMe)$ containing $AlCl_3$, and by the action of $ClCONH_2$ and $AlCl_3$ on C_6H_4OMe dissolved in CS_2 (Gattermann, *A* 244, 62, *B* 23, 1197) It crystallises from water in needles or plates The anilide $C_6H_4(OMe)CONPhH$ [169°] is formed by the action of phenyl cyanate on anisole containing $AlCl_3$ (Leuchart a Schmidt, *B* 18, 2338) The nitrile $C_6H_4(OMe)CN$, [62°], (254°), is formed by heating the amide alone or with PCl_5 (Henry, *Z* [2] 6, 209, *B* 2, 667), and by heating $C_6H_4(OMe)CHNOH$ with $AcCl$ at 115° (Miller, *B* 22, 2791) It crystallises in needles, v sol alcohol and ether Hydroxylamine converts it into $C_6H_4(OMe)C(NH_2)NOH$ [123°]

Ethyl derivative $C_6H_4(OEt)CO_2H$ [195°] Formed from its ether EtA' (275°) which is got from *p*-oxy benzoic acid, KOH, and EtI (L a F) Got also by oxidising the ethyl derivative of phloretic acid with chromic acid mixture (Körner a Corbetta, *B* 7, 1781), and by boiling the sulphate of *p* diazo-benzoic acid with alcohol (Griess, *B* 21, 980) Needles Yields AgA' crystallising in needles The amide $C_6H_4(OEt)CONH_2$ [202°] (G), [206°] (P), is formed by the action of cyanic acid or $ClCONH_2$ on C_6H_4OEt in presence of $AlCl_3$ (Gattermann, *A* 244, 68, *B* 23, 1197), and by adding $NaOHAq$ to *p*-ethoxy-benzamidine hydrochloride (Pinner, *B* 28, 2954) The anilide $C_6H_4(OEt)CONPhH$ [170°] is formed by the action of phenyl cyanate

on phenetole in presence of $AlCl_3$ (L a S) The nitrile $C_6H_4(OEt)CN$ [69°] (258°), is got from $C_6H_4(OEt)NH_2$ by Sandmeyer's reaction (Pinner, *B* 23, 2953) It is volatile with steam

Ethylene derivative The amide $C_6H_4(OC_2H_4CONH_2)_2$ [280°] is formed by the action of $ClCONH_2$ on $C_6H_4(OPh)_2$ in CS_2 , in presence of $AlCl_3$ (Gattermann, *A* 244, 69)

Allyl derivative $C_6H_4OC_2H_4CO_2H$ [123°] Formed from its ether EtA' [109°] (260°) which is got by heating *p* oxy-benzoic ether with KOH and allyl iodide at 120° (Seichilone, *G* 12, 451)

Phenyl derivative $C_6H_4(OPh)CO_2H$ [160°] Obtained by the action of phenol on the sulphate of *p* diazo benzoic acid (Griess, *B* 21, 980), and also by the action of boiling alcoholic potash upon $C_6H_4(OPh)CO_2Ph$, a white sublimate [73°-78°] got by strongly heating *p* oxybenzoic in a current of hot CO_2 (Klepl, *J pr* [2] 28, 200)

Phenoxy-ethyl derivative $C_6H_4(OPh)OC_2H_4CO_2H$ [196°] Satiny needles (from alcohol) (Wagner, *J pr* [2] 27, 227) Its ether EtA' [81°] is crystalline

Nitro-phenoxy-ethyl derivative $C_6H_4(NO_2)OC_2H_4OC_2H_4CO_2H$ The *o*-compound [207°] forms an ether EtA' [103°] crystallising from alcohol in plates, and may be reduced to $C_6H_4(NH_2)OC_2H_4OC_2H_4CO_2H$ [155°] The *p*-isomeride [218°] forms a salt NaA' 3aq and an ether EtA' [181°] crystallising in minute needles

Amide $C_6H_4(OH)CONH_2$ [162°] Needles (containing aq) Forms the sodium compounds $C_6H_4(ONa)CONH_2$ and $C_6H_4(ONa)CONH_2Cl$ [206°] Yields *p* oxy benzyl alcohol on reduction with sodium amalgam (Hutchinson, *B* 24, 175)

Anilide $C_6H_4(OH)CONPhH$ [197°] Yellow plates, v sol alcohol

Piperidine $C_6H_4(OH)CONC_4H_{10}$ [210°] Prisms (from dilute alcohol) (Schotten, *B* 21, 2254)

Nitrile $C_6H_4(OH)CN$ *p*-Cyanophenol [113°] Formed by distilling ammonium *p* oxy benzoate with P_2O_5 (Hartmann) Formed also from *p* amido phenol by Sandmeyer's reaction (Ahrens, *B* 20, 2954), and by the action of NH_3 on *p*-oxy benzide Thin trimetric laminae, $a b c = 855 \ 12 \ 308$ M sol hot water Forms an acetyl derivative $C_6H_4(OAc)CN$, [57°], (266°), crystallising in white needles

Anhydride $C_6H_4O_2$ *p*-Oxybenzide Left in the retort after distilling *p*-oxy benzoic acid below 350° (Klepl, *J pr* [2] 25, 525, 28, 194) White amorphous powder, blackening at 350° without melting Insol alcohol Reconverted into *p* oxy-benzoic acid by boiling $KOHAq$, not attacked by NH_3 , or Na_2CO_3Aq Heated in sealed tubes with PCl_5 it yields $C_6H_4Cl_2CCl_2$

Anhydride $C_6H_4O_2$ *o*-Cyanophenol $CO_2H.C_6H_4.O.CO.C_6H_4.O$ [261°] A product of the action of heat on *p* oxy benzoic acid Minute needles, v sol alcohol Quickly converted by alkalis into *p*-oxy-benzoic acid Yields NaA' , BaA' , and $C_6H_4AcO_2$, [217°]

Anhydride $C_6H_4O_2$ *o*-Cyanophenol $CO_2H.C_6H_4.O.CO.C_6H_4.OH$ [280°] 8 (alcohol) 45 in the cold, 13 at 78° Accompanies *p* oxybenzide. Crystalline powder Con

verted by potash into *p* oxy benzoic acid Gives NaA' and $C_6H_5(OAc)_2$ [230°]

Anhydride $C_6H_4O_3$. Formed from the acid and $POCl_3$ (Schiff, *B* 15, 2588) Insoluble powder:

Di-oxy benzoic acid $C_6H_4O_4$, 1c
 $\lambda_{max}(OH)_2CO_2H$ [3 2 1] *Pyrocatechin carboxylic acid* Mol w 154 [204°] Formed in small quantity, together with protocatechuic acid by heating pyrocatechin (1 pt) with ammonium carbonate (4 pts) and water (5 pts) at 140° (A Miller, *C J* 41, 398, *A* 220, 116) Formed also by heating iodosalicylic acid with KOH Needles (containing 2aq), m sol water, v sol alcohol and ether $FeCl_3$ gives a blue colour not destroyed by excess, but changed to violet red by Na_2CO_3 Gives a flocculent ppt with $Pb(OAc)_2$ — BaA', 5aq prisms (S of BaA') 1 at 18°

Isomeride of PROTOCATECHUIC ACID

Di oxy benzoic acid $C_6H_4(OH)_2CO_2H$ [5 3 1] (a) *Resorcylic acid* [222°] (B a S), [233°] (B) Formed by fusing 3 di sulpho benzoic acid with potash (Barth a Senhofer, *A* 159, 222) Formed also from bromo sulpho benzoic acid by potash-fusion (Böttinger, *B* 8, 374) Prisms or needles (containing 1½ aq), m sol cold water Gives no colour with $FeCl_3$, Conc H_2SO_4 at 140° forms a red solution whence water ppts green flakes of anthrachrysone $C_{14}H_8O_4$ Yields resorcin on fusion with potash

Salts — NaA' aq — BaA', 4aq — CuA', 6½aq — CdA', 4½aq — AgA' aq crystalline pp

Ethyl ether EtA' [below 100°] Prisms

Methyl ether of the methyl derivative $C_6H_4(OH)(OMe)CO_2Me$ (315°) Formed, together with $C_6H_4(OMe)_2CO_2Me$ from 3 di oxy benzoic acid, MeI, and KOH (Meyer, *M* 8, 430) Oil

Di methyl derivative $C_6H_4(OMe)_2CO_2H$ [176°] Formed by methylation and also by oxidation of the di methyl ether of orcin (Tiemann a Streng, *B* 14, 2002) White needles, sol hot water — AgA' crystalline pp

Methyl ether of the di methyl derivative $C_6H_4(OMe)_2CO_2Me$ [81°] (298°) Four sided prisms (M)

Di ethyl derivative $C_6H_4(OEt)_2CO_2H$ [88°] Prisms Forms only $C_6H_4(OEt)_2CO_2Et$

Di oxy benzoic acid $C_6H_4(OH)_2CO_2H$ [4 2 1] (B) *Resorcylic acid* [205°] S 26 at 17° H F 188, 100 H Cp 676, 900 (Stohmann, *J pr* [2] 40, 132)

Formation — 1 From $C_6H_5Me(OH)(SO_3H)$ [1 2 4] by heating with KOH (Ascher, *A* 161, 11) 2 From toluene disulphonic acid by oxidation and potash fusion (Blomstrand, *B* 5, 1088, Fahlberg, *Am* 2, 196) — 3 By oxidation of its aldehyde or of umbelliferone (Tiemann a Reimer, *B* 12, 997, 13, 2358) — 4 By heating resorcin with ammonium carbonate and water at 125° (Brunner a Senhofer, *B* 13, 2356) — 5 By oxidising morin with HNO_3 (Benedict a Hazura, *M* 5, 170) — 6 By warming $C_6H_4(OH)_2CS_2H$ with acid (Lippmann, *M* 9, 306, 10, 620)

Preparation — 20 pts of resorcin are heated for an hour and a half with a solution of 100 pts of potassium or sodium hydric carbonate in 200 grms of water, the yield is 80 p.c of the resorcin (Bistrzycki a Kostanecki, *B* 18, 1984)

Properties — Crystallises from ether in needles (containing 8aq) and from water in

prisms (containing ¼aq, 1½aq, or 2½aq) Decomposes at its melting point into CO_2 and resorcin $FeCl_3$ colours its solution dark rose red Bleaching powder gives a violet tint, changing to brown By treating the acid with $C_6H_4(OH)_2CO_2H$ [5 2 1] and Ac_2O and distilling the product there is formed euxanthone $[4 \frac{1}{2}] C_6H_4(OH) < \overset{CO}{O} > C_6H_4(OH) [2 \frac{5}{2}]$ (Graebe, *B* 22, 1405)

Salts — KA' aq — BaA', 4aq — BaA', 7aq — CuA', 8aq — AgA'

o Methyl derivative

$C_6H_4(OH)(OMe)CO_2H$ [4 2 1] Formed by oxidising $C_6H_5(OAc)(OMe)CHO$ (Tiemann a Parisius, *B* 13, 2354) Crystalline Sol water. Gives no colour with $FeCl_3$

p Methyl derivative

$C_6H_4(OMe)(OH)CO_2H$ [4 2 1] [154°] S 7 at 20° Got by partial methylation of the acid (T a P), and also by the action of CO_2 on $C_6H_4(ONa)(OMe)$ at 215° (Korner a Bertoni, *Rendiconti d R Istit Lombardo*, 13, 741, *B* 14, 847) Needles, sol hot water Gives a reddish-violet colour with $FeCl_3$ — NaA' aq — KA' — BaA', 4aq — PbA', 2aq

Di-methyl derivative $C_6H_4(OMe)_2CO_2H$ [108°] Got by methylation (T a P) and by oxidation of the di methyl derivative of (B) methyl umbellie acid (Pechmann, *B* 16, 2126, 17, 2133) Needles, sl sol cold water — CuA', — PbA' — AgA' white pp

Di ethyl derivative $C_6H_4(OEt)_2CO_2H$ [99°] Got from the aldehyde (Tiemann a Lewy, *B* 10, 2215)

Di oxy benzoic acid $C_6H_4(OH)_2CO_2H$ [6 2 1] [c 147°] Formed, together with the (4, 2, 1)-isomeride, by heating resorcin with ammonium carbonate and water (B a S) Obtained also from the methyl derivative of the nitrile (Lobry de Bruyn, *R T C* 2, 205) Needles Decomposes on fusion into CO_2 and resorcin $FeCl_3$ gives a violet colour, changed to blue by excess. Bromine water gives tri bromo resorcin — BaA' aq — CuA', 8aq — AgA' crystalline pp

Di-methyl derivative $C_6H_4(OMe)_2CO_2H$ [179°] Tables (from alcohol)

Nitrile of the di-methyl derivative $C_6H_4(OMe)_2CN$ [118°] (310°) Formed from $C_6H_4(OMe)(NO_2)CN$ by boiling with MeOH and KOH Crystals Gives with nitric acid a nitro-compound $C_6H_4(NO_2)_2CN$ [111°] The corresponding nitriles $C_6H_4(OMe)(OEt)CN$ [66°] and $C_6H_4(OEt)_2CN$ [122°] crystallise from alcohol, the former in trimetric crystals, $abc = 796 \cdot 1 \cdot 1 \cdot 65$, and the latter in di metric crystals $a = 1 \cdot 565$ (Lobry de Bruyn, *R T C* 3, 383)

Di oxy benzoic acid $C_6H_4(OH)_2CO_2H$ [5 2 1]. *Genusic acid Hydroquinone carboxylic acid*. [197°]

Formation — 1 By fusing iodo salicylic acid [196°] or biomo salicylic acid with KOH (Laute-mann, *A* 120, 299; Bakowsky a Leppert, *B* 8, 789, Müller, *A* 220, 124, P F Frankland, *C J* 37, 750) — 2 From oxy amido-benzoic acid (Goldberg, *J pr* [2] 19, 371) — 3 By fusing genusin with potash (Hlawetz a Habermann, *A* 175, 66, Tiemann a Miller, *B* 14, 1988) — 4 By digesting $KHCO_3$ (4 pts) with hydroquinone (1 pt), and water (4 pts) (Senhofer a Sarlay, *M* 2, 448)

Properties—Needles or prisms, v sol water, alcohol, and ether. FeCl_3 colours its solution blue. Reduces Fehling's solution on heating. Split up on distillation into CO_2 and hydroquinone. Benzamidine forms a compound [266°] (Pinner, *B* 28, 2939).

Salts— NaA' , 5½ aq. Deliquescent prisms. KA' aq— CaA' , 7 aq— BaA' , 8. S 40 at 18°— PbA , 2 aq— CuA , 2½ aq.

Ethyl ether EtA' [75°] Crystals

m-Methyl derivative

$\text{C}_6\text{H}_4(\text{OH})(\text{OMe})\text{CO}_2\text{H}$ [251] [142°] S 17 at 10°, 9 at 100°. Formed by oxidising the acetyl-methyl derivative of gentisic aldehyde $\text{C}_6\text{H}_3(\text{OAc})(\text{OMe})\text{CHO}$ with KMnO_4 , and saponifying the product (Tiemann & Miller, *B* 14, 1997). Formed also by the action of CO_2 at 225° on $\text{C}_6\text{H}_4(\text{ONa})(\text{OMe})$ (Korner & Bertoni). Needles. Its solution is coloured blue by FeCl_3 .

Di-methyl derivative $\text{C}_6\text{H}_3(\text{OMe})_2\text{CO}_2\text{H}$ [76°] Formed by oxidising $\text{C}_6\text{H}_3(\text{OMe})_2\text{CHO}$. Needles— AgA' small white needles.

Tri-oxy-benzoic acid v GALLIC ACID

Tri-oxy benzoic acid $\text{C}_6\text{H}_2(\text{OH})_3\text{CO}_2\text{H}$

Tri-methyl derivative $\text{C}_6\text{H}_2(\text{OMe})_3\text{CO}_2\text{H}$ [109°] Formed by oxidising the tri methyl derivative of æsculetic acid (Will, *B* 16, 2113).

Tri-ethyl derivative $\text{C}_6\text{H}_2(\text{OEt})_3\text{CO}_2\text{H}$ [134°] Formed by oxidation of the tri ethyl derivative of (α) or (β) æsculetic acid with KMnO_4 (W). Slender needles.

Tri oxy-benzoic acid

$\text{C}_6\text{H}_2(\text{OH})_3\text{CO}_2\text{H}$ [4321] Mol w 170 [206°–220°] S 13 at 12.5° HF 231,300 HCp 633,700 (Stohmann). Formed, together with pyrogallol dicarboxylic acid, by heating pyrogallol with ammonium carbonate (Senhofer & Brunner, *M* 1, 474, Kostanecki, *B* 18, 3202, Schiff, *A* 245, 85). Needles (containing ½ aq), sol alcohol. FeCl_3 colours its dilute solution violet. Bleaching powder and nitric acid do the same. Lime and baryta water gives a bluish pp. Reduces ammoniacal AgNO_3 in the cold. H_2SO_4 does not form rufigallic acid (difference from gallic acid). POCl_3 forms an acid $\text{C}_6\text{H}_2\text{O}_3$, greatly resembling tannin. It is an astringent yellow powder and gives $\text{Ba}(\text{C}_6\text{H}_2\text{O}_3)_2$ and $\text{C}_6\text{H}_2\text{Ac}_2\text{O}_3$.

Salts— KA' aq— NaA' 2 aq— BaA' , 5 aq— CaA' , 4 aq— Pb , $\text{C}_6\text{H}_2\text{O}_3$, 1½ aq white flocculent pp. **Methyl ether** $\text{C}_6\text{H}_2(\text{OH})_2\text{CO}_2\text{Me}$ [152°] Needles (containing 2½ aq).

Tri-methyl derivative $\text{C}_6\text{H}_2(\text{OMe})_3\text{CO}_2\text{H}$ [99°] Crystals (Will, *B* 21, 2020).

Methyl ether of the tri methyl derivative $\text{C}_6\text{H}_2(\text{OMe})_3\text{CO}_2\text{Me}$ (281°) Oil.

Ethyl ether EtA' [102°] Colourless crystals (containing aq) melting at 86° when hydrated. FeCl_3 gives a greenish-brown colour (Will & Albrecht, *B* 17, 2100, Schiff, *A* 245, 40).

Tri-ethyl derivative $\text{C}_6\text{H}_2(\text{OEt})_3\text{CO}_2\text{H}$ [100°] Formed by oxidation of the tri ethyl derivative of daphnetic acid with KMnO_4 (Will, *B* 17, 1088, 2099). Silky needles, sl sol cold water— BaA' — AgA' . Gives $\text{C}_6\text{H}_2(\text{OEt})_3$ when heated.

Ethyl ether of the tri-ethyl derivative $\text{C}_6\text{H}_2(\text{OEt})_3\text{CO}_2\text{Et}$. Oil. Formed by ethylating the acid.

Tri oxy benzoic acid $\text{C}_6\text{H}_2(\text{OH})_3\text{CO}_2\text{H}$.

Phloroglucin carboxylic acid. Formed by boiling phloroglucin (1 pt) with KHCO_3 (4 pts) and water (4 pts) (Will & Albrecht, *B* 17, 2103, 18, 1323). Colourless crystals (containing aq), sol alcohol and ether. Split up by boiling water into CO_2 and phloroglucin. FeCl_3 gives a transient blue colour. Alcohol and HCl yields CO_2 and the diethyl ether of phloroglucin. On heating with POCl_3 it gives an isomeride of tannin, which is, however, not very astringent and gives no colour with FeCl_3 (Schiff).

Tri methyl derivative?

$\text{C}_6\text{H}_2(\text{OMe})_3\text{CO}_2\text{H}$ **Asaronic acid** [144°] (300°) Formed by oxidising asarone on heating aqueous KMnO_4 (Butlerow & Rizza, *J R* 19, 3). Needles. On distillation with lime it yields $\text{C}_6\text{H}_2(\text{OMe})_3$ (246°).

References—BROMO, BROMO-NITRO, CHLORO, CHLORO IODO, CHLORO NITRO, IODO, and IODO NITRO OXY BENZOIC ACIDS.

O OXY-BENZOIC ALDEHYDE $\text{C}_6\text{H}_4\text{O}_2$ v

$\text{C}_6\text{H}_4(\text{OH})\text{CHO}$ **Salicylic aldehyde** Mol w 122 [c –20°] (1965°) SG 2 11671 (Bruhl) μ_D 1.5960 R_D 54.53 Occurs in the blossom of the meadow sweet (*Spiraea ulmaria*) (Etting, *A* 35, 247) and in *Crepis fetida* (Wicke, *A* 91, 374).

Formation—1 By oxidation of saligenin or salicin (Piria, *A* 30, 153)—2 In the products of the dry distillation of guinic acid (Wohler, *A* 51, 146)—3 By mixing chloroform (15 pts), phenol (10 pts), NaOH (20 pts), and water (35 pts), at 50°, boiling with inverted condenser, distilling off excess of chloroform, acidifying and distilling with steam (Tiemann & Reimer, *B* 9, 423, 824).

Properties—Liquid, with pleasant odour, v sl sol water, miscible with alcohol and ether. Turns red in air. Added to 'Vermouth' and other liqueurs. Produces epileptic convulsions in dogs (Laborde & Magnan, *J Ph* [5] 16, 448). FeCl_3 colours its aqueous solution violet. Does not reduce Fehling's solution. Combines with KHSO_4 forming $\text{C}_6\text{H}_4\text{O}(\text{OH})\text{SO}_3\text{K}$ crystallising in needles (Bertagnini, *A* 85, 193).

Reactions—1 Yields o oxy benzoic acid on oxidation—2 Sodium amalgam reduces it to saligenin (Beilstein & Reinecke, *A* 128, 179)—3 Heated with ZnCl_2 and HOAc it forms red amorphous $\text{C}_6\text{H}_4\text{O}_3$, insol water, which dissolves in alkalis with violet red colour but is insol acids. It yields amorphous $\text{C}_6\text{H}_4\text{AcO}_3$ (Bourquin, *B* 17, 502)—4 Ac_2O (2 mols) at 180° forms $\text{C}_6\text{H}_2(\text{OAc})_2\text{CH}(\text{OAc})$, [100°] which splits up on distillation into Ac_2O and $\text{C}_6\text{H}_2(\text{OAc})_3\text{CHO}$ (255°). On treatment with soda the compound $\text{C}_6\text{H}_2(\text{OAc})_2\text{CH}(\text{OAc})_2$ yields $\text{C}_6\text{H}_2(\text{OH})_2\text{CH}(\text{OAc})_2$, [104°] (Barbier, *C R* 90, 87)—5 **Acetyl chloride** yields 'disalicyl aldehyde' $\text{C}_6\text{H}_4\text{O}_3$, [180°]. The same body is formed by the action of BrCl , succinyl chloride, and PCl_5 (Osahours, *A* 78, 228, Perkin, *A* 145, 299, Zwenger, *A Suppl* 8, 42). It crystallises from alcohol in long needles, and is converted by Br in HOAc to $\text{C}_6\text{H}_2\text{Br}_2(\text{OH})\text{CHO}$, [166°], together with $\text{C}_6\text{H}_2\text{Br}(\text{OH})\text{CHO}$ [105°] and $\text{C}_6\text{H}_2\text{Br}_2(\text{OH})\text{CHO}$ [88°] (Bradley, *B* 22, 1134). 6 **Zinc dust** and HOAc form $\text{C}_6\text{H}_2\text{O}_3$, crystallising in needles [82°] (Tiemann, *B* 19, 357)—7 **Bromine** forms $\text{C}_6\text{H}_2\text{Br}_2(\text{OH})\text{CHO}$ and $\text{C}_6\text{H}_2\text{Br}_2\text{O}$ (Werner, *Bi* [2] 46, 277)—8. **Cyanogen**

bromide forms crystalline $C_6H_5NO_2$ (Cahours, *A* 108, 322)—9 *Ammonium cyanide* and alcohol form $C_6H_5N_3O_2$ [143°] and $C_6H_5N_3O_2$ [168°] (Haarmann, *B* 6, 841)—10 *Sodium succinate* and acetic anhydride form, on heating, dicoumarin $C_6H_5 \cdot O \cdot CO \cdot CO \cdot O$ $\begin{array}{c} \diagup \\ \text{CH} \end{array} \begin{array}{c} \diagdown \\ \text{CH} \end{array} \begin{array}{c} \diagup \\ \text{CH} \end{array} \begin{array}{c} \diagdown \\ \text{CH} \end{array} \cdot C_6H_5$ (Fittig, *B* 18, 2523)—11 *Thioglycolic acid* and zinc chloride form $C_6H_5(OH)CH(SCH_2CO_2H)_2$ [148°] (Bongartz, *B* 21, 480)—12 Gaseous or alcoholic ammonia gives rise to 'hydrosalicylamide' $C_6H_5(OH)CHNCH(C_6H_5OH)NCHC_6H_5OH$ which forms yellow crystals [145°] (Etting, *A* 35, 249, Herzfeld, *B* 10, 1270) It is insoluble in water, soluble in cold alcohol, and is decomposed by boiling acids and alkalis into NH_3 and *o*-oxybenzoic aldehyde. It is converted by alcoholic ammonium sulphide into crystalline $C_{12}H_{11}N_2SO_2$, and by HCl and HCl into two isomeric crystalline 'hydrocyanals' $C_6H_5N_3O_2$ (Beilstein, *A* 136, 170) Hydrosalicylamide forms the salts $FeC_{12}H_{11}N_2O_2NH_4$ and $Cu_2(C_{12}H_{11}N_2O_2)_2 \cdot 2NH_3$ —13 *Methylamine* gas forms $C_6H_5(OH)CHNMe$ an oil [229°], resolved by acids and alkalis into NH_3 and salicylic aldehyde (Dennstedt & Zimmermann, *B* 21, 1553)—14 *Ethylamine* forms the homologous C_6H_5NO (237°)—15 *Aniline* forms $C_6H_5(OH)CHNPh$ [50.5°] (Schischkoff, *C* 45, 272, Emmerich, *A* 241, 344) It forms a crystalline compound with HCl *p*-Nitroaniline forms the compound $C_6H_5(NO_2)NCHC_6H_5OH$ [115°]—16 *Di-methyl-p-phenylene diamine* forms in like manner $C_6H_5(OH)CHN(C_6H_5NMe_2)$ [134°] (Nuth, *B* 18, 573) Di-methylaniline and $ZnCl_2$ form $C_6H_5(OH)CH(C_6H_5NMe_2)_2$ —17 *Benzidine* in weak alcoholic solution forms the compound $C_{12}H_{11}NCHC_6H_5OH$, crystallising from benzene in colourless needles [260°] Di-amido ditolyl forms the homologous $C_{12}H_{12}NCHC_6H_5OH$ [202°] (Schiff & Vanni, *A* 258, 374)—18 *Ethylene diamine* forms $C_6H_5NCHC_6H_5OH$ [126°] (Mason, *B* 20, 271)—19 *Ethylene aniline* forms $C_6H_5(NPh)CHC_6H_5OH$ [116°] (Moos, *B* 20, 733)—20 *p*-Toluidine gives rise to $C_6H_5MeNCHC_6H_5OH$ [100°] (Jaillard, *Z* 1865, 440)—21 *Phenylene m-diamine hydrochloride* yields $C_6H_5(NCHC_6H_5OH)_2$, forming crystalline $B_2H_2PtCl_4$ (Schiff, *A* 253, 329) Tolyene *m*-diamine yields homologous $C_6H_5NCHC_6H_5OH$ [109°]—22 *Tolyene o-diamine* forms a compound $C_{12}H_{12}N_2O_2$ [106°–110°] and azurine $C_{12}H_{12}N_2O_2$ [250.5°] which exhibits blue fluorescence in alkaline solution (Ladenburg, *B* 11, 596)—23 (8) *Naphthylamine* reacts forming $C_{10}H_7NCHC_6H_5OH$ [121°] (Emmerich, *A* 241, 351)—24 *Urea* in aqueous solution gives crystalline $(NH_2CO)_2CHC_6H_5OH$ whence $Cu(C_6H_5N_3O_2)_2$ (Schiff, *A* 151, 199)—25 *m*-Amido-benzoic acid forms the compound $CO_2H \cdot C_6H_4NCHC_6H_5OH$ [190°] which forms an amide [186°] (Schiff, *A* 210, 114)

Salts— $KC_6H_5O_2$, aq yellow tables *S* (alcohol) 5 (Michael, *Am* 1, 309)— $NaHA' \cdot \frac{1}{2}aq$ — $BaA' \cdot 2aq$ — $Pb(OH)A'$ — CuA' , brownish green crystals
Acetyl derivative $C_6H_5(OAc)CHO$ [37°] (253°) Formed from $C_6H_5(ONa)CHO$ in ether by adding Ac_2O (Perkin, *A* 148, 208, 150, 82)
Butyryl derivative $C_{12}H_{11}O_2$ (260°–270°). Oil

Benzoyl derivative $C_6H_5(OBz)CHO$ Oil
Glucoside *v* HELICIN

Methyl derivative $C_6H_5(OMe)CHO$ [35°] (238°). Formed from $C_6H_5(ONa)CHO$, MeI, and MeOH (Perkin, *A* 145, 302, *C* 55, 550, Voswinckel, *B* 15, 2024) Thick prisms, nearly insoluble in water, soluble in alcohol, very soluble in ether. Alcoholic HCl and H_2S form the (8) thioaldehyde $C_{12}H_{12}S_2O_2$ [224°] While at -10° the (a)-isomeride $C_{12}H_{12}S_2O_2$ [157°] is formed (Baumann & Fromm, *B* 24, 1446) Alcohol and colourless ammonium sulphide yield $C_{12}H_{12}S_2O_2$ [85°–88°] When KCy followed by HCl is added to the ethereal solution of $C_6H_5(OMe)CHO$ there is formed $C_6H_5(OMe)CH(OH)CN$ [71°] whence alcoholic NH_3 at 70° yields $(C_6H_5(OMe)CHCy)_2NH$ [123°], and alcoholic aniline at 100° gives $C_6H_5(OMe)CH(NHPh)CN$ [61°] (V) Ethylene diamine at 120° forms crystalline $C_{12}H_{12}(NCHC_6H_5OMe)_2$

Ethyl derivative $C_6H_5(OEt)CHO$ [7°] (249°) (Göttig, *B* 10, 8, Perkin, *A* 145, 306, *C* 55, 551) With alcoholic NH_3 it yields crystalline $N_2(CHC_6H_5OEt)_2$, which is converted by heating at 165° into an amorphous isomeride yielding crystalline $B_2H_2PtCl_4$ (Perkin, *A* 145, 308) Forms with aniline only $C_{12}H_{12}(OEt)CHNPh$ (Schiff, *A* 150, 195), and with ethyl aniline only $C_{12}H_{12}(OEt)CH(NEtPh)_2$ Aqueous urea forms crystalline $C_{12}H_{12}N_2O_2$, aq

Isobutyl derivative $C_6H_5(OC_4H_9)CHO$ (265°) Oil (Baumann & Fromm, *B* 24, 1448) Alcoholic HCl and H_2S form the (a) [142°] and (8) [163°] isomerides $C_{12}H_{12}S_2O_2$. Ammonium sulphide yields $C_{12}H_{12}S_2O_2$ [52°–56°]

Benzyl derivative $C_6H_5(OC_6H_5)CHO$ [46°]

Derivatives of *o*-oxybenzoic orthoaldehyde

$C_6H_5(OH)CH(OAc)_2$ [104°] Formed from the aldehyde and Ac_2O at 150° (Perkin, *A* 146, 371°) Tables (from alcohol) $C_6H_5(OAc)CH(OAc)_2$ [101°] Needles (from alcohol)

$C_6H_5(OMe)CH(OAc)_2$ [75°] Prisms $C_6H_5(OEt)CH(OAc)_2$ [89°] Prisms, insoluble in Aq

Oxim $C_6H_5(OH)CHNOH$ [57°] White crystals, resolved by warm HCl aq into its components (Lach, *B* 16, 1782, 17, 1572) Ac_2O converts it into acetyl-*o*-oxybenzonitrile. The compound $NHPhCO \cdot O \cdot C_6H_5$, $CHNO \cdot CO \cdot NHPh$ [115°] is formed by phenyl cyanate (Goldschmidt & Schulthess, *B* 22, 3102)— $B'HCl$ — $C_6H_5(ONa)CHNO \cdot Na_3aq$ small pearly scales

Derivatives of the oxim

$C_6H_5(OMe)CHNOH$ [92°] With phenyl cyanate it yields $C_6H_5(OMe)CHNO \cdot CO \cdot NHPh$ [105°] (Goldschmidt, *B* 23, 2741)

$C_6H_5(OMe)CHNOEt$ Oil

$C_6H_5(OEt)CHNOEt$ Oil

$C_6H_5(OH)CHNO \cdot C_6H_5$ The (a) isomeride [63°] is formed from *o*-oxybenzoic aldehyde and (a) benzyl-hydroxyamine, while the (8) isomeride [100°] is obtained by using (8) benzyl-hydroxyamine (Beckmann, *B* 23, 3319)

Phenylhydrazide $C_6H_5(OH)CHN \cdot NHPh$ [143°] Colourless needles (from alcohol) (Fischer, *B* 17, 575, Rossing, *B* 17, 3003). Yields $C_6H_5(OAc)CHN \cdot NaOPh$ [138°], which forms a crystalline dibromide converted by boiling alcohol into $C_6H_5Br_2(OAc)CHN \cdot NHPh$ [168°], which

yields $C_6H_5Br(OAc)CHN_2AcPh$ [158°] and $C_6H_5Br(OH)CHN_2HPh$ [148°]

m-Oxy-benzoic aldehyde $C_6H_4(OH)CHO$ [81] [104°] (240°) Formed by reduction of *m* oxy-benzoic acid in acid solution by sodium-amalgam (Sandmann, B 14, 969) Obtained also by oxidation and diazotisation from *m* amido-cinnamic acid (Luff, B 22, 294) Prepared from *m*-nitro-benzoic aldehyde by reduction and treatment of the amido compound with nitrous acid (Tiemann & Ludwig, B 15, 2043) Needles (from water) Excess of Ac_2O forms $C_6H_4(OAc)CH(OAc)_2$ crystallising in plates [76°]

Acetyl derivative $C_6H_4(OAc)CHO$ (263°)

Formed from the K salt and Ac_2O Oil

Methyl derivative $C_6H_4(OMe)CHO$ (230°)

Oxim $C_6H_4(OH)CHNOH$ [87.5°] Soft silky needles (Clemm, B 24, 826)

Phenyl hydrazide

$C_6H_4(OH)CHN_2HPh$ [181°] Prisms, ν sol alcohol (Rudolph, A 248, 102)

p-Oxy-benzoic aldehyde $C_6H_4(OH)CHO$ [4.1] [115°] Formed by heating its methyl derivative with $HClAq$ at 200° (Bucking, B 9, 527) Prepared, together with the *o*-isomeride, by the action of chloroform and alkalis on phenol (Tiemann & Reimer, B 9, 824, 10, 63)

Properties — Needles (from water), ν sol alcohol and ether Not volatile with steam $FeCl_3$ gives a slight violet tint to its aqueous solution Reduces ammoniacal $AgNO_3$ With $NaHSO_3$ it forms the crystalline compound $C_6H_4(OH)CH(OH)SO_3Na$ [112°]

Reactions — 1 Potash fusion forms *p* oxy-benzoic acid — 2 Sodium amalgam and water reduce it to $C_6H_4(OH)CH(OH)CH(OH)C_6H_4(OH)$ [222°] and the isomeric di-oxy isohydrobenzoin [198°] which forms the crystalline derivative $C_6H_4(OH)_2(C_6H_4OAc)_2$ [192°] (Herzfeld, B 10, 1268, Tiemann, B 19, 354) — 3 Bromine ppts $C_6H_5Br(OH)CHO$ [181°] and, when in excess, forms $C_6H_5Br(OH)CHNOH$ (Werner, Bl [2] 46, 278) — 4 Boiling Ac_2O (3 pts) forms $C_6H_4(OAc)CH(OAc)_2$ [94°] (Tiemann & Herzfeld, B 10, 64, Barbier, C R 90, 37) — 5 Heating with $HOAc$ and $ZnCl_2$ forms red amorphous $C_{10}H_8O_3$, which gives a violet solution in alkalis (Bourquin, B 17, 503) — 6 Ammonia forms an oily compound — 7 Aniline in ethereal solution forms $C_6H_4(OH)CHNPh$ [191°] — 8 *p* Toluidine yields the compound $C_6H_4(OH)CHNC_6H_5$ [218°] — 9 *Di-methyl-phenylene-diamine* gives rise to crystalline $C_6H_4(OH)CHNC_6H_4NMe_2$ decomposing at 240° (Nuth, B 18, 574) — 10 (β)-Naphthylamine forms $C_6H_4(OH)CHNC_{10}H_7$ [220°] (Emmerich, A 241, 356)

Acetyl derivative $C_6H_4(OAc)CHO$ (260°) (Barbier, Bl [2] 33, 52, C R 90, 37, (265°) (T & H) Formed from $C_6H_4(OK)CHO$ and Ac_2O Oil

Methyl derivative $C_6H_4(OMe)CHO$ **Anisic aldehyde** Mol ν 136 (248°) SG 1.228 Formed, together with anisic acid, by oxidation of anethol or oil of anise (Cahours, A Ch [3] 14, 484, 23, 354, Rossel, A 151, 25) Formed also by distilling calcium anisate with calcium formate (Piria, A 100, 105) and by methylation of *p*-oxy-benzoic aldehyde (T & H) Oil, forming with H_2SO_4 a crimson solution, turned violet on heating With $NaHSO_3$ it forms crystalline $C_6H_4NaSO_3$ aq (Bertagnini, A. 85,

268) **Reactions** — 1 Alcoholic potash forms the corresponding alcohol and acid — 2 Alcoholic HCl and H_2S form $C_6H_5S_2O_3$ [183°] and, at -10°, an isomeride [127°] (Baumann & Fromm, B 24, 1442) Alcoholic H_2S forms the thio-aldehyde [75°-77°], while alcoholic ammonium sulphide forms a polymeric thioanisic aldehyde [92°] and the disulphide $(C_6H_4(OMe)CH)_2S_2$ (B' & F) — 3 Sodium amalgam forms two 'hydrazones' $C_6H_4(OMe)CH(OH)CH(OH)C_6H_4(OMe)$, melting at 172° and 125° (Samosadsky, Z 1867, 678, 1868, 643) Boiling dilute H_2SO_4 converts the isomeride [172°] into $C_6H_5H_2O_3$ [95°] (Rossel, A 151, 36) Zinc and hydrochloric acid form $C_6H_4(OMe)CH(OH)$ and crystalline $C_6H_4O_3$ [215°] — 4 Aqueous HCy (28 p) forms $C_6H_4(OMe)CH(OH)CN$ [68°], which yields $C_6H_4(OMe)CH(OH)CO_2H$ on saponification (Tiemann & Friedlander, B 14, 1976) — 5 Alcoholic KCy yields anisoin — 6 Succinic acid yields $C_6H_4(OMe)CHCH_2CO_2H$ and $C_6H_4(OMe)CHCH_2C(CO_2H)CHC_6H_4OMe$ (Fit tig, B 18, 2523) — 7 Di thio glycol gives rise to $C_6H_4(OMe)CHS_2C_6H_5$ [65°] (Easbender, B 21, 1476) — 8 Aqueous ammonia produces 'anishydramide' $N(CH_2C_6H_4OMe)_2$ [120°] converted at 170° into crystalline 'anisine' $C_6H_4N_2O_3$, which forms the salts $BHCl$ aq and $B_2H_4PtCl_4$ (Bertagnini, A 88, 128) — 9 Ethylene aniline forms $C_6H_4(NCH_2C_6H_4OMe)_2$ [111°] (Mason, B 20, 272) — 10 Aniline gives rise to crystalline $C_6H_4(OMe)CHNPh$ — 11 Ethylene aniline forms $C_6H_4(OMe)CH(NPh)_2$ C_6H_4 [164°] (Moos, B 20, 733) — 12 *o* Toluidine reacts, forming $C_6H_4(OMe)CHNC_6H_4Me$ [32°] The *p* isomeride [92°] is also crystalline (Steinhart, A 241, 340) — 13 Phenylene-*di*-methyl-*p* diamine forms $C_6H_4(OMe)CHNC_6H_4NMe_2$ [148°] (S, cf Nuth, B 18, 574) — 14 Toluylene-*o* diamine hydrochloride forms $C_{12}H_{12}N_2O_3$ [152°-156°] (Ladenburg, B 11, 1660) — 15 (β) Naphthylamine yields $C_6H_4(OMe)CHNC_{10}H_7$ [98°] — 16 Acetamide at 120°-180° forms $C_6H_4(OMe)CH(NHAc)$ [180°] (Schuster, A 154, 80) — 17 Benzamide gives $C_6H_4(OMe)CH(NHBz)_2$ [192°] — 18 Urea forms crystalline $C_6H_4(OMe)CH(NHCO NH)_2$ and $C_{10}H_{12}N_2O_3$ — 19 Carbamic ether and HCl give $C_6H_4(OMe)CH(NHCO_2Et)$ [172°]

Oxim $C_6H_4(OH)CHNOH$ [65°] Formed from the aldehyde and hydroxylamine (Lach, B 16, 1785) White needles Converted by Ac_2O and by $AcCl$ into $C_6H_4(OH)CN$ Yields $C_6H_4(ONa)CHNONa3aq$

Methyl derivative of the oxim $C_6H_4(OMe)CHNOH$ (a) - Isomeride [62°] Formed from anisic aldehyde and hydroxylamine (Westenberger, B 16, 2993, Goldschmidt & Polonowska, B 20, 2407, 22, 3102, 23, 2163; Beckmann, B 21, 768, 23, 1687, Miller, B 22, 2790) White plates, ν sol hot water Tastes sweet Heated with Ac_2O and HCl it gives $C_6H_4(OMe)CN$ [81°] $NaOEt$ and benzyl chloride yield the (a) benzyl ether [46.5°] Ac_2O forms $C_6H_4(OMe)CHNOAc$ [48°] (Hantzsch, B 24, 41), crystallising in prisms Phenyl cyanate forms $C_6H_4(OMe)CHNOCONHPh$ [82°] $NaOMe$ and MeI form $C_6H_4(OMe)CHNOMe$ [43°] (246°) — (b) - Isomeride [130°] Ppd as hydrochloride by passing HCl into an ethereal solution of the (a) isomeride Slender needles Has no taste. With $NaOEt$ and benzyl chloride it yields the

(8)-benzyl ether [107°] The acetyl derivative $C_6H_5(OMe)CHNOAc$ [64°] is converted by Na_2CO_3 aq into the nitrate [60°]

Phenyl-hydrazide $C_6H_5(OH)CHN.HPh$ [178°] Tufts of needles (Rudolph, A 248, 102)

Phenyl-hydrazide of the methyl derivative $C_6H_5(OMe)CHN.HPh$ [121°]

c Di-oxy benzoic aldehyde m Methyl derivative $C_6H_5(OMe)(OH)CHO$ [3 3 1] (264°-268°) Formed, together with vanillin, by the action of chloroform on a solution of guaiacol in dilute NaOH (Tiemann & Koppe, B 14, 2020) Liquid, volatile with steam, sol alcohol, ether, and benzene, nearly insol water $FeCl_3$ colours its alcoholic solution violet

Di-oxy benzoic aldehyde $C_6H_5(OH)_2CHO$ [4 2 1] (8) *Resorcylic aldehyde* [135°] Formed by the action of chloroform and NaOH aq on resorcin (Tiemann & Lewy, B 10, 2212) Needles (from water), v sol water, alcohol, and ether $FeCl_3$ colours the aqueous solution reddish-brown Readily resinsified

o Methyl derivative $C_6H_5(OH)(OMe)CHO$ [4 2 1] [153°] Formed, together with the *p* methyl derivative, by the action of chloroform and NaOH on $C_6H_5(OH)(OMe)$ [1 3] (Tiemann & Parrissius, B 13, 2365) Colourless plates, sl sol. water Gives white crystalline pps with ammoniacal $AgNO_3$, and with $Pb(OAc)_2$ Yields an acetyl derivative $C_6H_5(OAc)(OMe)CHO$ [86°]

p Methyl derivative $C_6H_5(OMe)(OH)CHO$ [4 2 1] [63°] Formed by partial methylation of the aldehyde White plates, nearly insol water $FeCl_3$ colours its alcoholic solution reddish violet Gives pps with ammoniacal $AgNO_3$, and lead acetate

Di-methyl derivative $C_6H_5(OMe)_2CHO$ [68°] Obtained by methylation, and also by oxidation of the di-methyl derivatives of (a) and (8) umbellic acid with $KMnO_4$ (Will, B 16, 2117) Needles (from dilute alcohol)

Di-ethyl derivative [72°]

Phenyl hydrazide $C_6H_5(OH)CHN.HPh$ [c 158°] Needles (Rudolph, A 248, 104)

Di-oxy benzoic aldehyde

$C_6H_5(OH)_2CHO$ [5 2 1] *Genticic aldehyde* [99°] Formed by boiling hydroquinone with chloroform and aqueous (18 p c) NaOH (Tiemann & Muller, B 14, 1986) Flat yellow needles, v sol water Gives a transient blue colour with $FeCl_3$ Yields gentisic acid on fusion with potash Alcoholic aniline forms the amide $C_6H_5(OH)CHNPh$, crystallising in red needles

m Methyl derivative

$C_6H_5(OMe)(OH)CHO$ [5 2 1] [4°] (248°) V D (H=1) 75.7 (obs) Formed from methyl hydroquinone $C_6H_5(OH)(OMe)$ [1 4], chloroform, and NaOH aq Liquid, volatile with steam, sl sol water Gives a bluish green colour with $FeCl_3$ Aniline yields $C_6H_5(OMe)(OH)CHNPh$ [59°], crystallising in red needles The acetyl derivative $C_6H_5(OMe)(OAc)CHO$ [68°] crystallises in needles, and is converted by boiling Ac_2O into $C_6H_5(OMe)(OAc)CH(OAc)$

Di-methyl derivative $C_6H_5(OMe)_2CHO$ [51°] Volatile with steam Not coloured by $FeCl_3$

m-Ethyl derivative

$C_6H_5(OEt)(OH)CHO$ [5 2 1] [52°] (230°) Yellow prisms, nearly insol water Coloured violet by $FeCl_3$ Yields $C_6H_5(OEt)(OAc)CHO$ [69°], (c 285°)

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Di-ethyl derivative $C_6H_5(OEt)_2CHO$ [80°] (c 283°) Needles (T & M, Hantzsch, J pr [2] 22, 468)

Di-oxy-benzoic aldehyde

$C_6H_5(OH)_2CHO$ [4 3 1] v *PROTocatechuic aldehyde*

Tri-oxy benzoic aldehyde Tri-ethyl derivative $C_6H_5(OEt)_3CHO$ [4 3 2 1] [70°] Formed by oxidising the tri-ethyl derivative of daphnetic acid with $KMnO_4$ (Will & Jung, B 17, 1088)

Tri-oxy benzoic aldehyde Tri-ethyl derivative $C_6H_5(OEt)_3CHO$ [95°] Formed by oxidation of the tri-ethyl derivative of (a) or (8) ascouletic acid with alkaline $KMnO_4$ (Will, B 16, 2112) Large crystals, insol water

Tri-oxy benzoic aldehyde Tri-methyl derivative $C_6H_5(OMe)_3CHO$ [114°] Formed by oxidation of asarone (Butlerow & Rizza, J R 19, 3) Needles, v sol hot water

References—Bromo-, Chloro-, and Iodo OXY BENZOIC ALDEHYDE

m-OXY-BENZOPHENONE $C_6H_5CO C_6H_5(OH)$ [116°] Formed by the action of nitrous acid upon *m* amido-benzophenone (Geigy & Koenigs, B 13, 2402) Needles

p Oxy-benzophenone $C_6H_5CO C_6H_5OH$ [1 4] *p-Benzoyl phenol* Formed by heating phenol with $BzCl$ and $ZnCl_2$ (Grucarevitch & Merz, B 6, 1245) Obtained also from *p* amido benzo phenone by the diazo reaction (Doebner & Weiss, B 14, 1840, A 210, 275) Needles or plates

Acetyl derivative [81°] Needles (from alcohol) (Doebner & Stackmann, B 10, 1970)

Benzoyl derivative [113°]

Methyl derivative $C_6H_5CO C_6H_5OMe$ [62°] Four sided prisms (Rennie, C J 41, 227) Possesses two oxims $C_6H_5C(NO) C_6H_5OMe$, a stable oxim [116°] yielding $BHCl$, an acetyl derivative [53°], and a benzyl ether [74°], and an unstable oxim [140°], which yields $BHCl$ [124°], an acetyl derivative [135°], and a benzyl ether [60 5°] (Schafer, A 264, 158, Hantzsch, B 24, 53)

Ethyl derivative $C_6H_5CO C_6H_5OEt$ [39°] (above 300°) Formed from C_6H_5OEt , benzoyl chloride, and $AlCl_3$ (Gattermann, Ehrhardt, & Marsch, B 23, 1206)

Di-oxy benzophenone $CO(C_6H_5OH)_2$ [60°] (c 335°) Formed by heating diphenylene ketone oxide with alcoholic potash at 180° (Richter, J pr [2] 28, 273, Graebe & Feer, B 19, 2607) Prisms or plates (from ligroin) —KHA" yellow crystals (from alcohol) —K₂A" crystals, v e sol water

Acetyl derivative $C_6H_5AcO_2$ [96°] (G & F), [85°] (R) Prisms (from alcohol)

Benzoyl derivative $C_6H_5BzO_2$ [104°]

Methyl derivative $C_6H_5MeO_2$ [69°]

Di-methyl derivative $C_6H_5Me_2O_2$ [98°] (R), [104°] (G & F) Prisms Forms the oxim $C(NO)H(C_6H_5OMe)_2$ [188°]

Di-ethyl ether $C_6H_5Et_2O_2$ [109°] Needles (from dilute alcohol) Yields the phenyl hydrazide $C(N.HPh)(C_6H_5OEt)_2$ [114°]

Oxim $C(NO)H(C_6H_5OH)_2$ [99°]

Phenyl-hydrazide $C(N.HPh)(C_6H_5OH)_2$ [152°]

op-Di-oxy-benzophenone

$[2 1] C_6H_5(OH)CO C_6H_5(OH)$ [1 4] [144°] Formed by heating salicylic acid with phenol

Y Y

and SnCl_4 at 120° (Michael, *B*, 14, 656, *Am* 5, 83) Large yellow plates, sl sol water — $\text{Ag}_2\text{A}''$ aq

Acetyl derivative $\text{C}_{15}\text{H}_{11}\text{Ac}_2\text{O}_2$ [88°]

Di-p-oxy-benzophenone $\text{CO}(\text{C}_6\text{H}_4\text{OH})_2$ [210°]

Formation — 1 From $\text{CH}_3(\text{C}_6\text{H}_4\text{OBz})_2$ by oxidation and saponification (Gail, *A* 194, 834)

2 By fusing phenol-phthalein with potash (Baeyer a Burkhardt, *B* 11, 1299, *A* 202, 126)

3 By heating aurin with water at 240° (Caro a. Graebe, *B* 11, 1348), or rosaniline with water at 270° (Liebermann, *B* 6, 951, 11, 1435) —

4 By the action of HNO_3 on di-*p*-amido-benzophenone (Staedel a Sauer, *B*, 11, 1747)

Properties — Long needles, m sol hot water On treatment with PCl_5 followed by phenol and H_2SO_4 it yields aurin Bromine forms $\text{C}_{15}\text{H}_7\text{Br}_2\text{O}_2$ [214°]

Acetyl derivative $\text{C}_{15}\text{H}_{11}\text{Ac}_2\text{O}_2$ [148°]

Benzoyl derivative $\text{C}_{15}\text{H}_7\text{Bz}_2\text{O}_2$ [182°]

Methyl derivative $\text{C}_{15}\text{H}_{11}\text{Me}_2\text{O}_2$ [144°]

Needles (Bösler, *B* 14, 328) Yields $\text{C}_{15}\text{H}_{11}\text{Br}_2\text{O}_2$ [181°] and an oxim [133°]

Ethyl derivative $\text{C}_{15}\text{H}_{13}\text{Et}_2\text{O}_2$ [147°]

Di-ethyl derivative $\text{C}_{15}\text{H}_{13}\text{Et}_2\text{O}_2$ [147°] (Gail, [181°] (Gattermann, *B* 22, 1181)

(8) *Di-oxy-benzophenone* $\text{C}_{15}\text{H}_{10}\text{O}_2$ [162°]

Formed from di-nitro benzophenone [149°] by reduction and application of the diazo reaction (Staedel a Sauer, *B* 18, 836) Needles

Acetyl derivative $\text{C}_{15}\text{H}_9\text{Ac}_2\text{O}_2$ [90°]

Benzoyl derivative $\text{C}_{15}\text{H}_5\text{Bz}_2\text{O}_2$ [102°]

Di-oxy-benzophenone $\text{C}_6\text{H}_4\text{CO C}_6\text{H}_4(\text{OH})_2$ [145°] Formed from di benzoyl pyrocatechin, BzCl , and ZnCl_2 (Doebner, *A* 210, 261) Needles (containing 1/2 aq)

Benzoyl derivative $\text{C}_{15}\text{H}_5\text{Bz}_2\text{O}_2$ [95°]

Di oxy benzophenone $\text{C}_6\text{H}_4\text{CO C}_6\text{H}_4(\text{OH})_2$

Benzoresorcin [144°] Formed from resorcin, BzCl , and ZnCl_2 at 120° (Doebner a Stachmann, *B* 11, 2270) Needles, sol hot water

Benzoyl derivative $\text{C}_{15}\text{H}_5\text{Bz}_2\text{O}_2$ [141°]

Tri oxy benzophenone

$\text{C}_6\text{H}_4(\text{OH})\text{CO C}_6\text{H}_4(\text{OH})_2$ [124] [133°] Formed by heating salicylic acid with resorcin at 200° (Michael, *B* 14, 658), or by heating oxy diphenylene ketone oxide with NaOH at 270° (Graebe, *A* 254, 291) Plates, sl sol water

Tetra-oxy-benzophenone

$\text{CO}(\text{C}_6\text{H}_4(\text{OH})_2)_2$ [125], [202°] is EUXANTHONIC ACID (*q* v)

Hexa-oxy-benzophenone $\text{CO}(\text{C}_6\text{H}_4(\text{OH})_3)_2$

Anhydride $\text{C}_{15}\text{H}_3\text{O}_6$ *Anhydropyrogallolketone* Formed by fusing gallein with alkalis (Buchka, *A* 209, 270) Brown powder

Acetyl derivative $\text{C}_{15}\text{H}_3\text{Ac}_3\text{O}_6$ [237°]

p OXY-BENZOPHENONE o-CARBOXY-

LIC ACID *Methyl derivative* $\text{C}_{15}\text{H}_9\text{O}_4$ *o* $\text{C}_6\text{H}_4(\text{OMe})\text{CO C}_6\text{H}_4\text{CO}_2\text{H}$ *Anisole phthalic acid* [148°] Formed by the action of phthalic anhydride (50 g) on anisole (150 g) in presence of AlCl_3 (80 g) (Nourissou, *B* 19, 2103) Colourless crystals (from toluene) Split up by potash-fusion into benzoic and *p*-oxy benzoic acids Yields a bromo-derivative [198°] Conc H_2SO_4 forms *m* oxy-anthraquinone Distillation with zinc dust gives anthracene — NaA'' — KHA'' — CaA'' , 2aq — BaA'' , 4aq white needles — AgA''

Di oxy benzophenone carboxylic acid

$\text{C}_6\text{H}_4(\text{OH})_2\text{CO C}_6\text{H}_4\text{CO}_2\text{H}$ [200°] Got by fusing

fluorescein with NaOHAq (Baeyer, *A* 183, 23) Crystals (containing aq)

DI-OXY-BENZOPHENONE SULPHONIC

ACID $\text{C}_6\text{H}_4(\text{OH})_2\text{CO C}_6\text{H}_4\text{SO}_3\text{H}$ Formed by heating $\text{C}_6\text{H}_4(\text{SO}_2\text{NH}_2)\text{CO}_2\text{H}$ with resorcin (Remsen, *Am* 9, 5, 872, 11, 73) Small plates (containing 2aq) — BaA'' — $\text{NH}_4\text{A}''$

OXY-BENZOYL ACETIC ACID $\text{C}_6\text{H}_4\text{O}_4$ *o*

$\text{C}_6\text{H}_4\text{CO OH}(\text{OH})\text{CO}_2\text{H}$ [125°] Formed from nitroso benzoyl-acetic ether $\text{Bz C}(\text{NOH})\text{CO}_2\text{Et}$ and NaOHAq (Baeyer a Perkin, *B* 16, 2183, *C* 7, 245) Small prisms (from water) — AgA''

OXY-BENZOYL BROMIDE *Methyl deri-*

vative $\text{C}_6\text{H}_4(\text{OMe})\text{COBr}$ Formed from anisic aldehyde by cautious treatment with bromine (Cahours, *A* *Ch* [8] 14, 486) Silky crystals, resolved by KOHaq into potassium anisate and potassium bromide

p-OXY-BENZOYL CHLORIDE *Methyl*

derivative $\text{C}_6\text{H}_4(\text{OMe})\text{COCl}$ (262°) *SG* 1 261 Formed from anisic acid and PCl_5 (Cahours, *A* *Ch* [3] 23, 351) Oil, converted by water into anisic acid

p OXY BENZURIC ACID $\text{C}_6\text{H}_4\text{NO}_2$ *o*

$\text{C}_6\text{H}_4(\text{OH})\text{CO NH CH}_2\text{CO}_2\text{H}$ [c 228°] Occurs in the urine of dogs to which *p* oxy benzoic acid or hydro *p* coumaric acid has been administered (Baumann a Herter, *H* 1, 260, Schotten, *H* 7, 26) Prisms, m sol water

OXY DIBENZYL v OXY DI PHENYL ETHANE

o OXY-BENZYL ALCOHOL $\text{C}_6\text{H}_4\text{O}_2$ *o*

$\text{C}_6\text{H}_4(\text{OH})\text{CH}_2\text{OH}$ *Salgenin* Mol w 124 [82°] *S* 7 at 22° *S* (benzene) 19 at 18°

Formed by the hydrolysis of salicin (Prins, *A* 56, 37), by reducing *o* oxy benzoic aldehyde with sodium amalgam (Beilstein, *A* 128, 179), and by heating phenol with CH_2Cl_2 and aqueous NaOH at 100° (Greene, *Am* 2, 19) Tables, v e sol hot water FeCl_3 gives a blue colour Dilute H_2SO_4 forms saliretin $\text{C}_{15}\text{H}_{14}\text{O}_4$ or $\text{C}_{15}\text{H}_{12}\text{O}_5$, a yellowish powder, insol water (Gerhardt, *A* *Ch* [3] 7, 215, Beilstein, *A* 117, 84, Kraut, *A* 156, 124) On heating with glycerin at 100° it forms saliretone $\text{C}_{15}\text{H}_{12}\text{O}_4$ [121.5°] crystallising from water (Giacosa, *J* *pr* [2] 21, 221)

Methyl derivative $\text{C}_6\text{H}_4(\text{OMe})\text{CH}_2\text{OH}$ (248°) *SG* 23 1120 (Cannizzaro a Kerner, *B* 5, 436)

Ethyl derivative $\text{C}_6\text{H}_4(\text{OEt})\text{CH}_2\text{OH}$ (265°) Solidifies at 0° (Botsch, *M* 1, 621)

m Oxy-benzyl alcohol [3 1] $\text{C}_6\text{H}_4(\text{OH})\text{CH}_2\text{OH}$

[67°] (c 300°) A product of the action of sodium-amalgam on *m* oxy benzoic acid in acid solution (Van der Velden, *J* *pr* [2] 15, 163) White mass, v sol hot water FeCl_3 gives a violet colour

Acetyl derivative $\text{C}_6\text{H}_4(\text{OH})\text{CH}_2\text{OAc}$

[55°] (295°–302°) Crystalline, v sl sol water

Di acetyl derivative $\text{C}_6\text{H}_4(\text{OAc})\text{CH}_2\text{OAc}$ (c 290°). Oil, sl alcohol and ether

p Oxy-benzyl alcohol $\text{C}_6\text{H}_4(\text{OH})\text{CH}_2\text{OH}$

[110°] Prepared by slowly adding 40 pts of 8 p sodium amalgam to a solution of 1 pt para-oxy benzaldehyde in 10 pts water and 5 pts alcohol, kept slightly acid with H_2SO_4 (Biedermann, *B*, 19, 2378). Thin white needles v. sol water, alcohol, and ether, sl sol benzene and chloroform, nearly insol ligroin Dissolves in conc H_2SO_4 with a splendid violet colour.

Mono-acetyl derivative

$C_6H_5(OH)CH_2OAc$ [84°], small yellowish needles, v sol al ohol and ether, sl sol water

Di acetyl derivative

$C_6H_5(OAc)CH_2OAc$ [75°], small needles, v sol alcohol and ether, nearly insol water

Methyl derivative $C_6H_5(OMe)CH_2OH$
Anisic alcohol Mol w 198 [45°] Formed, together with anisic acid, by mixing anisic aldehyde with alcoholic potash (Cannizzaro, A 98, 188, 137, 246, G 2, 61) Obtained also by methylation (Biedermann, B 19, 2376) White needles $HClAq$ forms only $C_6H_5(OMe)CH_2Cl$ whence $NaOMe$ forms the compound $C_6H_5(OMe)CH_2OMe$ (226°)

Di-oxy benzyl alcohol *Ethyl derivative* $C_6H_5(OEt)(OH)CH_2OH$ [52 1] [84°] Formed by adding 5 pc sodium amalgam to the corresponding aldehyde suspended in water The product is acidified and shaken with ether (Hantzsch, J pr [2] 22, 475) Large thick tablets, changing at 100° into a brown amorphous mass Acids also resinify it

Di oxy-benzyl alcohol $C_6H_5(OH)_2CH_2OH$ [43 1] **Methyl derivative** $C_6H_5(O)_2CH_2OH$ [43 1] *Vanillyl alcohol* [115°] Formed by the action of sodium-amalgam on vanillin (Tiemann, B 8, 1125, 9, 415) Formed also by the action of emulsion on the glucoside $C_6H_5(OC_6H_4O_2)(OMe)CH_2OH$ [120°], a crystalline body (containing aq) prepared by reduction of glucosyl vanillin (Tiemann B 18, 1595) Prisms, v sol alcohol

Methylene derivative

$C_6H_5(O_2CH_2)CH_2OH$ *Piperonyl alcohol* [51°] Got by reducing piperonal $C_6H_5(O_2CH_2)CHO$ with sodium amalgam and hot water (Fittig a Remsen, A 159, 138) Long crystals, m sol hot water

Reference — CHLORO OXY BENZYL ALCOHOL

OXY BENZYL AMINE $C_6H_5(OH)CH_2NH_2$ [125°] Formed by heating its methyl derivative with $HClAq$ at 150° (Goldschmidt a Ernst, B 23, 2744) and by the action of dilute H_2SO_4 and zinc dust on $C_6H_5(OH)CH_2NHNH_2C_6H_5COH$ (Tiemann, B 23, 3017) Groups of white needles (from ether) Readily sublimates Ferric chloride colours its solution deep violet blue — $B'HCl$ — $B'H_2PtCl_2$, 2aq [197°] Golden needles

Acetyl derivative $C_6H_5(OH)CH_2NHAc$ [140°] Colourless needles, sol alkalis

Methyl derivative $C_6H_5(OMe)CH_2NH_2$ (224° at 724 mm) Formed by reducing the oxim $C_6H_5(OMe)CH_2NOH$ in alcoholic solution with sodium amalgam and $HOAc$ (Goldschmidt a Ernst, B 23, 2742) Liquid, v sol water Yields $C_6H_5(OMe)CH_2NHAc$ [97°] — $B'HCl$ [150°] — $B'H_2PtCl_2$, 2aq [187°] Golden plates

p Oxy benzylamine $C_6H_5(OH)CH_2NH_2$, aq [95°] Formed from p amido benzylamine, $NaNO_2$, and HCl (Salkowski, B 22, 2143) Plates — $B'HCl$ — $B'H_2PtCl_2$, 2aq flat needles

Methyl derivative $C_6H_5(OMe)CH_2NH_2$, (222°) (S), (235°) (G a P) Formed by reducing hydroanisamide in alcoholic solution with sodium amalgam (Steinhart, A 241, 335) Obtained also by reduction from the oxim $C_6H_5(OMe)CH_2NOH$ (Goldschmidt a Polonowski, B 20, 2407) Liquid, sol water, volatile with steam — $B'HCl$ [280°] — $B'HHgCl_2$, aq [300°] Scales — $B'H_2PtCl_2$ [210°] Bright

yellow needles Absorbs CO_2 from the air, forming a compound crystallising in needles [110°] (cf Cannizzaro, A 117, 240)

Acetyl derivative $C_6H_5(OMe)CH_2NHAc$ [96°]

Di-oxy di-benzyl amine $NH(CH_2C_6H_5)_2$, [170°] Formed by reducing hydrosulcalyamide in alcoholic solution with sodium amalgam (Emmerich, A 241, 349) Needles, v sl sol water Gives an oily nitrosamine — $B'H_2PtCl_2$

Di p-oxy di-benzyl amine *Di-methyl derivative* $(C_6H_5(OMe)CH_2)_2NH$ [34°] Formed by the action of $C_6H_5(OMe)CH_2Cl$ on alcoholic ammonia, and also by reducing $(C_6H_5(OMe)OH)_2N_2$ White needles Yields a nitrosamine [80°] — $B'HCl$ [243°]. Flat prisms. — $B'H_2PtCl_2$, 2aq

O OXY BENZYL-ANILINE

$C_6H_5(OH)CH_2.NPhH$ *Phenyl-o amido cresol* [106°] Formed by reducing o oxybenzylidene-aniline with sodium amalgam (Emmerich, A 241, 344) Needles or plates, sl sol water Its nitrosamine is oily — $B'HCl$ [181°] — $B'H_2PtCl_2$ [184°] M sol water

p-Oxy benzyl aniline [208°] Formed in like manner (E) White needles — $B'H_2PtCl_2$

Methyl derivative $C_6H_5(OMe)CH_2.NPhH$ [65°] Formed by reducing $C_6H_5(OMe)CH_2NPh$ (Steinhart, A 241, 337) Prisms Gives a nitrosamine [104°] — $B'HCl$ [163°] — $B'H_2PtCl_2$

DI OXY DI BENZYL BENZENE

$C_6H_5(CHPh)_2$ [171°] Formed by reducing $C_6H_5(COPh)_2$ with sodium amalgam (Wehnen, B 9, 310) Satiny needles (from dilute alcohol) Yields $C_{20}H_{16}AcO_2$ [97°] and $C_{20}H_{16}Ac_2O_2$ [144°]

OXY o BENZYL BENZOIC ACID $C_6H_5O_2$, s.e. $C_6H_5CH(OH)C_6H_5CO_2H$ *Benzhydryl carb-oxylic acid* — KA' amorphous — Baa' . From the anhydride and baryta

Anhydride $C_6H_5CH<\overset{O}{\underset{O}{C}}>CO$ [115°]

Formed by reducing o benzoyl benzoic acid with zinc and HCl (Rotering, J 1875, 596) White insoluble powder

Ezo Oxy m-benzyl-benzoic acid

Formed by reducing m benzoyl benzoic acid with sodium amalgam (Senff, A 220, 242) Satiny needles in hemispherical groups (from hot water). Reduced by $HIAq$ at 170° to m benzyl-benzoic acid — NaA' 4aq — CaA' 5aq — AgA' aq

Oxy-p benzyl-benzoic acid [165°] Formed by reducing p benzoyl benzoic acid (Zincke, A. 161, 102) Needles (from hot water) — NH_4A' — NaA' — KA' — CaA' 5aq — BaA' — AgA' pp

Methyl ether MeA' [110°] Prisms

Ethyl ether EtA' Oil

Oxy benzyl-benzoic acid

$C_6H_5CH_2C_6H_5(OH)CO_2H$ [140°] Formed from benzyl phenol, sodium, and CO_2 (Paterno a Fileti, G 3, 287) Needles (from water), sl sol hot water — AgA' curdy pp, sl sol hot water

DI-OXY BENZYL ETHYL KETONE CARB-OXYLIC ACID *Methylene derivative*

$C_6H_5(O_2CH_2)CH_2CO_2C_6H_5CO_2H$ *Paperketonic acid* [84°] Formed by heating di bromo-piperhydronic acid with aqueous Na_2CO_3 (Weinstein, A 237, 38) Silky needles (from OS_2) — CaA' — AgA' flocculent pp

OXY-BENZYLIDENE-ACETONAMINE a. **ACETONAMINE**

o-OXY-BENZYLIDENE-m-AMIDO-BENZOIC ACID $C_8H_7(OH)CHN(C_6H_4)CO_2H$ [190°]
Formed from salicylic aldehyde and aqueous *m*-amido-benzoic acid (Schriff, *A* 210, 114)
Yellowish needles, *v* *e* sol alcohol

Amide $C_{11}H_9N_3O_2$ [186°] Converted by boiling benzoic aldehyde into crystalline $C_{11}H_9N_3O_2$, whence Ac_2O yields $C_{11}H_9Ac_2N_3O_2$ [220°] The glucosyl derivative of the amide $C_{11}H_{11}O_6$, $C_6H_4CHN(C_6H_4)CONH_2$ [113°] is formed by the action of helicin on *m*-amido-benzamide

Di-oxy-benzylidene-o-amido-benzoic acid
 $C_8H_7(OH)CHN(C_6H_4)(OH)CO_2H$ [5 2 1] [245°]
Formed from oxy amido benzoic acid and salicylic aldehyde. Needles, *v* *e* sol alcohol

DI-OXY-BENZYLIDENE DI-AMIDO-DI-PHENYL $C_{12}H_9N_2[CH(C_6H_4)OH]_2$ [145°] Formed from di-*o*-amido-diphenyl and salicylic aldehyde (Reuland, *B* 23, 3012) Yellow plates

o OXY-BENZYLIDENE-ANILINE
 $C_8H_7(OH)CHNPh$ [51°] *v* Oxy-benzoic ALDEHYDE The *p* isomeride melts at 191°

OXY-BENZYLIDENE ANTHRONE *Ethyl ether* $C_8H_7 \left\langle \begin{smallmatrix} C(Ph OEt) \\ CO \end{smallmatrix} \right\rangle C_6H_4$ [173°]

Formed from bromo benzylidene anthrone and $NaOEt$ (Bach, *B* 23, 2529) Yellow plates, *v* sol ether

OXY - BENZYLIDENE - DICARBAMIC ETHER *Methyl derivative* $C_{11}H_{13}N_2O_4$ *vs* $C_6H_4(OMe)CH(NHCO_2Et)_2$ [172°] Formed from anisic aldehyde and carbamic ether (Bischoff, *B* 7, 1078) Needles (from dil alcohol)

DI-o-OXY-BENZYLIDENE-ETHYLENE-DIAMINE $C_{11}H_{11}N_2O_2$ *vs* $C_6H_4(NCH_2C_6H_4OH)_2$ [126°] Formed from ethylene diamine and *o*-oxy-benzoic aldehyde (Mason, *B* 20, 271) The dimethyl derivative [c 113°] and its *p* isomeride $C_6H_4(NCH_2C_6H_4OMe)_2$ [111°] are both crystalline

o-OXY-BENZYLIDENE-MALONIC ACID *Methyl derivative* $C_8H_7(OMe)CH(CO_2H)_2$ [178°] Formed by heating a mixture of malonic acid, $C_6H_4(OMe)CHO$, and $HOAc$ at 100° (Stuart, *C* J 53, 142)

OXY-BENZYLIDENE NAPHTHYLAMINE
 $C_{12}H_9(OH)CHNC_6H_4$, The *o* [121°] and *p* [220°] compounds are formed from (8) naphthylamine and the corresponding oxy benzoic aldehydes (Emmerich, *A* 241, 350)

o-OXY-BENZYLIDENE DITHIOGLYCOLLIC ACID $C_8H_7(OH)CH(SCH_2CO_2H)_2$ [148°] Formed from *o*-oxy-benzoic aldehyde, thioglycollic acid, and $ZnCl_2$ (Bongartz, *B* 21, 478)

o OXY-BENZYLIDENE p TOLUIDINE
 $C_8H_7(OH)CHNC_6H_4$ [100°] (Jaillard, *J* 1865, 428, *v* OXY-BENZOIC ALDEHYDE)

OXY BENZYLIDENE DI UREA
 $C_8H_7(OH)CH(NH.CO NH_2)_2$, *v* *o*-OXY-BENZOIC ALDEHYDE

OXY-BENZYL-MALONIC ACID *Ethyl derivative* $C_8H_7CH(OEt)CH(CO_2H)_2$ [c 120°] Formed from benzylidene malonic acid and cold alcoholic KOH (Claisen a Crismer, *A* 213, 141) Crystalline. Split up at 120° into alcohol and benzylidene malonic acid [192°] — K_2A — Ag_2A

Isomeride *v* BENZYL-TARTRONIC ACID

OXY BENZYL-METHYL-ETHYL-PYRIMIDINE

INE $CH_2Ph C \left\langle \begin{smallmatrix} N CMe \\ N C(OH) \end{smallmatrix} \right\rangle OEt$ [193 5°]

Formed from phenyl acetamidine hydrochloride, ethyl-acetoacetic ether, dilute (10 p c) $NaOH$, and alcohol (Pinner, *B* 22, 1623) Needles, *m* sol water, *v* *e* sol alcohol

Di-oxy-benzyl-methyl-ethyl pyrimidine

$CH(OH)Ph C \left\langle \begin{smallmatrix} N CMe \\ N C(OH) \end{smallmatrix} \right\rangle OEt$ [148°–152°]

Formed from oxy phenyl acetamidine, acetoacetic ether, and $NaOHAq$ (Pinner, *B* 23, 2951)

OXY BENZYL METHYL-PYRIMIDINE

$CH_2Ph C \left\langle \begin{smallmatrix} N CMe \\ N C(OH) \end{smallmatrix} \right\rangle CH$ [175°] Formed from phenyl acetamidine, acetoacetic ether, alcohol, and dilute (10 p c) $NaOH$ (Pinner, *B* 22, 1622) Prisms, *m* sol hot water

Oxy-benzyl-di methyl pyrimidine

$CH_2Ph C \left\langle \begin{smallmatrix} N CMe \\ N C(OH) \end{smallmatrix} \right\rangle CMe$ [181°] Formed from phenyl acetamidine, methyl acetoacetic ether, and $NaOHAq$ (P) *M* sol water

Oxy-di-benzyl-methyl-pyrimidine

$CHPh C \left\langle \begin{smallmatrix} N CMe \\ N C(OH) \end{smallmatrix} \right\rangle CCH_2Ph$ [192°] Formed from phenyl acetamidine, benzyl acetoacetic ether, and $NaOHAq$ (P) Needles, *insol* water

Di-oxy-benzyl-methyl-pyrimidine $C_{11}H_{11}N_2O_2$, *vs* $CH(OH)Ph C \left\langle \begin{smallmatrix} N CMe \\ N C(OH) \end{smallmatrix} \right\rangle CH$ [216°]

Formed from oxy-phenyl acetamidine, $NaOHAq$ and acetoacetic ether in the cold (Pinner, *B* 23, 2949) Long needles, *v* *sl* sol water, *sl* sol alcohol, sol acids and alkalis — B/HCl [217°] Needles — $B/C_6H_5(NO_2)_2OH$ [175°] $AgC_6H_5N_2O_2$ white pp

Acetyl derivative

$CH(OAc)Ph C \left\langle \begin{smallmatrix} N CMe \\ N C(OH) \end{smallmatrix} \right\rangle CH$ [170°] Formed by boiling with Ac_2O Yields $AgC_6H_5N_2O_2$, B/HCl [188°], and $B/C_6H_5(NO_2)_2OH$ [160°]

Benzoyl derivative $C_{11}H_9BzN_2O_2$, [205°–208°] — B/HCl [240°] From the base and $BzCl$

Di-oxy-benzyl methyl pyrimidine [233°] Got from potassium methyl-uracil and benzyl chloride (Hagen, *A* 244, 1)

Di-oxy benzyl-di methyl pyrimidine

$CH(OH)Ph C \left\langle \begin{smallmatrix} N CMe \\ N C(OH) \end{smallmatrix} \right\rangle CMe$ [155°] Formed from oxy phenyl-acetamidine, $NaOHAq$, and methyl-acetoacetic ether (Pinner, *B* 23, 2951) — $B/HOAc$ needles — $AgC_6H_5N_2O_2$ white pp

OXY - BENZYL - (8) NAPHTHYLAMINE

The following compounds have been prepared by reducing the products of the action of (8) naphthylamine on the corresponding aldehydes (Steinhart, *A* 241, 341, Emmerich, *A* 241, 352) —

1 2] $C_6H_4(OH)CH_2NHC_6H_4$, [147°] — B/HCl [189°]

[1 2] $C_6H_4(OMe)CH_2N(NO)C_6H_4$, [165°]

[1 2] $C_6H_4(OMe)CH_2NH C_6H_4$, [92°] (223°)

[1 4] $C_6H_4(OH)CH_2NHC_6H_4$, [117°]

[1 4] $C_6H_4(OH)CH_2N(NO)C_6H_4$, [142°]

[1 4] $C_6H_4(OMe)CH_2NHC_6H_4$, [101°] — B/HOL

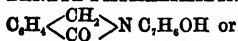
[195°] — B/HCl — *Nitrosamine*

$C_6H_4(OMe)CH_2N(NO)C_6H_4$, [138°]

OXY-BENZYL-ISOPHTHALIC ACID *Benz-hydryl-isophthalic acid Anhydride*

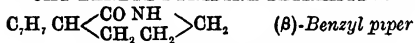
$C_6H_5CH<\begin{smallmatrix} C_6H_5(CO_2H) \\ CO \end{smallmatrix}>$. [207°]. Formed by the action of zinc and HClAq on benzoyl isophthalic acid (Zincke, *B* 9, 1763) Needles (from dilute alcohol) —BaA', 2½ aq —AgA' pulverulent pp. —EtA' [115°] An isomeric acid, obtained by reduction of benzoyl terephthalic acid, forms $Ca(C_{11}H_9O_4)_2$ 8 aq (Weber, *J* 1878, 408)

p-OXY-BENZYL-PHTHALIMIDINE



Formed from the amido compound by the diazo reaction (Hafner, *B* 23, 344) Red needles Converted by conc HClAq at 150° into a base $C_6H_5NO_2$

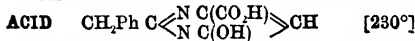
OXY BENZYL PYRIDINE TETRAHYDRIDE



done [118°] Formed by distilling δ amido α-benzyl valeric acid (Aschan, *B* 23, 3696) Pearly plates (from hot water) $B'O_2H_2(NO_2)_2 \cdot OH$ [97°] Crystals, sl sol water

Nitrosamine $C_6H_5(NO)NO$ [62 5°]

OXY BENZYL PYRIMIDINE CARBOXYLIC



Formed from phenyl acetamidine, oxalacetic ether, and (10 p c) NaOHAq (Finner, *B* 22, 1627) Prisms, v sl sol water

OXY BENZYL PYROTARTARIC ACID

$C_6H_5CH(OH)CH(CO_2H)CHMeCO_2H$ Phenylhomotartaric acid Formed from benzoic aldehyde, sodium pyrotartrate, and Ac_2O at 125° (Penfield, *A* 216, 119, Fittig a Liebmann, *A* 255, 257) The acid splits up, at the moment of liberation, into water and anhydride — $CaC_{11}H_9O_4$ 3 aq —BaA' 2 aq —AgA'

Anhydride C_6H_5O Phenylhomoparaconic acid [177°] Plates (from water) — $AgC_{11}H_9O_4$ crystals, m sol water —BaA' 2 aq. —CaA'

Isomeride

$C_6H_5CH(OH)CMe(CO_2H)CH_2CO_2H$ Formed together with the preceding acid Its salt $BaC_{12}H_{10}O_4$ is got by heating the anhydride with baryta water —CaA' aq —AgA' bulky flocculent pp

Anhydride $CHPh<\begin{smallmatrix} CMe(CO_2H) \\ O CO CH_2 \end{smallmatrix}>$. [124 5°] Yields $Ba(C_{12}H_{10}O_4)_2$, CaA', 2 aq, and AgA'

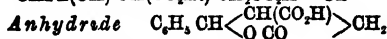
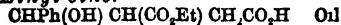
DI-OXY-BENZYL-QUINOLINE CARB-

OXILYIC ACID Ethyl derivative

$C_6H_5<\begin{smallmatrix} CO \\ N \end{smallmatrix}>C(C_6H_5)(CO_2H) \quad [147^\circ]$ Formed by reducing benzyl o nitro benzoyl malonic ether (Bischhoff, *B* 22, 386)

OXY BENZYL SUCCINIC ACID $C_{11}H_9O_4$ 1 e $PhCH(OH)CH(CO_2H)CH_2CO_2H$ Its salts are formed by warming the anhydride with bases — $CaC_{11}H_9O_4$ —BaA' 2 aq —AgA'

Ethyl ether



Phenyl-paraconic acid [99°] Formed by heating sodium succinate with benzoic aldehyde and Ac_2O (Fittig a Jayne, *A* 216, 108, 256, 63) Needles (from water), converted by NaOEt into

phenylitaconic acid. Yields $Ca(C_{11}H_9O_4)_2$ 2 aq, BaA', 2 aq, and AgA', and the ether EtA' (252° at 25 mm)

p OXY-BENZYL-THIOCARBIMIDE

$C_6H_5(OH)CH_2NCS$ From p oxy benzylamine, CS_2 , and $HgCl_2$ (Salkowski, *B* 22, 2144) Liquid, sol alkalis

p-OXY-BENZYL-THIO-UREA Methyl derivative $C_6H_5(OMe)CH_2NHCSNH_2$ [95°] (Goldschmidt a Polonowska, *B* 20, 2409)

Di oxy di benzyl-thio urea $C_6H_5(OMe)CH_2NHCSNH_2$ Di methyl derivative $C_6H_5(OMe)CH_2NHCSNH_2$ [150°]

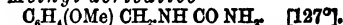
o OXY BENZYL p TOLUIDINE $C_{11}H_9NO$ 1 e $C_6H_5(OH)CH_2NH C_6H_4Me$ [116°] Formed by reducing o oxy benzylidene p toluidine in alcoholic solution with sodium amalgam (Emmerich, *A* 241, 346) Crystals Yields the methyl derivative $C_6H_5(OMe)CH_2NH C_6H_4Me$ [110°] and a tetra nitro derivative [168°] —B'HCl [147°] — $B'_2H_2PtCl_4$ reddish yellow needles

p Oxy benzyl p toluidine [186°] Yields $B'_2H_2PtCl_4$ and $C_6H_5(OMe)CH_2NH C_6H_4Me$ [68°], whence B'HCl [160°], $B'_2H_2PtCl_4$ and the nitrosamine $C_6H_5(OMe)CH_2N(NO) C_6H_4Me$ [108°]

p oxy benzyl o toluidine Methyl derivative [55°] Triangular plates (Steinhart, *A* 241, 340) Yields an oily nitrosamine

o-OXY-BENZYL-UREA $C_6H_5N_2O_4$ 1 e $C_6H_5(OH)CH_2NH CO NH_2$ [170°] Formed by warming o oxy benzylamine hydrochloride with potassium cyanate (Goldschmidt a Ernst, *B* 23, 2745) Prisms, v sol hot water

Methyl derivative



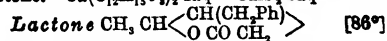
p Oxy benzyl-urea Methyl derivative [167°] Needles (Goldschmidt a Polonowska, *B* 20, 2409)

δ OXY α BENZYL-VALERIC ACID

$CH_2(OH)CH_2CH_2CH(CH_2Ph)CO_2H$ Formed from nitroso oxy benzyl pyridine tetrahydride and NaOHAq (Aschan, *B* 23, 3697). Liquid, m sol hot water

γ Oxy β-benzyl-valeric acid

$CH_2CH(OH)CH(CH_2Ph)CH_2CO_2H$ [76°] Formed by carefully adding dilute HCl to its Ca salt obtained from the lactone (Erdmann, *A* 254, 217) Prisms (containing aq) Melts at 56° when hydrated HClAq converts it into the lactone. — $Ca(C_{11}H_9O_4)_2$ 4 aq —CaA', 6 aq



Got by reducing benzyl acetyl propionic acid [99°] with sodium amalgam, and boiling the product with dilute H_2SO_4 . Large crystals

DI-OXY-BUTANE $CH_3(OH)CH_2(OH)$

n-Butylene glycol Mol w 90 (192°) SG 1.0189 Obtained from α-di-bromo-*n*-butane by boiling with baryta water (Grabowsky a Saytzeff, *A* 179, 325). Liquid, v sol water. Yields glycollic and glyoxylic acids on oxidation

Di-oxy-butane $C_4H_{10}O_2$ 1 e $CH_3(OH)CMe_2OH$ Isobutylene glycol (178°) SG 1.139 Formed from the bromide and K_2CO_3 (Nevolé, *C* R 83, 65, 146) and by the action of HClAq on isobutyl alcohol (Lwoff, *Bl* [2] 43, 112) Formed also in the alcoholic fermentation of sugar (Henninger a Sanson, *C* R 95, 91, 106, 208) Does not form an acetal with aldehyde (Loebert, *A* Ch, [6] 16, 58).

Di-oxy-butane $\text{CHMe}(\text{OH})\text{CHMe}(\text{OH})$ (184°)

Formed by heating *s*-butylene oxide with water at 100° (Eiltekoft, *J R* 14, 372) Liquid

Di-oxy-butane $\text{CHMe}(\text{OH})\text{CH}_2\text{CH}_2\text{OH}$ (*B*-

Butylene glycol. (207°) SG \pm 1.0259 Formed by reduction of a dilute, slightly acid, solution of aldol by sodium-amalgam (Kekulé, *B* 5, 56, *A* 162, 810, Wurtz, *C R* 97, 478) Thick liquid, miscible with water Ac_2O at 100° forms $\text{C}_4\text{H}_8(\text{OAc})_2$ (207°) SG \pm 1.055 HIAq yields $\text{C}_4\text{H}_8\text{I}_2$ SG \pm 2.291

Di-oxy-butane $\text{C}_4\text{H}_8(\text{OH})_2$ (184°) SG \pm 1.048 Obtained, *via* $\text{C}_4\text{H}_8(\text{OAc})_2$ (c 200°), from the crude $\text{C}_4\text{H}_8\text{Br}_2$ got from fusel oil (Wurtz, *A Ch*. [3] 55, 452) Liquid, miscible with water

Di-oxy-butane $\text{CH}_2(\text{OH})\text{CH}_2\text{CH}_2\text{CH}_2(\text{OH})$ *Tetramethylene glycol* (204°) SG 1.011 Formed by the action of dilute H_2SO_4 on tetramethylene diamine $\text{C}_4\text{H}_{10}(\text{NH}_2)_2$ (Dekkers, *R. T C* 9, 101)

Tri-oxy-butane

$\text{CH}_2\text{CH}(\text{OH})\text{CH}(\text{OH})\text{CH}_2(\text{OH})$ *Butenylglycerol* (178° at 27 mm) Formed from crotonic aldehyde by reduction, addition of bromine, and boiling the resulting $\text{CH}_3\text{CHBrCHBrCH}_2\text{OH}$ with water (Lieben & Zeisel, *M* 1, 832) Thick liquid with sweet taste Ac_2O yields $\text{C}_4\text{H}_8(\text{OAc})_3$ (262°) Yields $\text{C}_4\text{H}_8(\text{OH})_3\text{Cl}$, $\text{C}_4\text{H}_8(\text{OH})\text{Cl}_2$, and $\text{C}_4\text{H}_8\text{OCl}_2$ (Zikes, *M* 6, 348)

Tri-oxy-butane $\text{C}_4\text{H}_8\text{O}_3$ (240° at 18 mm) Formed from isobutyl iodide by chlorinating and heating the resulting tri-chloro butane with water at 170° (Prunier, *Bl* [2] 42, 261, *C R* 99, 193) Yields nearly solid $\text{C}_4\text{H}_8(\text{OAc})_3$

Tetra-oxy-butane *v* **ERYTHRITE**

OXY-BUTANE TRICARBOXYLIC ACID $\text{CMe}(\text{OH})(\text{CO}_2\text{H})\text{CH}(\text{CO}_2\text{H})\text{CH}_2\text{CO}_2\text{H}$

Lactone $\text{C}_4\text{H}_6\text{O}_4$ Formed by heating acetosuccinic ether with KCy and HCl (Rach, *A* 284, 36) At 180° it yields pyrocinchonic anhydride — BaA'' . — CaA'' — $\text{Ca}_2(\text{C}_4\text{H}_6\text{O}_4)_n$ — $\text{Ba}_2(\text{C}_4\text{H}_6\text{O}_4)_n$

Di-oxy-butane tetra-carboxylic ether $\text{CH}(\text{CO}_2\text{Et})_2\text{CH}(\text{OH})\text{CH}(\text{OH})\text{CH}(\text{CO}_2\text{Et})_2$ Formed by the action of malonic ether on glyoxal in presence of conc ZnCl_2 Aq (Polonowsky, *A* 246, 2) Oil, not volatile with steam

Tetra oxy butane tri carboxylic acid $\text{CO}_2\text{HCH}(\text{OH})\text{CH}(\text{OH})\text{CH}(\text{OH})\text{O}(\text{OH})(\text{CO}_2\text{H})_2$ [147°] Formed by oxidation of levulose carb-oxylic acid by dilute HNO_3 (Dull, *B* 24, 848) — $\text{K}_2\text{HA}''$ Large prisms — $\text{Ca}_2\text{A}''$, 6aq

OXY-BUTANE PHOSPHONIC ACID

$\text{O}_2\text{HCH}(\text{OH})\text{PO}(\text{OH})_2$ [168°] Formed from isobutyric aldehyde by successive treatment with PCl_3 and water (Fossek, *M* 5, 640) Trimetric crystals, $a b c = 97.1394$

OXY-ISOBUTANE SULPHONIC ACID $\text{CMe}(\text{OH})\text{CH}_2\text{SO}_3\text{H}$ Formed from ammonium sulphite solution and isobutylene bromide or $\text{CH}_3\text{Br CMe}_2\text{OH}$ (Guareschi & Garzino, *Ann. chim. farm.* [4] 6, 110, 9, 96) — BaA' , 1½aq — NaA' plates (from alcohol)

DI-OXY BUTYL-BENZENE $\text{C}_{10}\text{H}_{18}\text{O}_2$ *s.s* $\text{C}_4\text{H}_9\text{CH}(\text{OH})\text{C}_2\text{H}_5\text{CH}_2\text{OH}$ (200°) Formed by the action of sodium-amalgam on an alcoholic solution of $\text{C}_4\text{H}_9\text{CO}_2\text{C}_2\text{H}_5\text{CHO}$ which is got from phenyl propyl ketone by successive treatment with CrO_3Cl_2 and water (Burcker, *C R* 94, 220). Syrup. Forms only $\text{C}_{10}\text{H}_{18}(\text{OAc})_2$

ISOBUTYL o-OXY BENZOIC ACID

$\text{C}_4\text{H}_9(\text{C}_2\text{H}_5)(\text{OH})\text{CO}_2\text{H}$ [421] Formed from $\text{C}_4\text{H}_9(\text{C}_2\text{H}_5)\text{ONa}$ and CO_2 at 140° (Dobrzycki, *J pr* [2] 86, 391) Needles — CaA' , 6aq — BaA' , 2aq needles, v sol water

Ethers MeA' [54°] (266°) — EtA' (276°) Oil — $\text{C}_4\text{H}_9\text{A}'$ [68°] Formed from the acid, phenol, and POCl_3 Converted by long boiling into $\text{C}_{11}\text{H}_{18}\text{O}_2$ [158°]

DI OXY-BUTYLENE $\text{C}_4\text{H}_8(\text{OH})_2$ *Crotonylene glycol* (197°) SG \pm 1.0616, \pm 1.0465 Obtained by the action of boiling baryta on its formyl derivative which is formed when erythrite is distilled with formic acid (Henninger, *B* 5, 1060, *A Ch* [6] 7, 215) Liquid, sol water Yields $\text{C}_4\text{H}_8(\text{OAc})_2$ (208°)

DI-OXY-DI BUTYL DIKETONE

Anhydride $\begin{array}{c} \text{CH}_2 \quad \text{CH}_2 \\ \diagdown \quad \diagup \\ \text{O} \end{array} > \text{C} < \begin{array}{c} \text{CH}_2 \quad \text{CH}_2 \\ \diagup \quad \diagdown \\ \text{O} \end{array} \text{CHMe}$

Di methyl-oxetone (169.5° V) SG \pm 0.978 Formed, together with CO_2 , by heating its carb

oxylic acid $\begin{array}{c} \text{CH}_2 \quad \text{CH}(\text{CO}_2\text{H}) \\ \diagdown \quad \diagup \\ \text{O} \end{array} > \text{C} < \begin{array}{c} \text{CH}_2 \quad \text{CH}_2 \\ \diagup \quad \diagdown \\ \text{O} \end{array} \text{CHMe}$ (divalonic acid) [130°] which is got by the action of NaOHAq at 90° on 'divalolactone,' the product of the action of NaOHAq on valerolactone (Fittig, 256, 128) Liquid Volatile with steam

OXY BUTYL MALEIC ACID *Lactone*

$\text{CHPr} < \begin{array}{c} \text{C}(\text{CO}_2\text{H})_2 \\ \diagup \quad \diagdown \\ \text{O} \end{array} > \text{CH}$ *Propaconic acid*

[124°] Formed by distilling the bromide of propyl itaconic acid with steam, the acid remains behind and is extracted with ether (Fittig, *A* 256, 108) — Needles — BaA'

OXY BUTYL MALONIC ACID $\text{C}_4\text{H}_8\text{O}_4$ *s.s* $\text{C}_4\text{H}_9\text{C}(\text{OH})(\text{CO}_2\text{H})_2$ [110°–114°] Obtained from $\text{C}_4\text{H}_9\text{CCl}(\text{CO}_2\text{Et})_2$ and KOHAq (Conrad & Bischoff, *B* 13, 600, 14, 617, Guthzeit, *A* 209, 237) Deliquescent mass, v e sol water

DI OXY ISOBUTYL METAPYRAZOLE

$\text{C}_4\text{H}_7\text{N}_2\text{O}_2$ *s.s* $\begin{array}{c} \text{C}_4\text{H}_7\text{CH} - \text{N} \\ | \quad \diagup \\ \text{C}(\text{OH}) \quad \text{N} \end{array} > \text{C}(\text{OH})$ [210°]

Obtained by boiling with dilute HCl the product ($\text{C}_4\text{H}_7\text{CH}(\text{CN})\text{NHCO}_2\text{NH}_2$) of the action of urea upon valeric aldehyde-cyanhydrin Small white needles M sol alcohol and hot water, sl sol cold water Dissolves readily in alkalis (Pinner & Lufschütz, *B* 20, 2356)

OXY BUTYL SUCCINIC ACID The salts are got by the action of bases on the anhydride $\text{Ca}(\text{C}_4\text{H}_7\text{O}_4)_2$ 5aq — BaA'' 2aq — AgA'' *Anhydride* $\text{CHPr} < \begin{array}{c} \text{CH}(\text{CO}_2\text{H})_2 \\ \diagup \quad \diagdown \\ \text{O} \end{array} > \text{CH}_2$ *Propyl paraconic acid* [73.5°] Formed by heating butyric aldehyde with sodium succinate and Ac_2O (Schmidt, *A* 255, 68) Needles (from ligroin) Yields heptenoic acid and oxy heptenoic lactone on distillation — $\text{Ca}(\text{C}_4\text{H}_7\text{O}_4)_2$ 2aq — BaA' , — AgA' . *Ethyl ether* EtA' (c 214° at 96 mm)

Oxy-isobutyl-succinic acid *Salts.* — $\text{Ba}(\text{C}_4\text{H}_7\text{O}_4)_2$ 2aq — $\text{Ag}_2\text{A}''$

Anhydride Formed, like the preceding isomeric, using isobutyric aldehyde (Zanner, *A* 255, 86) — $\text{Ba}(\text{C}_4\text{H}_7\text{O}_4)_2$ 3aq — CaA' 2aq — AgA'

OXY-BUTYRAMIDINE $\text{C}_4\text{H}_9\text{N}_2\text{O}_2$ *s.s* $\text{CMe}(\text{OH})\text{C}(\text{NH})\text{NH}_2$ The crystalline hydrochloride is formed from oxy-butyrimido-ethyl ether and NH_3 (Pinner, *B* 17, 2009) It is v. e. sol water

α-OXY-*n*-BUTYRIC ACID $C_4H_7O_3$, 18
 $CH_3CH(OH)CO_2H$, Mol w 104 [43°] Formed by the action of moist Ag_2O upon bromo-*n*-butyric acid (Naumann, *A* 119, 115, Friedel & Machuga, *A* 120, 279), and by the action of HCl and HCl on propionic aldehyde (Prachibiteck, *B* 9, 1312) Formed also by reduction of ethylglyoxylic acid Deliquescent crystals Yields propionic acid on oxidation (Markownikoff, *A* 176, 309, Ley & Popoff, *A* 174, 81) — CaA' , 6aq ZnA' , 2aq S 2.35 at 18° — AgA' prisms

Ethyl ether EtA' (167° 1 V) SG 2 1.004, 12 995 (Schreiner, *B* 12, 177, *A* 197, 21). Yields ethyl glyoxylic ether on oxidation (Aristoff & Demjanoff, *C* 1887, 1157) Acetyl derivative $CH_3CO(OAc)CO_2Et$ (198°) (Gal, *A* 142, 373) Butyryl derivative (215°)

Methyl derivative $CH_3CO(OMe)CO_2H$ Formed from its Me and Et ethers which are made by the action of $NaOMe$ on bromo butyric ether (Duvillier, *C* 86, 47, 1026, 87, 981, 88, 598, *A* *Ch* [5] 17, 528) Liquid, sol water — AgA' — MeA' (150°–155° 1 V) — EtA' (160°) (D), (148° 1 V) (Schreiner, *A* 197, 16)

Ethyl derivative $CH_3CO(OEt)CO_2H$. Formed from its ether, which is made from bromo butyric ether and $NaOEt$ Liquid, v sol water — KA' — BaA' — AgA' — MeA' (157°) — EtA' (168°–174°) (D), (169° 1 V) (S)

Methyl derivative of the amide $CH_3CO(OH)CONH_2$ [78°] Formed from $CH_3CO(OH)CO_2Me$ and alcoholic NH_3 (D) Slender needles, v sol water

Ethyl derivative of the amide $CH_3CO(OEt)CONH_2$ [69°] Laminæ, sol water **β-Oxy-*n*-butyric acid**

$CH_3CH(OH)CH_2CO_2H$ Occurs in urine and blood of diabetic patients (Kulz, *Zeit Biol* 20, 165, 23, 329, Munkowski, *Er* 24, 153, Stadelmann, *Zeit Biol* 32, 456, Wolpe, *C* 1887, 277, Hugouenq, *Bl* [2] 47, 645, Deichmuller, Szymanski, *A* Tollens, *A* 228, 92) Formed by reducing acetoacetic acid with sodium amalgam (Wislicenus, *A* 149, 205), and from propylene chlorhydrin by successive treatment with KCy and KOH (Markownikoff, *A* 153, 237) Obtained also by oxidising aldol with moist Ag_2O (Wurtz, *C* 76, 1165) Thick syrup, volatile with steam Decomposes at 130° into water and (α)-crotonic acid When prepared from urine it is levorotatory, $[α]_D = -23.4$ — NaA' very deliquescent needles — ZnA' — CuA' — AgA'

Ethyl derivative $CH_3CH(OEt)CH_2CO_2H$ (c 215°) Formed by the action of HCl on the nitrile, which is formed by combining allyl cyanide with alcohol (Pinner, *B* 12, 2057)

Amide $CH_3CH(OEt)CH_2CONH_2$ [71°]

γ-Oxy-*n*-butyric acid $CH_3CH(OH)CH_2CH_2CO_2H$ Obtained by the action of boiling lime- or baryta water upon its lactone, which is formed by treating succinyl chloride, dissolved in $HOAc$ and ether, with sodium-amalgam (Saytzeff, *B* 6, 1255, *A* 171, 270, *J* pr [2] 25, 66, *Bl* [3] 37, 540) Formed also from $CH_3BrCH_2CH_2OH$ by successive treatment with alcoholic KCy and potash (Fruhling, *M* 8, 700), and by boiling oxy-ethyl-acetoacetic ether with conc baryta-water (Ochanlaroff, *A* 226, 825) Liquid, which volatilises in the cold Volatile with steam Sol water. Forms the

lactone slowly in the cold, more quickly on heating Chromic acid mixture oxidises it to succinic acid — KA' deliquescent tufts — NaA' — ZnA' (dried at 100°) — BaA' (dried at 110°) dendritic mass (from alcohol)

Lactone $CH_3CH_2CH_2CO$ (206°). SG

§ 1 1441, 1/2 1 1286 CE (0° to 16°) 00086 **Butyrolactone** Formed as above Formed also by heating the lactonic acid of γ oxy ethyl malonic acid at 120° (Röder, *A* 227, 22), and by heating γ chloro butyric acid at 200° (Henry, *C* 101, 1158) Mobile liquid, miscible with water, but separated therefrom by K_2CO_3 . May be converted into *n*-butyric acid by successive treatment with HI and sodium amalgam

α-Oxy isobutyric acid $(CH_3)_2C(OH)CO_2H$ **Acetonic acid** **Butylactic acid** [79°] (212°)

Formation —1 From acetone, HCl , and $HClAq$ (Stadeler, *A* 111, 320) —2 From bromo isobutyric acid and moist Ag_2O or Na_2CO_3Aq (Markownikoff, *A* 146, 339, 153, 228, 251, Fittig, *A* 200, 70) —3 From dimethyl oxalate by treatment with $ZnMe$, followed by water (Frankland & Duppa, *A* 133, 80, 135, 25) —4 By oxidising isobutyric acid with alkaline $KMnO_4$ (R Meyer, *A* 219, 240) —5 By oxidising di oxy pentane (amylene glycol) with diluted HNO_3 (Wurtz, *A* 107, 197) —6 By heating acetone chloroform with water at 180° (Willgerodt, *B* 15, 2807, *Bl* [2] 89, 157) Hygroscopic needles, v sol water, alcohol, and ether Volatile with steam Sublimes at 50° Yields acetone and acetic acid on oxidation Phenyl hydrazine at 160° forms a ψ-phenyl hydrazide [152°] converted by nitrous acid into a nitrosamine [98°] (Reissert & Kayser, *B* 22, 2926)

Salts — BaA' — ZnA' , 2aq hexagonal plates, sl sol water S 6 at 15° — AgA' stellate groups of nacreous scales S 7

Ethyl ether EtA' (151°)

Ethyl derivative $(CH_3)_2C(OEt)CO_2H$ (180°) SG § 1 0211, 1/2 1 0101 Formed from CMe_2BrCO_2H and $KOEt$ (Hell & Waldbauer, *B* 10, 449) — BaA' , aq — PbA' , aq — ZnA' — AgA' plates in sol water — EtA' (155°)

Isopropylidene derivative $C_6H_{12}O_4$ SG SG (197° uncor) VD 120.3 (calc 124) Formed by the action of KOH (8 mols) on acetone chloroform (2 mols) and acetone (1 mol), or upon a mixture of chloroform (1 mol) and acetone (2 mols) (Willgerodt, *B* 20, 2445, Engel, *C* 104, 688) Liquid, converted into oxy isobutyric acid by heating with water — CaA' 1/2 aq — BaA' 1/2 aq — PbA' (W), PbA' 2aq (E) — ZnA' aq small scales (W), ZnA' 2aq (E).

Nitrile v ACETONE CYANHYDRIN, vol i p 31 **Anhydride** $C_4H_4O_3$, s.s. $O(CMe_2CO_2H)_2$

Diethyl lacto acid A product of the action of alcoholic potash on chloro isobutyric acid (Balbiano, *J* 1873, 704, 1880, 789) Amorphous, v sol water — NaA' deliquescent

α,β-Di-oxy-butyric acid

$CH_3CH(OH)CH(OH)CO_2H$ [80°]. Formed by boiling bromo-oxy butyric acid with water (Kolbe, *J* pr [2] 25, 390), and by heating β-methylglycidic acid with water at 100° in sealed tubes (Melikoff, *J* R. 18, 526, *B* 21, 2055). Mass of slender needles, v sol water, not volatile with

steam.—AgA' needles Gives a silver mirror when boiled with water.

Ethyl ether EtA' (c 228°) Liquid.

Anhydride v. METHYL-GLYCIDIC ACID

γ-Di-oxy-butyric acid

$\text{CH}_2(\text{OH})\text{CH}(\text{OH})\text{CH}_2\text{CO}_2\text{H}$ *Butylglyceric acid* Formed from $\text{CH}_2(\text{OH})\text{CH}(\text{OH})\text{CH}_2\text{Cl}$ by successive treatment with KOy and dilute HNO_3 (Hanriot, *A Ch* [5] 17, 106) Formed also by warming its anhydride with water (Melikoff, *B* 15, 2587) Thick liquid, v sol water, alcohol, and ether —BaA', amorphous

Anhydride $\text{CH}_2\text{CH}(\text{CO}_2\text{H})\text{CH}_2\text{CO}_2\text{H}$ *Butylglycidic acid* Formed from (8) crotonic acid by successive treatment with HOCl and alcoholic potash Mobile liquid Unites with HCl, forming chloro oxy butyric acid [99°] The ether $\text{C}_2\text{H}_5\text{EtO}$, (145°–150°), SG 215 9931 is formed by the action of sodium amalgam on a mixture of epichlorhydrin and chloroformic acid (Kelly, *B*, 11, 2225)

Di-oxy-isobutyric acid $\text{CH}_2(\text{OH})\text{CMe}(\text{OH})\text{CO}_2\text{H}$ [100°] Formed from a methyl glycidic acid (q v) by warming with water at 100° (Melikoff, *J R* 15, 535) Prisms, v sol water —KA' 3aq small prisms

Tri-oxy-isobutyric acid $(\text{CH}_2(\text{OH}))_2\text{C}(\text{OH})\text{CO}_2\text{H}$ [116°] Formed from glycerose by successive treatment with HCl and HCl (Fischer a Tafel, *B* 22, 106) Prisms (from alcohol), insol benzene HIAq followed by zinc dust yields isobutyric acid —CaA', 4aq —PbA', aq needles, sl sol hot water

A tri-oxy-butyric acid is also formed in the oxidation of gallic acid and tannin by dilute HNO_3 (Böttinger, *A* 257, 248)

References —Bromo- and Chloro-oxybutyric acids.

β-OXY-*n*-BUTYRIC ALDEHYDE v. ALDOL.

α-OXY-ISOBUTYRIC ALDEHYDE

$\text{CMe}_2(\text{OH})\text{CHO}$ (c 90°) Obtained by boiling with HOAc the compound $\text{C}_{11}\text{H}_{22}\text{O}$, formed by the action of NaOC_2H_5 on iodoform (Gorboff, *J pr* [2] 41, 243) Mobile liquid, oxidised by Ag_2O to oxy-isobutyric acid [79°] Forms with water a hydrate [c 70°] Polymerises on standing, forming prisms [c 65°] v sol water

Di-isobutyl derivative of oxyisobutyric orthoaldehyde $\text{CMe}_2(\text{OC}_2\text{H}_5)_2\text{CH}(\text{OH})(\text{OC}_2\text{H}_5)$ (125°) SG 2 9041 One of the products of the action of iodoform (or iodine) on NaOC_2H_5 . Oil, with bitter taste Converted by HOAc into isobutyl acetate and oxyisobutyric aldehyde

OXY BUTYRIC IMIDO-ETHYL ETHER $\text{CMe}_2(\text{OH})\text{C}(\text{NH})\text{OEt}$ The hydrochloride $\text{C}_2\text{H}_5\text{NO}_2\text{HCl}$ is formed from acetone, HCl, alcohol, and HCl (Pinner, *B* 17, 2009) It is crystalline

OXYBUTYROCYNAMINE v GUANIDO BUTYRIC ACID

OXY-CAFFEINE v. CAFFEINE

OXY-CAMPHOR v. CAMPHOR and CAMPHOLENIC

OXY-CAMPHORAMIC ACID v. CAMPHORIC ACID

OXY-CAMPHORIC ACID v. CAMPHORIC ACID.

OXY-CAMPHORIC ANHYDRIDE v. CAM-

PHANIC ACID

OXY-CAMPHORONIC ACID v. CAMPHORONIC ACID.

OXY CAPROIC ACID v Oxy-HEXOIC ACID

OXYTRICARBALLYLIC ACID *Methyl derivatives* $\text{C}_2\text{H}_5(\text{OMe})(\text{CO}_2\text{H})_3$ Formed by heating $(\text{C}_2\text{H}_5)_3\text{C}(\text{OMe})\text{CO}_2\text{H}$ with dilute HNO_3 (Schatzky, *J R* 17, 85) Syrup —CaH₂A', 3aq —BaH₂A', 2aq

OXY-CARBOSTYRIL v Di OXY-QUINOLINE

OXY-CARBOXYLIC ACID v BENZENE TRI-QUINONE

OXY-*o* CARBOXY-PHENYL-ACETIC ACID

$\text{CO}_2\text{H}\text{C}_6\text{H}_4\text{CH}(\text{OH})\text{CO}_2\text{H}$ Formed from $\text{CO}_2\text{H}\text{C}_6\text{H}_4\text{CO}\text{CO}_2\text{H}$ and sodium amalgam (Scherks, *B* 18, 381) Quickly changes, when set free, to the anhydride (phthalide carboxylic acid) [149.5°]

Oxy-carboxy-di-phenyl acetic acid

$\text{C}_6\text{H}_5(\text{CO}_2\text{H})\text{CPh}(\text{OH})\text{CO}_2\text{H}$ [c 80°] Formed by warming benzil *o* carboxylic acid with alkalis (Graebe, *B* 21, 2003) —KA' 2aq

Di-oxy-di-carboxy-phenyl acetic acid *Triethyl-ether* $\text{C}_2\text{H}_5(\text{OH})_2(\text{CO}_2\text{Et})_2\text{CH}\text{CO}_2\text{Et}$ [98°] Formed from acetone dicarboxylic ether and sodium (Cornelius a Pechmann, *B* 19, 1448) Needles (from alcohol), insol water

γ-OXY-*o* CARBOXY-PHENYL-BUTYRIC ACID *Lactone* $\text{C}_{11}\text{H}_{12}\text{O}_4$ [121°] Formed by reducing the anhydride of carboxy benzoyl propionic acid with sodium amalgam (Roser, *B* 17, 2773) —Ag $\text{C}_{11}\text{H}_{12}\text{O}_4$ —Ag $\text{C}_{11}\text{H}_{12}\text{O}_4$

An isomeric lactone [140°] is obtained by reducing phthalyl propionic acid (Gabriel a Michael, *B* 11, 1681) —Ba $\text{C}_{11}\text{H}_{12}\text{O}_4$ —Ba $(\text{C}_{11}\text{H}_{12}\text{O}_4)_2$ —Ag $\text{C}_{11}\text{H}_{12}\text{O}_4$

OXY-CARBOXY-PHENYL-PROPIONIC ACID *Lactone* $\text{C}_{10}\text{H}_{10}\text{O}_4$

$\text{C}_6\text{H}_5\text{CH}(\text{CO})\text{CH}_2\text{CO}_2\text{H}$ Formed by reducing

phthalyl acetic acid with sodium amalgam, and acidifying (Gabriel a Michael, *B* 10, 1558, 2200) Needles (containing aq) Yields Ag $\text{C}_{10}\text{H}_{10}\text{O}_4$, Ag $\text{C}_{10}\text{H}_{10}\text{O}_4$, and Ba $\text{C}_{10}\text{H}_{10}\text{O}_4$ 2aq

Tri oxy-carboxy-phenyl propionic acid

Lactone $[4\frac{1}{2}] \text{C}_6\text{H}_5(\text{OH})_2\text{CH}(\text{CO})\text{CH}_2\text{CO}_2\text{H}$

[228°] Formed by reducing its di methyl derivative (meconic acetic acid) with HI and P (Liebermann a Kleemann, *B* 19, 2293) Tables Gives a blue colour with FeCl₃

Ethyl ether EtA' [c 131°] Crystalline

OXYCHRYSOQUINONE v CHRYSOQUINONE

OXYCINCHENE v CINCHENE

OXYCINCHONIDINE v CINCHONIDINE

OXY-CINNAMIC ACID v COUMARIC ACID

α-Oxy cinnamic acid $\text{C}_6\text{H}_5\text{CH}(\text{CO}_2\text{H})\text{CO}_2\text{H}$

Formed in small quantity in the preparation of the isomeric phenylglycidic acid by the action of alcoholic potash on $\text{C}_6\text{H}_5\text{CHBrCH}(\text{OH})\text{CO}_2\text{H}$ (Plöchl, *B* 16, 2821) Readily decomposes, yielding phenyl-acetic aldehyde and di-oxy-phenyl propionic acid

Phenyl derivatives

$\text{C}_6\text{H}_5\text{CH}(\text{COPh})\text{CO}_2\text{H}$ [180°] Formed by heating sodium phenoxy-acetate with benzoil aldehyde and Ac_2O (Oghaloro, *O J*, 40, 276) Prisms —AgA' prisms (from water)

(8, 2, 1)-Di-oxy-cinnamic acid

$[8\frac{2}{2}]\text{C}_6\text{H}_5(\text{OH})_2\text{CHCHCO}_2\text{H}$

Anhydride $\text{C}_6\text{H}_5(\text{OH})_2\text{CHCHCO}_2\text{H}$ *m-Oxy-*

coumarin [280°-285°] Formed by heating pyrocatechin with ihalic acid and H_2SO_4 (Bezari, *B* 15, 34) Needles, sl sol cold water Reduces salts of Au, Cu, and Ag

(4, 3, 1) Di-oxy-cinnamic acid *v* **CAFFEIC ACID**

(5, 2, 1)-Di-oxy cinnamic acid
[5 2 1] $C_6H_4(OH)_2CH CH CO_2H$

Anhydride [5 1] $C_6H_4(OH) < \begin{smallmatrix} CH & CH \\ & O & CO \end{smallmatrix}$

[250°] Formed by heating a mixture of hydroquinone, malic acid, and H_2SO_4 (Pechmann & Welsh, *B* 17, 1648) Needles, v sol alcohol Yields an acetyl derivative [147°] crystallising in needles The methyl derivative [108°] is got from [5 2 1] $C_6H_4(OMe)(OH)CHO$ by boiling with Ac O and NaOAc (Tiemann & Muller, 14, 1996)

o Methyl derivative

[5 2 1] $C_6H_4(OH)(OMe)CH CH CO_2H$ [180°] Obtained from $C_6H_4(NH_2)(OMe)CH CH CO_2H$ by the diazo reaction (Schnell, *B* 17, 1387) Crystals

Di-methyl derivative

$C_6H_4(OMe)_2CH CH CO_2H$ [143°] Obtained by methylating the *o* methyl derivative Needles Yields di methyl gentisic aldehyde on oxidation with $KMnO_4$

(4, 2, 1) Di oxy-cinnamic acid *v* **UMBELLIC ACID**

ap di oxy cinnamic acid Phenyl-methyl derivative $C_6H_4(OMe)CH C(OPh)CO_2H$ [200°] A product of the action of anisic aldehyde and Ac_2O on sodium phenoxy acetate (Valentini, *G* 14, 147) Rectangular tablets, sol hot alcohol —MeA' [100°] Laminæ

Anhydride of the phenyl derivative $C_{11}H_{10}O_3$ [113°] Formed from salicylic aldehyde, Ac_2O , and sodium phenoxy acetate (Ogliaro, *C* 1887, 1164) Yellow prisms, v sl sol hot water

Tri oxy cinnamic acid *v* **ÆSCULETIC ACID** and **DAPHNETIN**

Tetra-oxy cinnamic acid *Di methyl methylene ether* $C_6H_4(O_2CH_2)(OMe)_2CH CH CO_2H$ [196°] Formed from apiole aldehyde, Ac_2O , and NaOAc (Camician & Silber, *B* 22, 2485) Small yellow needles (from hot alcohol), sl sol ether and hot water

OXY-CINNAMIC ALDEHYDE *v* **COUMARIC ALDEHYDE**

Di-oxy cinnamaldehyde *Methyl derivative* *v* **FERULIC ALDEHYDE**

OXY-CINNOLINE

$C_8H_7N_2O$ *vs* $C_8H_7N < \begin{smallmatrix} C(OH) & CH \\ & N \end{smallmatrix}$ [225°] Formed by heating its carboxylic acid at 260° (Richter, *B* 16, 681) Small prisms, v sol alcohol and ether, sl sol water May be sublimed Sol Na_2CO_3 Aq —B' H_2PtCl_6 small prisms

Oxy-cinnoline carboxylic acid

$C_8H_7N < \begin{smallmatrix} C(OH) & C & CO_2H \\ & N \end{smallmatrix}$ [c 265°] Formed from *o* diazo phenyl propionic acid by heating with water at 70° Colourless needles or scales, sol HCl Aq, sl sol alcohol, nearly insol water

OXY-CITRAGONIC ACID $C_6H_8O_4$ *vs*

$O < \begin{smallmatrix} CMe & CO_2H \\ & CH & CO_2H \end{smallmatrix}$ [162°] Formed from citraconic acid by successive treatment with HCl and alkalis (Morawski, *J pr* [3] 10, 69, 11,

480, Scherko, *A* 227, 233, Michael, *J pr* [2] 40, 171, Melikoff & Feldmann, *A* 263, 89) Prisms (containing aq) Hot water converts it into propionic aldehyde and CO_2 With HBr it forms C_6H_5BrO , [156°] HCl yields, in like manner, CO_2H $CClMe$ $CH(OH)CO_2H$ [162°]

Salts — $(NH_4)A'$ — $(NH_4)HA'$ — KHA' — BaA'' 4aq — SrA'' 4aq — PbA'' 4aq

Ethyl ether EtA' (255°) S G § 1 1376; § 2 11167 CE (0°-22°) 0008507

OXY-CITRIC ACID $C_6H_8O_7$ *vs*

$C_6H_4(OH)_2(CO_2H)_2$ Occurs in beetroot (Lippmann, *B* 16, 1078) Formed from asconitic acid by successive treatment with HCl and lime-water (Pawolleck, *A* 178, 150) Deliquescent mass, v *e* sol water —Salts BaA''' , 5aq — Ca_2A''' , 9aq — Ca_3A''' , 10aq — $Cd_4(C_6H_4O_7)_2$, 6aq — $Cu_4(C_6H_4O_7)_2$, 2aq — EtA''' oil with bitter taste.

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OXYCOMENIC ACID *v* **COMENIC ACID**

OXY CONICEINE *v* **CONINE**

OXY-COPAIVIC ACID *v* **COPAIVIC ACID**

OXY COUMARIC ACID *v* **DI OXY CINNAMIC ACID**

OXY-COUMARILIC ACID *v* **COUMARILIC ACID**

OXY-COUMARIN *v* *Anhydride of Di-oxy-cinnamic acid and UMBELLIFERONE*

Di-oxy coumarin *v* **DAPHNETIN**

OXY COUMARONE *v* **COUMARONE**

β OXY CROTONIC ACID *Methyl derivative* $CH_3C(OMe)CH CO_2H$ [128 5°] Formed from β chloro crotonic acid and NaOMe (Friedrich, *A* 219, 327, 334) Crystals, insol water

Ethyl derivative $CH_3C(OEt)CH CO_2H$ [137 5°] Formed from $CH_3CClCH CO_2Et$ and alcoholic potash Prisms, v sl alcohol and ether —KA' plates —KA' 3aq needles —EtA' [30°] Decomposed by dilute H_2SO_4 into acetone, CO_2 , and alcohol

Phenyl derivative $CH_3C(OPh)CH CO_2H$ [150°] Formed by heating sodium β chlorocrotonate or β chloro isocrotonate with NaOPh (Autenrieth, *A* 254, 240) Crystals, v sl sol hot water Yields $CH_3C(OPh)CH_2$ (162°) on heating

The above compounds may be viewed as derivatives of acetoacetic acid (*q* v)

β Oxy-isocrotonic acid This acid is not known in free state, for, like the preceding isomeride, it would at once change to acetoacetic acid

Methyl ether of the methyl derivative $CH_3C(OMe)CH_2CO_2Me$ (175°) S G § 1 0235 Formed from $CH_3CClCH_2CO_2Me$ and NaOMe (Enke, *A* 256, 205)

Methyl ether of the ethyl derivative $CH_3C(OEt)CH_2CO_2Me$ [12°] (195°) S G § 1 999

Ethyl ether of the ethyl derivative $CH_3C(OEt)CH_2CO_2Et$ [29°] (195° cor) Formed from β chloro isocrotonic ether and NaOEt (Koll, *A* 249, 324) Plates

Propyl derivative of the methyl ether $CH_3C(OPr)CH_2CO_2Me$ (280°) S G 961

Isobutyl derivative of the methyl ether $CH_3C(OOC,H)CH_2CO_2Me$ (258°) S G 980

α -Oxy-isocrotonic acid *Nitrile* $CH_3CHCH(OH)CN$ Formed from acrolein,

KCy, and HOAc (Lobry de Bruyn, *R. T. C.* 4, 223) Oil

Isomerides *v* OXY METHACRYLIC ACID and ACETOACETIC ACID

OXYCUMENE *v* CUMENOL

Di-oxy cumene *Di-methyl derivative* $C_6H_4(C_2H_5)(OMe)_2$ (246°) Formed by the action of sodium on an alcoholic solution of $C_6H_5(C_2H_5)(OMe)$ (268°), which is got by boiling the isomeric methyl ether of eugenol with alcoholic potash (Ciamician a Silber, *B.* 23, 1164) Colourless oil

m-Di-*o*-oxy ψ -cumene

$C_6H_5(CH_3)(CH_2OH)_2$ [4 3 1] [77°] Formed by boiling *m* di *o* bromo ψ cumene with aqueous Na_2CO_3 . V sol water and alcohol, sl sol ether (Hjelt a Gadd, *B.* 19, 867)

Isomeride *v* HYDRO- ψ -CUMOQUINONE

Tri-oxy-cumene. *Di-methyl derivative* $C_6H_5(C_2H_5)(OMe)_3(OH)$ (278°) Formed by reduction of iso-apiol (Ciamician a Silber, *B.* 23, 2286) Thick liquid, sol KOH aq.

Isomeride *v* PROPYL PYROGALLOL

OXY-CUMIDINE *v* AMIDO CUMENOL.

OXY *n*-CUMINIC ACID

$C_6H_5Pr(OH)CO_2H$ [3 2 1] [94°] Formed from *o*-propyl phenol, Na, and CO_2 (Spica, *J.* 1878, 585) —BaA', 2aq —PbA', 2aq —AgA' pp

Oxy *n* cuminic acid

$C_6H_5Pr(OH)CO_2H$ [5 2 1] [98°] Formed from *p*-propyl phenol, Na, and CO_2 (Spica) Coloured violet by $FeCl_3$ —BaA', 3aq —PbA', 2aq —AgA'

Oxy-cuminic acid $C_6H_5Pr(OH)CO_2H$ [4 2 1] [93°] *Formation* —1 By fusing carvacrol with potash (Jacobsen, *B.* 11, 1063) —2 By fusing isocymene sulphonic acid with potash (Jacobsen, *B.* 12, 432) —3 From amido cuminic acid by the diazo-reaction (Widman, *B.* 19, 270)

Properties —Flat needles or leaflets. Converted by HCl at 190° into *m* propyl phenol [26°] (228° i V) $FeCl_3$ gives a reddish violet colour. Yields CaA', and BaA'.

Oxy cuminic acid $C_6H_5Pr(OH)CO_2H$ [4 3 1] [141°] *Formation* —1 From amido cuminic acid (Cahours, *A. Ch.* [3] 53, 338, Lippmann a Lange, *B.* 13, 1662) —2 Among the products got from thymol by potash fusion (Barth, *B.* 11, 1571) —3 By oxidation of potassium cumyl-sulphate $C_6H_5MePr O SO_3K$ [1 4 3] or cumyl-phosphate $C_6H_5MePr O PO_3K_2$ [1 4 3] with alkaline $KMnO_4$, and saponification of the resulting sulphate or phosphate (Heymann a Königs, *B.* 19, 3306)

Properties —Slender needles, sl sol cold water. Not attacked by conc HCl aq at 200°

Salts —NaA' 2aq — $Na_2C_6H_5O_3$ 1aq —BaA' 2aq —CdA' 2aq —AgA' white pp

Ethyl ether EtA' [75°] Prisms

Ethyl derivative $C_6H_5Pr(OEt)CO_2H$ [140°] Crystalline powder —CdA' —AgA'

Oxy *o* cuminic acid $C_6H_5Pr(OH)CO_2H$ [3 2 1] [72°] Formed, together with $C_6H_5Pr(OH)(CO_2H)_2$ [295°], from isopropyl-phenol, Na, and CO_2 (Filet, *G.* 16, 126) Colourless needles $FeCl_3$ colours it violet

Oxy-*o*-cuminic acid $C_6H_5Pr(OH)CO_2H$ [5 2 1]. [121°] Obtained from *p*-isopropyl-phenol, sodium, and CO_2 (Paterno, *J.* 1878, 806) Formed also by fusing *m*-isocymenol with KOH (Jesurun, *B.* 19, 1415) Flat needles $FeCl_3$ colours its solution deep bluish-violet. —CaA' 2 —BaA' 2 —AgA'.

An isomeride [166°–170°] is got by fusing isocymene sulphonic acid with potash (Jacobsen, *B.* 12, 433)

Oxy ψ -cuminic acid

$C_6HMe_3(OH)CO_2H$ [6 5 3 2 1] [181°] Formed from sodium ψ cumenol and CO_2 (Krohn, *B.* 21, 884) Needles, v sol ether

Oxy ψ -cuminic acid

$C_6HMe_3(OH)CO_2H$ [1 3 4 5 6] [148°] Formed by fusing durenol with potash (Jacobsen a Schnapauff, *B.* 18, 2844) Small needles, nearly insol cold water HCl aq at 200° yields ψ cumenol [93°] —CaA' 2aq prisms, sl sol cold Aq

Di-oxy ψ cuminic acid

$C_6HMe_3(OH)_2CO_2H$ [6 4 3 5 2 1] [210°] Formed by reducing ψ cumoquinone carboxylic acid with zinc dust and aqueous NaOH (Nef, *B.* 18, 3498, A 237, 13, C J 53, 428) Needles, m sol hot water. Reduces ammoniacal $AgNO_3$. Its alkaline solution turns violet in air

Ethyl ether EtA' [109°] Needles

OXY- ψ CUMINIC ALDEHYDE $C_{10}H_{12}O_2$ *ie* $C_6HMe_3(OH)CHO$ [6 5 3 2 1] [106°] A product of the action of chloroform and NaOH aq upon ψ cumenol (Auwers, *B.* 17, 2976) Needles

DI-OXY DI ψ CUMYL $C_{18}H_{20}O_2$ *ie*

$C_6HMe_3(OH)C_6HMe_3(OH)$ [173°] Formed in the preparation of ψ cumenol from ψ cumidine by the diazo reaction, and also by oxidation of ψ -cumenol with dilute HNO_3 (Auwers, *B.* 17, 2982, 18, 2659) Crystals. Yields a di bromo derivative [187°] and a di methyl derivative [126°]

a OXY CUMYL-ACETIC ACID $C_{11}H_{14}O_3$ *ie*

$C_6H_5PrCH(OH)CO_2H$ [158°] Formed from cuminic aldehyde, HCy, and HCl (Raab, *B.* 8, 1148) Needles —BaA' 4aq —PbA' 2

Oxy di cumyl acetic acid

$(C_6H_5C_2H_5)_2C(OH)CO_2H$ [120°] Formed from $C_6H_5C_2H_5CO CO C_6H_5C_2H_5$, by potash-fusion (Boesler, *B.* 14, 326) Needles —BaA'

OXY-CUMYL-ACRYLIC ACID $C_{11}H_{14}O_3$ *ie* $C_6H_5(C_2H_5)(OH)CHCHCO_2H$ The (4, 2, 1) acid [176°] and the (4, 3, 1) isomeride [206°] are formed by the diazo reaction from the corresponding amido-acids. Both are crystalline (Widman, *B.* 19, 268, 417)

DI OXY DI CUMYL PYRAZINE DICARB OXYLIC ACID

[1 4 3] $CO_2H.C_6H_5Pr.N < \begin{smallmatrix} C(OH) & OH \\ CH & C(OH) \end{smallmatrix} > N.C_6H_5Pr.CO_2H$ [3 4 1]

Formed by boiling $CH_2Cl.CO.NH.C_6H_5Pr.CO.Me$ with alcoholic potash (Abenius, *J. pr.* [2] 40, 440) Insoluble powder —EtA' [193°]

OXYCYANONINE *v* CYANONINE

OXYCYANETHINE *v* CYANETHINE

OXY CYCLOPIN $C_{11}H_{16}O_{11}$ A glucoside in Cape Tea split up by dilute acids into oxy cyclopico acid $C_{11}H_{16}O_{12}$ and glucose (Greenish, *Ph.* [3] 11, 569)

DI-OXY DIGUMYL $C_{26}H_{30}O_2$ *ie*

$C_6H_5Me(C_2H_5)(OH)C_6H_5Me(C_2H_5)(OH)$ [165 5°] Formed by oxidising thymol with iron alum (Dianin, *J. R.* 14, 136) Prisms or tables (containing aq), v sol alcohol

OXY CUMYL-ACRYLIC ACID $C_{11}H_{14}O_3$ *ie* [2 5 6 1] $C_6H_5Me(C_2H_5)(OH)CHCHCO_2H$ The anhydride [53°] [220°–230°] is formed by heating thymol with malic acid and H_2SO_4 (Fechmann a. Welsh, *B.* 17, 1647)

The isomeride (2, 5, 4, 1)-acid 280° is formed from thymolic aldehyde, NaOAc, and Ac_2O

(Kotek, *B* 16, 2104) It yields a crystalline methyl derivative [141°]

DI OXY-DI CYMYL-ETHANE $C_{22}H_{40}O_2$, *s.e.* $CH_3CH(C_6H_5, OH)_2$ [185°] Formed from thymol, paraldehyde, chloroform, and $SnCl_4$ (Steiner, *B* 11, 287) Needles (from benzene). Yields $C_6H_5(OEt)_2$ and $C_6H_5(OAc)_2$ [100°]

DI OXY-DI CYMYL-ETHYLENE $CH_2C(C_6H_5, OH)_2$ [171°] Formed, together with the preceding body, from $CCl_4CH(C_6H_5, OH)_2$ by boiling with alcohol and zinc-dust (Jaeger, *B* 7, 1198, *C* *J* 31, 263) Oxidised by alkaline K_2FeO_4 to $C_6H_5O_2$ [215°] and $C_6H_5O_2$ [215°].

DI OXY DI CYMYL SULPHIDE $(C_6H_5MePr(OH))_2S$ [153°] (Tassinari, *G* 17, 83)

DI OXY DECANE $C_{10}H_{20}O_2$ *Decylene glycol* *Diamylene glycol* The diacetyl derivative, formed from di bromo decane and $AgOAc$, yields on saponification with KOH the anhydride $C_{10}H_{18}O$ (diamylene oxide) (Bauer, *J* 1862, 450)

Di-oxo-decane $CMePr(OH)CMePr(OH)$ *Methyl propyl pinacone* (c 223°) Formed from methyl *n* propyl ketone, water, and Na (Friedel, *J* 1869, 513) Dilute H_2SO_4 converts it into the pinacol $C_{10}H_{20}O$ (185° uncor) (Szymanski, *B* 19, 1532) The isomeric $CEt_2(OH)CEt_2(OH)$ [28°] is formed in like manner from di ethyl-ketone (Schramm, *B* 16, 1594)

Tetra oxy-decane Anhydride $\langle CMe(OH)CH_2CMe \rangle O$ (270°) Formed from $\langle CMe(OH)CH_2CMe \rangle O$ (270°) Formed from $(CH_3CO)_2CH_2$ and sodium amalgam in presence of a dilute acid (Combes, *A Ch* [6] 12, 230) Liquid, v sl sol water

γ OXY DECOIC ACID *Lactone* $C_{10}H_{18}O_2$, *s.e.* $C_6H_5CH \langle \begin{smallmatrix} CH_2CH_2 \\ OCO \end{smallmatrix} \rangle$ (281° uncor) Formed from bromo decanoic acid and Na_2CO_3 (Schnee gans, *A* 227, 92) Yields $Ba(C_{10}H_{17}O_2)_2$ aq and $AgC_{10}H_{17}O_2$

Oxy decanoic acid $CH_3PrCH(OH)CHPrCO_2H$ [120°] Formed from isovaleric ether and Na (Wohlbruck, *B* 20, 2332) Needles, v sol hot water— BaA'_2 — AgA'_2 $AgOH$ amorphous.

Oxy decanoic acid $C_{10}H_{18}O_2$, *s.e.* $C_6H_5CH(OH)CH(C_6H_5)CO_2H$ [120°] Formed by the action of alcoholic potash at 110° on $C_6H_5CH(OEt)CH(C_6H_5)CO_2Et$, which constitutes half the product of the action of sodium on isovaleric ether (Hantzsch, *A* 249, 64)

DI OXY-DECYLENE *Di-valeryl-derivative* $C_6H_5C(O)COC_6H_5$ $C(O)COC_6H_5$ C_6H_5 (155°–165° at 12 mm) (Klinger & Schmutz, *B* 24, 1275)

DI-OXY-DODECANEC $CEtPr(OH)CEtPr(OH)$ (255°) Formed from ethyl propyl ketone, water, and Na (Oechsner de Coninck, *Bl* [2] 25, 10) Liquid The isomeric 'methyl butyl pinacones' $CHMeEtCMe(OH)CMe(OH)CHMeEt$ (249°) and $CMe, CMe(OH)CMe(OH)CMe$ [69°] are prepared in like manner (Wishoenus, *A* 219, 810, Friedel & Silva, *J* 1873, 340)

TRI-OXY-DODECANEOIC ANHYDRIDE $C_{12}H_{22}O_5$, *s.e.* $\begin{smallmatrix} CH_2CH_2 \\ CHEtO \end{smallmatrix} \rangle C \langle \begin{smallmatrix} CH_2CHEt \\ COO \end{smallmatrix} \rangle$ *Di-hexolactone* (above 800°) Formed from oxy-hexanoic lactone and $NaOEt$ (Fittig, *A* 256, 135). Liquid, sl sol water

DI-OXY-DODECINENE $C_{12}H_{20}O_2$, *s.e.* $C_6H_5CH_2CMe(OH)CMe(OH)CH_2C_6H_5$ (266°).

SG § 963 *CE* (0°–24°) 00082 Formed from allyl acetate (Kablukoff, *J R* 1887, 518)

OXY-DODECOIC ACID $(C_6H_5)_2C(OH)CO_2H$ *Diamoxalic acid* [122°] Formed from oxalic ether, isomyl iodide, and zinc (Frankland & Duppa, *A* 142, 8) Yields EtA' (262°) and C_6H_5A' (280°–290°)

OXY-DURYL-ACETIC ACID [64321] $C_6HMe, CH(OH)CO_2H$ [156°] Formed from tetra methyl-phenyl glyoxylic acid, alcohol, and sodium-amalgam (Claus & Foecking, *B* 20, 3100, *J pr* [2] 33, 232) Prisms— NaA'_2 aq— CaA'_2 8aq— BaA'_2 8aq v sol water The isomeric (5, 4, 3, 2, 1)-acid [160° uncor] yields KA'_2 4aq, BaA'_2 3aq, and CaA'_2 2aq

DI-OXY-ENNANE $C_9H_{18}O_2$, *s.e.* $PrCH(OH)CH(OH)CH_2Pr$ [80°] (232°) Formed by the action of alcoholic potash on a mixture of isobutyric aldehyde and valeric aldehyde (Fossek, *M* 5, 120, Swoboda, *M* 11, 384) Prisms (from water) On boiling with dilute H_2SO_4 it yields a pinacol $C_9H_{18}O_2$ (274°) Cold conc H_2SO_4 forms $C_9H_{16}O$ (150°)

Di acetyl derivative $C_9H_{16}O_4$ (242°) **OXY-ENNENOIC ACID** $C_9H_{16}O_3$ *Oxy vinyl-iso heptonic acid* Found among the products got by passing CO over $NaOAc$ mixed with sodic iso amylate at 180° (Poetsch, *A* 213, 78) Thick honey yellow oil, not volatile with steam— $C_6H_5NaO_2$ 8aq

Methyl ether $C_9H_{16}MeO_2$ (c 250°) **OXY-ENNOIC ACID** $CP_2(OH)CH_2CO_2H$ Formed by oxidising $CP_2(OH)C_6H_5$ with $KMnO_4$ (Schirokoff, *J pr* [2] 23, 197) Syrup, sl sol hot water— CaA'_2 aq— BaA'_2 aq S (of BaA'_2) 93 at 20°— PbA'_2 S 16 at 195°— AgA' prisms

Oxy ennoic acid $PrCH_2CH_2CEt(OH)CO_2H$ Crystalline— BaA'_2 — AgA'

Ethyl ether EtA' (225°) *SG* 12 940 A product of the action of isomyl iodide and zinc on oxalic ether (Frankland & Duppa, *A* 142, 6, Beilstein, *Bn* 1, 529)

OXY ERUCIC ACID $C_{22}H_{42}O_2$ Formed from di bromo behenic acid $C_{22}H_{42}Br_2O_2$ and moist Ag_2O (Haussknecht, *A* 143, 51) Oil, forming amorphous salts

OXY ETHANE PHOSPHONIC ACID $C_2H_5PO_2$, *s.e.* $CH_3CH(OH)PO(OH)_2$ [76°] Formed from aldehyde (4 mols) and PCl_3 (1 mol), followed by cold water (Fossek, *M* 7, 32) Crystals decomposing at 120°— CaA'

OXY ETHANE SULPHONIC ACID *s.e.* *ETHIONIC ACID*

Oxy ethane disulphonic acid $C_2H_5(OH)(SO_3H)_2$ Formed, together with ethane tri-sulphonic acid, by boiling tri-bromo ethane with a saturated solution of ammonium sulphite (Monari, *B* 18, 1347)— Na_2A' 3½aq— $(NH_4)_2A'$ ½aq— BaA' 2aq very soluble powder

Oxy-ethane disulphonic acid $SO_3HCH_2CH(OH)SO_3H$ Formed from is ethionic acid and H_2SO_4 at 100° (Meves, *A* 143, 196, Engelhardt & Latschnoff, *Z* 1868, 271)— KA' ½aq needles, v sol water

OXY-ETHENYL-AMIDO-PHENYL-MERCAPTAN C_6H_5NSO , *s.e.* $C_6H_5 \langle \begin{smallmatrix} N \\ S \end{smallmatrix} \rangle CH_2OH$ [175°] Prepared by heating chloro-acetic acid with amido-phenyl-mercaptan (Hofmann, *B* 18,

1284) Long fine needles Insol water, sol. alcohol and caustic alkalis

DI-OXY-ETHENYL-*o*-PHENYLENE DIAMINE $C_8H_8N_2O_2$ [above 280°] Obtained by reduction of *o*-nitro oxaniline acid with SnCl₂ (Aschan, B. 18, 2939) Sublimes in plates Sol acetic acid, sl sol water, alcohol, and ether, insol benzene, ligroin, and chloroform It has weak acid properties, forming unstable salts —BaA₂ 2aq white crystalline pp

OXY-ETHYL-ACETOACETIC ETHER

$CH_3COCH(CO_2H)CH_2CH_2OEt$ Formed from acetoacetic ether, NaOEt, and glycol chlorhydrin (Chanlaroff, A. 226, 326) Liquid, yielding γ -oxy butyric acid on boiling with baryta and alcohol

Oxy-di-ethyl-acetoacetic ether *Methyl derivative* $CH_3(OMe)COCEt_2CO_2Et$ (c 188°) Formed, together with an oil $C_8H_{11}O_2$ (c 131°) by the action of NaOMe on chloro di-ethyl-acetoacetic ether (James, A. 231, 240)

Di-oxy-di-ethyl-acetoacetic ether *Di-methyl derivative* $CH(OMe)_2COCEt_2CO_2Et$ (c 195°) Formed together with the compound $CH(OMe)_2COCHEt_2$ (184°), by the action of NaOMe on $CHCl_2COCEt_2CO_2Et$

OXY-ETHYL DIALLYLAMINE

$(C_2H_5)_2NCH_2CH_2OH$ (197°) Formed from di-allylamine and CH_2ClCH_2OH (Ladenburg, B. 14, 1879) Liquid base

OXY-ETHYL-*p*-AMIDO-BENZOIC ACID $CH_2OHCH_2NH_2C_6H_4CO_2H$ (187°) Formed from *p*-amido-benzoic acid and ethylene oxide (Ladenburg, B. 6, 129) Prisms, sl sol cold water —HA/HNO₃ crystals

OXY-ETHYL-*o*-AMIDO-PHENOL

$CH_2(OH)CH_2NH_2C_6H_3OH$ (290°–310°) Formed from *o*-amido phenol and glycol chlorhydrin (Knorr, B. 22, 2095) Liquid, sol water Successive treatment with fuming HCl and NaOHAq forms the anhydride $C_8H_7O_2<NHCH_2$ (268°)

Methyl derivative

$CH_2(OH)CH_2NH_2C_6H_3OMe$ (305° i V) Formed from *o*-anisidine

OXY-ETHYL-AMIDO-QUINOLINE DIHYDRIDE *v* **HYDRAZIDO-PHENYL-PROPIONIC ACID**

OXY-ETHYL-AMINE $CH_2(OH)CH_2NH_2$ *Amido ethyl alcohol* Formed by the action of ammonia on ethylene oxide or on glycol chlorhydrin (Wurtz, C. R. 49, 898, 53, 338) Formed also by the action of conc H_2SO_4 on bromo-ethyl phthalimide, by the action of HNO₃ on vinylamine, and by the action of AgNO₃ (2 mols) on bromo ethylamine hydrobromide (1 mol) (Gabriel, B. 21, 569, 2666)

Salts —B'HCl [below 100°] —B'₂H₂PtCl₂ —B'HBz hygroscopic crystals —B'HNO₃ [55°] —B'₂C₆H₄(NO₂)₂OH [159°] Yellow needles (from alcohol)

Picrate of the acetyl derivative $CH_2(OH)CH_2NHAcC_6H_4(NO_2)_2OH$ [169°] (Gabriel, B. 22, 2222)

Benzoyl derivative $CH_2(OH)CH_2NHBz$ —B'HBz [148°] —B'₂H₂PtCl₂ —Picrate. [195°] Crystalline pp

Phenyl derivative $CH_2(OPh)CH_2NH_2$ (229°). Formed from phenyl oxyethyl phthalamic acid and conc HClAq (Schmidt, B. 22, 8256). Yields an acetyl derivative [78°] and a

benzoyl derivative [98°] (Schreiber, B. 24, 189). —B'HCl [215°] —B'₂C₆H₄(NO₂)₂OH —Platinichloride B'₂H₂PtCl₂ golden needles

p-Tolyl derivative (243°) Yields a benzoyl derivative [134°] and B'HC [240°], B'₂H₂PtCl₂, and B'₂C₆H₄N₂O

Oxy-tri-ethyl-amine $C_6H_5NO_3$ Formed from $NEt_3CH_2CH_2OH$ (161°) NO from NEt_3H and glycol chlorhydrin (Ladenburg, B. 14, 1878, 15, 1144) Liquid, miscible with water

Cinnamoyl derivative

$NEt_3CH_2CH_2OOCOC_6H_5$ Formed from the cinnamate of the base and dilute HCl Yields B'HAuCl₄ and B'₂C₆H₄(NO₂)₂OH

Ethyl-chloride $NEt_3ClCH_2CH_2OH$ Prisms (Wurtz, A. Suppl. 7, 88) —Gold salt $C_6H_5NOAuCl$ golden plates

Di-oxy-ethyl-amine *Di-ethyl derivative* $CH_2(NH_2)CH(OEt)_2$ *Amido alcohol* (162°) Formed from chloro acetal and alcoholic NH₃ (Wohl, B. 21, 616, Wolff, B. 21, 1482) Colourless liquid, v e sol water —B'₂H₂PtCl₂ hexagonal plates, sl sol cold alcohol

Di-oxy-di-ethyl-amine $NH(CH_2CH_2OH)_2$ Formed from ethylene oxide or $ClCH_2CH_2OH$ and ammonia (Wurtz) —B'₂H₂PtCl₂ tables

Tri-oxy tri-ethyl-amine $N(CH_2CH_2OH)_3$ A product of the action of ethylene oxide on ammonia (Wurtz) With glycol chlorhydrin it yields $N(OH)(CH_2CH_2OH)_2$ Both are syrupy

Tetra-oxy-di-ethyl-amine *Tetra-ethyl derivative* $NH(CH_2CH_2OEt)_2$ (260°) S 14 Formed from chloro acetal and NH₃Aq at 130° (Wolff, B. 21, 1484) —B'₂H₂PtCl₂ [121°]

***o*-OXY-ETHYL AMINE *acv* - TRI SULPHONIC ACID** $C_6H_5NS_3O_6$ *v* $CH_3C(HO)(SO_3H)CH(SO_3H)NH(SO_3H)$ The sodium salt Na_3A''' 3aq is formed by shaking nitroso acetone $CH_3COCH(NOH)$ with a 30 p c solution of sodium bisulphite, and crystallises out on standing as a crystalline powder consisting of white needles It is v sol water, insol alcohol

By warming with dilute acids it is split up into methyl-glyoxal (pyruvic aldehyde) CH_3COCHO , sodium sulphate, NH₃, and SO₂ (Pechmann, B. 20, 2543)

OXY-ETHYL-ANILINE $NHPhCH_2CH_2OH$ (286°) SG 2 1 110 Formed from aniline and ethylene oxide (Demole, B. 6, 1024, Ladenburg, B. 6, 181, Knorr, B. 22, 2092) Liquid, with strong reducing properties —B'₂H₂PtCl₂ red dish brown crystals

o **Oxy-di-ethyl-aniline** $C_6H_5NEtO_2CH_2OH$ (268°) Formed from ethyl aniline and ethylene chlorhydrin at 100° (Laun, B. 17, 677) Heavy oil

Di-oxy-di-ethyl-aniline $NPh(CH_2CH_2OH)_2$ (above 350°) Formed from the preceding and glycol chlorhydrin Treatment with HClAq at 170°, followed by cold caustic soda solution, yields $NPh<\begin{smallmatrix} C_6H_5 \\ C_6H_5 \end{smallmatrix}>O$ [58°] (270°). The compound

$NPh<\begin{smallmatrix} C_6H_5 \\ C_6H_5 \end{smallmatrix}>O_2$ [67°] is formed by the action of alcoholic AgNO₃ on the corresponding sulphur compound (Holzmann, B. 20, 1640)

OXY-ETHYL-BENZENE *v* **ETHYL PHENOL** *Di-oxy-ethyl-benzene* $C_6H_5CH_2CH_2OH$ *Styrolene alcohol* [68°] (278° i V) Formed from styrene dibromide and boiling aqueous K₂CO₃ (Zincke, A. 216, 298). Needles (from

benzene ligroin) Converted by boiling dilute H_2SO_4 into oily $\text{C}_8\text{H}_{10}\text{O}$ (260° at 50 mm)

Di-acetyl derivative $\text{C}_8\text{H}_8\text{Ac}_2\text{O}_2$ (274° i.v.)

Di benzoyl derivative [97°] Needles *Di-oxy-ethyl benzene* $\text{C}_8\text{H}_8\text{Et}(\text{OH})_2$ [134] (295°) Formed from *p* ethyl phenol sulphonic acid by potash fusion (Sempotowski, *B* 22, 2667) Coloured green by FeCl_3

Di-oxy-tri ethyl-benzene $\text{C}_8\text{H}_6\text{Et}_3(\text{OH})_3$ [185°] Formed by the action of HCl on $\text{C}_8\text{H}_6\text{Et}_3(\text{OEt})(\text{OH})$ (165° uncor at 20 mm) which is a product of the action of alcoholic potash and EtI on resorcin (Herzig & Zeisel, *M* 11, 306) White needles (from dilute alcohol) Yields $\text{C}_8\text{H}_6\text{Et}_3(\text{OEt})(\text{OAc})$ [65°] crystallising in monoclinic prisms

OXY-ETHYL BENZOIC ACID $\text{C}_8\text{H}_8\text{O}_2$ *s.s.* $\text{C}_8\text{H}_7\text{Et}(\text{OH})(\text{CO}_2\text{H})$ [120°] Formed from sodium *o*-ethyl phenol and CO_2 (Beilstein & Kuhlberg, *A* 156, 213) Coloured violet by FeCl_3

Oxy-ethyl benzoic acid [112°] Formed in like manner from sodium phlorol (Oliveri, *G* 13, 267) Needles (from water) — BaA^1 , aq scales β *Oxy ethyl benzoic acid*

$\text{CH}_3\text{CH}(\text{OH})\text{C}_6\text{H}_4\text{CO}_2\text{H} - \text{AgA}^1$ needles *Anhydride* $\text{C}_8\text{H}_6\text{O}_2$ (276°) V D 74 (obs) Formed by reducing acetophenone carboxylic acid with sodium amalgam and acidifying (Gabriel & Michael, *B* 10, 2205) Crystalline at 0°

Tri oxy-ethyl-benzoic acid Methylene derivative $\text{CH}_3\text{CH}(\text{O})\text{C}_6\text{H}_4(\text{CH}_2\text{CH}_2\text{OH})\text{CO}_2\text{H}$ 'α *Oxyethylperonylcarboxylic acid*' [146°] Formed from the corresponding amido-acid $\text{CH}_3\text{O}\text{C}_6\text{H}_4(\text{C}_6\text{H}_4\text{NH})\text{CO}_2\text{H}$ (Perkin, jun, *C* 57, 1021) Crystals Converted at 120°-130° into the anhydride $\text{C}_{10}\text{H}_8\text{O}_4$ [127°] — $\text{AgC}_{10}\text{H}_8\text{O}_4$, — $\text{KA}^1 - \text{NH}_4\text{MeA}^1$ crystals

TRI OXY TRI-ETHYL BORATE $\text{B}(\text{OC}_2\text{H}_4\text{OH})_3$, [162°] Formed from BCl_3 and glycol (Counciler, *J pr* [2] 18, 392) Deliquescent crystals

DI OXY ETHYL-BUTYL GLYOXALINE $\text{C}_8\text{H}_8\text{CH}(\text{CO})\text{NEt}$ [135°] (295°) Formed from isobutyl hydantoin, KOH , and EtBr (Pinner, *B* 22, 685) Silky needles

OXY-ETHYL CARBAMIC ACID Anhydride $\text{CH}_3\text{OCH}(\text{NH})\text{CO}$ [91°] Formed from β bromo ethyl-amine and Ag_2CO_3 (Gabriel, *B* 21, 568) Flat crystals (from alcohol), v sol water

OXY-ETHYL CONIINE v CONIINE

DI-OXY ETHYLENE DIQUINOLINE OCTO-HYDRIDE $\text{C}_{20}\text{H}_{18}\text{N}_4\text{O}_2$ *s.s.* $\text{OHCH} - \text{OCH}_2\text{CH}_2\text{CH}_2\text{OCH}(\text{OH})\text{O} - \text{NCH}_2\text{CH}_2\text{N} - \text{OCH}(\text{OH})\text{CH}$ [233°] Formed by heating oxy-quinoline tetrahydride with ethylene bromide (Kohn, *C* 49, 509) Small silky prisms

DI-OXY-ETHYLIDENE-DI-NAPHTHOQUINONE $\text{C}_8\text{H}_6\text{CH}(\text{CO})\text{C}(\text{OH})\text{CO} - \text{CO} > \text{C}_8\text{H}_6$ *Phenyl hydrazide*

$\{\text{C}_{10}\text{H}_7(\text{OH})\text{O}(\text{N}_2\text{HPh})\}_2\text{CHMe}$ [258°] Got by heating the phenyl hydrazide of oxy (α) naphthoquinone with alcohol and aldehyde at 100°

(Zincke & Thielen, *B* 21, 2205) Small red needles, v sl sol alcohol

TETRA OXY-TETRA ETHYLIDENE-PHOSPHONIUM CHLORIDE $(\text{CH}_3\text{C}(\text{OH}))_4\text{PCl}$ [112°] Formed from aldehyde, PH_3 , and HCl (Messenger & Engels, *B* 21, 328) Dimetric crystals, with unpleasant smell, v sol alcohol, insol ether Decomposed by water The corresponding $(\text{CH}_3\text{C}(\text{OH}))_4\text{PBr}$ [88°] is also crystalline The iodide $(\text{CH}_3\text{C}(\text{OH}))_4\text{PI}$ [65°] is formed from aldehyde and phosphonium iodide (Girard, *A Ch* [6] 2, 11)

DI-OXY-ETHYLIDENE-METAPYRAZOLE

$\text{CH}_3\text{CH}(\text{C}=\text{N})\text{C}(\text{OH})\text{CH}(\text{C}=\text{N})\text{CH}_3$ or $\text{CH}_3\text{CH}(\text{C}=\text{N})\text{CO} - \text{OC}(\text{NH})\text{CH}_3$ *Ethylidene-metapyrasolone*

Formed by heating chloro-crotonyl-urea, $\text{CH}_3\text{CH}(\text{C}=\text{N})\text{CO}\text{NH}\text{CO}\text{NH}_2$, to its melting-point, HCl being evolved Needles M sol. hot water (Pinner & Lifschutz, *B* 20, 2350)

α OXY-ETHYL MALONIC ACID $\text{CET}(\text{OH})(\text{CO}_2\text{H})_2$ *Ethyltartronic acid* [98°] Prepared by saponification of chloro-ethyl malonic ether with baryta water (Conrad, *B* 14, 618) Decomposes at 180° into α-oxy-butyric acid and CO_2

β Oxy-ethyl malonic acid $\text{CH}_3\text{CH}(\text{OH})\text{CH}(\text{CO}_2\text{H})_2$ Formed from ethylidene malonic acid and baryta water (Kommenos, *A* 218, 163) Syrup — AgA^1 amorphous

γ Oxy ethyl malonic acid $\text{CH}_3(\text{OH})\text{CH}_2\text{CH}(\text{CO}_2\text{H})_2$ — BaA^1 1½ aq Got by boiling the lactonic acid with baryta — AgA^1

Lactonic acid $\text{CH}_3\text{CH}(\text{CO}_2\text{H})\text{O} - \text{CO}$

Formed by boiling bromo ethyl malonic acid with water (Röder, *A* 227, 19) Obtained also by boiling vinaconic acid with diluted H_2SO_4 — BaA^1

DI-OXY DI-ETHYL OXIDE

$\text{O}(\text{CH}(\text{OH})\text{CH}_3)_2$ Alkoyl derivatives of this body are formed by the action of sodium salts on di chloro di ethylether (Geuther, *A* 226, 223, 245, 101)

Formyl derivatives (c. 180°) SG 11 1134

Acetyl derivative $\text{O}(\text{CH}(\text{OAc})\text{CH}_3)_2$ (192°) SG 11 1071, 22 1067 Colourless liquid, slowly decomposed by water into aldehyde and acetic acid

Propionyl derivative (c 218°) SG 22 1037

Butyryl derivative (c 238°) SG 22 994

Benzoyl derivative $\text{C}_8\text{H}_8\text{O}_2$ Needles *Succinyl derivative* $\text{O}(\text{CMeH})_2\text{C}_4\text{H}_4\text{O}_2$ *Di-methyl ether* $\text{O}(\text{CH}(\text{OMe})\text{CH}_3)_2$ (127°) SG 11 958 Formed from *aa* di-chloro-di-ethyl oxide, MeOH , and NaOMe Oil decomposing on standing into di methyl acetal and aldehyde

Di-ethyl ether (153° cor) SG 14 891

Di-propyl ether (184° cor) SG 14 895

Di-isobutyl ether (175° cor) SG 14 879

Di-isoamyl ether (227°) SG 14 874

DI-OXY-DI-ETHYL-DIPHENYL Di-ethyl ether [341] $\text{C}_6\text{H}_5\text{Et}(\text{OEt})\text{C}_6\text{H}_5\text{Et}(\text{OEt})$ [134]

[120° cor]. Formed from di amido di-ethyl-di-phenyl, alcohol, and nitrous acid (Schultz, *B* 17, 475) White plates, sl sol cold alcohol

Tetra-oxy-ethyl-diphenyl Tetra-acetyl derivative $C_{12}H_8Et(OAc)_4$ [138°] Formed from the tetra-ethyl derivative by successive treatment with $HIAg$ and Ac_2O (Herzig, *M* 11, 418) Needles (from alcohol)

Tetra-ethyl derivative $C_{12}H_8Et(OEt)_4$ [92°] A product of the action of EtI and alcoholic potash on tetra oxy diphenyl (Herzig, *M* 11, 417) Plates, v sol alcohol

OXY-ETHYL-PHTHALIMIDE $C_{10}H_8NO_4$ i.e. $C_6H_4(CO)_2NCH_2CH_2OH$ [127°] Formed from bromo-ethyl phthalimide and $KOHAc$ (Gabriel, *B* 21, 571) Needles or plates, sol hot water

Phenyl derivative $C_6H_4(CO)_2NCH_2CH_2OPh$ [180°] Formed from $PhOC_6H_4Br$ and potassium phthalimide (Schmidt, *B* 22, 3255) On warming with potash it yields the acid $C_{10}H_8NO_4$ [125°]

p-Tolyl derivative [135°] Crystals Yields a di nitro derivative [88°] (Schreiber, *B* 24, 190)

OXY ETHYL PIPERIDINE $C_7H_{13}NO$ i.e. $C_4H_8NCH_2CH_2OH$ (199°) Formed by heating piperidine with ethylene chlorhydrin (Ladenburg, *B* 14, 1876) Liquid, sol water — B^*HauCl , plates

Phenyl acetyl derivative $C_{11}H_{11}NO_2$ Oil, formed by heating phenylacetic acid with oxyethyl piperidine and HCl — B^*HauCl , [c 100°] — B^*HI — B^*HI , Brown needles

Benzoyl derivative $C_{11}H_{11}NO_2$ — B^*HI — $B^*H_2PtCl_6$, silky needles

Oxy-ethyl-piperidine $C_8H_{10}(C_2H_5OH)_2N$ [32°] (227°) Formed by reducing oxy-ethyl pyridine with sodium amalgam (Ladenburg, *B* 22, 2586) Hygroscopic mass Yields an oily nitrosamine — $B^*H_2PtCl_6$ [158°] Large crystals

OXY ETHYL PYRIDINE $C_7H_7NCH_2CH_2OH$ (179° at 25 mm) SG ± 1.111 Formed from (a) methyl pyridine and formic aldehyde (Ladenburg, *B* 22, 2584) Syrup, v sol water, not volatile with steam Yields vinyl pyridine on distillation — $B^*H_2PtCl_6$ [170°] prisms, m sol cold water

An isomeride $C_7H_7N(CH(OH)CH_3)$ is got by distilling barium a oxy pyridyl propionate (Hardy a Calmels, *Bl* [2] 48, 280)

Oxy-di-ethyl pyridine $NC_2H_5Et(C_2H_5OH)_2$ (148° at 18 mm) Formed from methyl ethyl-pyridine (aldehyde collidine) and formic aldehyde at 165° (Prausnitz, *B* 23, 2725) Oil — $B^*H_2PtCl_6$ [159°] Red cubes

OXY-ETHYL-PYRIDINE TETRAHYDRIDE $CH_3CH_2COCH_2CH_2CH_2CH_2NH$ [68°] Formed by heating β -amido heptonic acid at 220° (Aschan, *B* 23, 3694) Plates, smelling like conine

OXY-ETHYL-PYRIDINE CARBOXYLICACID $NC_2H_5Et(OH)CO_2H$ Formed from comanic acid and ethylamine (Ost, *J pr* [2] 29, 380) Prisms (containing jag), v sol water

Di oxy-ethyl-pyridine carboxylic acid $NC_2H_5Et(OH)CO_2H$ [c 210°] Formed from ethylamine and comenamic acid (Meunel, *J pr* [2] 32, 178) Prisms (containing jag) Coloured violet by $FeCl_3$. Reduces ammoniacal $AgNO_3$, Ac_2O at 160° forms acetyl ethyl pyromecanic

acid $C_7H_7NO_3$, whence boiling water forms $C_7H_7NO_3$ [166°]

Ethyl ether EtA' [115°] Needles (from water) — $EtA' \cdot HCl$ — $Ba(C_2H_5Et.NO)_2$, needles

OXY ETHYL-QUINAZOLINE DIHYDRIDE $C_8H_8 \begin{matrix} CH_2 \\ \diagup \quad \diagdown \\ NH \quad CO \end{matrix} \begin{matrix} CH_2 \\ \diagup \quad \diagdown \\ NH \quad CO \end{matrix}$ Benzylene ethyl urea [95°]

Formed from oxy-tolyl-ethyl thio-urea, alcohol, and HgO (Soderbaum a Widman, *B* 22, 2937). Needles (from alcohol) — $B^*H_2PtCl_6$, [c 205°] — B^*HauCl , golden scales

(Py 3 4) **OXY-ETHYL QUINOLINE**

$C_8H_8 \begin{matrix} CH \quad CH \\ \diagup \quad \diagdown \\ NH \quad CO \end{matrix}$ Ethyl pseudo carbostyryl

[55°] Formed together with (Py 3) ethoxy quinoline by the ethylation of carbostyryl (Py 3 oxy quinoline) (Friedlander a Weinberg, *B* 18, 1528) Glistening white needles Sl sol water, v sol other solvents Scarcely volatile with steam Dissolves in strong HCl

(Py 1, 3) Di oxy (Py 2)-ethyl quinoline Formed from chloro oxy ethyl quinoline and HCl (Rügheimer a Schramm, *B* 21, 301) Prisms, v e sol alcohol

DI OXY-ETHYL ISOQUINOLINE

$C_8H_8 \begin{matrix} CH_2 \quad CO \\ \diagup \quad \diagdown \\ CO \quad NH \end{matrix}$ [105°] Formed by distilling o carboxy phenyl acetic acid with ethyl amine (Pulvermacher, *B* 20, 2493) Needles, v sol alcohol With diazobenzene chloride it yields $C_8H_8ONEt \cdot N_2Ph$ [139°]

Di-oxy di ethyl-isoquinoline

$C_8H_8 \begin{matrix} CH_2 \quad CO \\ \diagup \quad \diagdown \\ CO \quad NH \end{matrix}$ [144°] Formed from di oxy isoquinoline (vol 1 p 706) $NaOEt$ and EtI (P) White plates With alcoholic potash and EtI it produces di-oxy tri ethyl-isoquinoline

$C_8H_8 \begin{matrix} CH_2 \quad CO \\ \diagup \quad \diagdown \\ CO \quad NH \end{matrix}$ [50°] (309°)

(Py 8, 2) **OXY ETHYL-QUINOLINE DIHY**

DRIDE $C_8H_8 \begin{matrix} CH_2 \quad CH_2 \\ \diagup \quad \diagdown \\ NH \quad CO \end{matrix}$ Ethylhydrocarbo

styryl [88°] Formed from phenyl valeric acid by nitration and reduction (Baeyer a Jackson, *B* 13, 119) Crystals Occurs also in an unstable modification [76°] PCl_5 gives chloro ethyl quinoline and $C_{11}H_{11}NO$ [168°]

(B 2) Oxy (Py 4) ethyl quinoline tetrahydryde **Methyl derivative** C_8H_7NO **Ethylthalline** (287°) Formed from thallin and EtI (Skraup, *M* 6, 779) Thick liquid — B^*HCl — B^*EtI [133°] — $(B^*EtCl) \cdot PtCl_6$, orange needles

(B 4) Oxy-(Py 4) ethyl-quinoline tetrahydryde $CH \quad CH \quad CH_2 \quad CH_2$ [76°] Prepared by heating (B 4) oxy quinoline tetrahydryde with EtI or $EtBr$ (O Fischer, *B* 16, 717, Fischer a Renouf, *B* 17, 756, Fischer a Kohn, *B* 19, 1044, *O J* 49, 608) Monoclinic prisms or tables, sol alcohol and ether, sl sol water $FeCl_3$ colours its alcoholic solution brown

— B^*HCl **Kaurne A** Trimetric prisms, $a, b, c = 595.1, 957$, v sol water Febrifuge — B^*EtI [160°] Prisms (from alcohol)

Ethyl ether C_8H_7NO [88°] (270°). Formed from the ethyl ether of (B 4) oxy quinoline tetrahydryde and $EtBr$ at 180° Silky plates, insol water — B^*EtI . [187°] Prisms — $B^*Et \cdot PtCl_6$, [188°]

Acetyl derivative $C_{11}H_{11}AcNO$ [64°].

(B. 1) Oxy (*Py* 4) ethyl-quinoline tetrahy-
 dride $\begin{array}{c} \text{CH C(OH) C CH}_2\text{CH}_2 \\ \text{CH CH—C NEtCH}_2 \end{array}$. [73°] Formed
 by ethylating the corresponding oxy-quinoline
 tetrahyrdr (Riemerschmid, *B* 16, 724) Crys-
 tals, sol alcohol—B/HCl aq

(Py 3) OXY-(Py 2) ETHYL-QUINOLINE-
DIHYDRIDE (Py 2) CARBOXYLIC ETHER
CH₃ CH₂ CO₂ E⁺

$$(\text{?})\text{C}_6\text{H}_4 \begin{cases} \text{CH}_2\text{CEtCO}_2\text{Et} \\ | \\ \text{N}-\text{C}(\text{OH}) \end{cases} \cdot \quad [114^\circ] \quad \text{Obtained}$$

by reduction of *o* nitro benzyl ethyl-malonic ether with zinc and acetic acid (Lellmann & Schleich, *B* 26, 440) Colourless silky needles

OXY-ETHYL SUCCINIC ACID *C₆H₁₀O₈
—BaA''3aq —CaA''3aq —Ag₂A'' *Anhydride*

$$\text{C}_6\text{H}_5\text{O}_4 \text{ i.e. } \text{CHMe} \left\langle \begin{array}{c} \text{O CO CH}_2 \\ \text{CH}(\text{CO}_2\text{H}) \end{array} \right\rangle \quad \text{Methyl}$$

paraconic acid [84°] Formed from aldehyde, succinic acid, and Ac_2O (Fittig & Frankel, A 255, 17) Small plates $-\text{BaA}'$, $3\frac{1}{2}\text{aq}$ $-\text{CaA}'$, $2\frac{1}{2}\text{aq}$ $-\text{AgA}'$

Reference — TRI CHLORO OXY-ETHYL-SUCCINIC ACID

DI OXY DI ETHYL SULPHIDE

$\text{S}(\text{CH}_2\text{CH}_2\text{OH})_2$. Formed from ethylene chlorhydrin and aqueous K_2S (V Meyer, *B* 19, 3259)
Syrup With PCl_5 it yields $\text{S}(\text{C}_2\text{H}_4\text{Cl})_2$, whence K_2S yields insoluble $\text{C}_2\text{H}_4\text{S}_2$. A soluble $\text{C}_2\text{H}_4\text{S}_2$ is formed from $\text{C}_2\text{H}_4(\text{SNa})_2$ and ethylene bromide

OXY ETHYL *p* TOLUENE C_8H_9NO *re*
 $C_8H_7NHCH_2CH_2OH$ [37°] (287°) Formed
 from *p*-toluidine and ethylene oxide (Demole,
B 7, 635, *A* 173, 123) Crystals, sol
 water — $B'H_2PtCl_4$ [148°] — $B'H_2SO_4$ [111°] —
 $B'H_2C_2O_4$ [122°]

Di-oxy di-ethyl *p* toluidine $C_7H_7N(C_2H_4OH)_2$
(339°) —B' H_2PtCl_4 .

OXY-ETHYL UREA *p*-Tolyl-derivative
 $\text{NH}_2\text{CO NH C}_6\text{H}_4\text{OC}_2\text{H}_5$, [158°] Formed from
 tolyloxyethylamine hydrochloride and potassium
 cyanate solution at 100° (Schreiber, *B* 24, 193)

OXY-FLAVOLINE v **FLAVENOL**

OXY-FLUORENE CARBOXYLIC ACID

$\text{C}_6\text{H}_4\text{CO}_2\text{H} \xrightarrow{\text{C}_6\text{H}_5(\text{CO}_2\text{H})} \text{CH}(\text{OH})$ [203°] Formed by reducing diphenylene ketone carboxylic acid with zinc dust and ammonia (Graebe, *A* 247, 283). Crystals (from hot water or benzene).

OXYGEN O (Older names were *vital air*, *pure air*, *dephlogisticated air*) At w 15.96 (v *infra*) Mol w 31.92 Boils -181.4° at 740 mm (Olszewski, *W* 31, 58) Has not been solidified (v *infra*) SG (liquid, compared with water at 4°) 899 at -130° (Wroblewski, *W* 20, 860), 7555 at -129.6° , 8788 at -139.3° , 8544 at -137.5° , 8772 at -139.4° , 1124 $^\circ$ at -181.4° [=b p] (Olszewski, *M* 5, 124, *W* 31, 58), Wroblewski (*Lc*) gives SG for temperature from -118° to -200° as $=1.212 + 0.0428T - 0.000529T^2$; where T = absolute temperature SG (gaseous, at 3000 atmos. referred to water $=1$) 1.1054 (Amagat, *C R* 107, 522) Critical temp c -113° , critical pressure c 50 atmos VD 15.96 (v *infra*) S.H.p (equal weight of water = 1) 21751 from 18° to 207° (Regnault, *Acad* 26, 1), S.H.v (equal wt of water = 1) 1551 (Clausius, *Mechan. Wärmetheorie*, 1, 62) CE (21° to 98° at 760 mm) 0086748 (Jolly, *P. Jubelbd* 82) S 04399 at 0° . 04286 at 0° . 03802 at 10° . 03415 at

15°, -0.3103 at 20°, 0.2616 at 30° (Winkler, B. 22, 1764, S is given for each degree from 0° to 30°, and the results are compared with those of Bunsen) S (alcohol, 0° to 24°) 23897 (Carus, A 94, 184, cf Timofejeff, Z P C 6, 141) Compressibility coefficient = 0.0025 at 1000–1500 atmos, 0.0016 at 1500–2000 atmos, 0.00115 at 2000–2500 atmos, 0.00091 at 2500–3000 atmos (Amagat, C R 107, 522) O exhibits several very different emission spectra under different conditions, for measurements of lines v B A 1884 432 The absorption spectrum of O at pressures up to 90 atmos has been examined by Laveing Dewar (P M [5] 26, 286, Pr 46, 222), for the absorption spectrum of liquid O v Olzsewski, W A B 95 [2nd part], 257 (cf L a D Pr 46, 422, Janssen, C R 101, 649, 102, 1332, Egoroff, C R 101, 1143)

Occurrence.—Uncombined in the atmosphere, of which it forms one fifth by volume. Compounds of O are very numerous, and occur in enormous quantities. O forms eight-ninths of water by weight, and about one half of silica, alumina, and chalk, which are the three most plentiful constituents of the solid earth, almost every widely distributed rock or mineral contains O, the principal exceptions being rock salt, fluorspar, blende, galena, and pyrites. As O is slightly sol in water, this gas is found uncombined in all natural waters. O is a constituent of all living tissues, according to Wurster (*B* 21, 1525), O_2H_2 occurs in the juices of plants. According to H. Draper, O occurs in the sun (*Am S* [3] 14, 89, cf J O Draper, *ibid* 16, 256).

O was prepared for the first time by Priestley, in 1774, by heating HgO , a year later the gas was discovered by Scheele. Lavoisier was the first to recognise the importance of the discovery of O, and to study its chemical properties.

References to older memoirs on Oxygen — Priestley, 'Experiments and Observations on different kinds of Airs' [London, 1775-1777], 2, 29, 3, 1, Priestley, 'Experiments and Observations relating to various branches of Natural Philosophy' [London, 1779], 1, 192, Scheele, 'Abhandlungen von der Luft und dem Feuer' [Upsala & Leipzig, 1777], also *Crells Annal* 1785, 2, 229, 291, Lavoisier's memoirs appeared in *Acad* from 1774 to 1788, Berzelius, *Lehrbuch der Chemie* [3rd ed 1838, 1, 5. 46

Formation—The processes wherein O is formed may be divided into groups I *From air*—1 BaO is heated in air to dull redness, when BaO₂ is formed, the pressure is decreased, and the BaO₂ is heated, when O is evolved, and BaO remains—2 Hg is heated in air at c 300° when HgO is slowly formed, on strongly heating HgO, O and Hg are produced—3 MnO₂ is heated with NaOH in air, Na₂MnO₄ and H₂O are produced ($4\text{NaOH} + 2\text{MnO}_2 + \text{O}_2 = 2\text{Na}_2\text{MnO}_4 + 2\text{H}_2\text{O}$), the product is heated to dull redness in steam, when NaOH, Mn₂O₃, and O are formed ($2\text{Na}_2\text{MnO}_4 + 2\text{H}_2\text{O} = 4\text{NaOH} + \text{Mn}_2\text{O}_3 + 3\text{O}$), by again passing air over the residue, Na₂MnO₄ is reformed (Tessie du Motay, *D P J* 186, 230)—4 When air is pressed into water, more O than N is dissolved, by reducing pressure on the water O and N are evolved, by pressing the evolved gases again

into water, again reducing pressure, and pressing the gases into a fresh quantity of water, and repeating these processes eight times, nearly pure O (about 2.3 p c N) is obtained (Mallet, *D P J.* 199, 112). For details of the methods for obtaining O from air *see* DICTIONARY OF APPLIED CHEMISTRY II *From Oxides*—5 Several metallic oxides give off O when heated, *eg* HgO , MnO_2 , PbO_2 , Au_2O_3 , Bi_2O_3 .—6 When water is electrolysed, O separates at the positive electrode.—7 O is obtained by passing a mixture of H_2O and Cl through a red hot tube, the issuing gas is passed through NaOH to absorb HCl and excess of Cl .—8 When steam is passed over CuCl_2 at $100^\circ\text{--}200^\circ$ an oxychloride of Cu is formed which is reduced to CuCl_2 at $c\ 400^\circ$ with evolution of O (Vogel, *W J* 1861 177, Mallet, *C R* 64, 226, 66, 349).—9 O is obtained by the reaction of $\text{H}_2\text{O}_2\text{Aq}$ with Ag_2O ($\text{Ag}_2\text{O} + \text{H}_2\text{O}_2 = \text{H}_2\text{O} + 2\text{Ag} + \text{O}_2$).—10 MnO_2 , PbO_2 , CrO_3 , BaO_2 , Bi_2O_3 , and other metallic peroxides yield sulphates, H_2O , and O when heated with conc H_2SO_4 .—11 CO_2 is decomposed by fresh parsley, mint, and other green plants, when the plant is placed in water charged with CO_2 and exposed to direct sun light III. *From compounds other than oxides*.—12 KClO_3 is heated, either alone or mixed with MnO_2 or Fe_2O_3 ,—13 CaCl_2O_2 is heated, when CaCl_2 and O are formed, or dry Cl is passed over CaO heated to redness.—14 Conc HNO_3 is dropped into a red hot Pt flask, $2\text{HNO}_3 = \text{H}_2\text{O} + \text{O} + 2\text{NO}_2$.—15 Conc H_2SO_4 is dropped on to red-hot Pt, or on to red-hot bricks, the gases produced are passed through milk of lime, $\text{H}_2\text{SO}_4 = \text{H}_2\text{O} + \text{SO}_2 + \text{O}$ (Deville a Debray, *C R* 51, 822).—16 By strongly heating dry ZnSO_4 , O and SO_2 are evolved and ZnO remains (D a D, *lc*).—17 $\text{Ba}(\text{NO}_3)_2$ yields O, mixed with N, when strongly heated.—18 A mixture of O and N, containing about 60 p c O is obtained by heating NaNO_3 with two parts dry ZnO (*v* Pepper, *C N* 6, 218).—19 Conc H_2SO_4 is heated with $\text{K}_2\text{Cr}_2\text{O}_7$ or KMnO_4 , sulphates of K and Cr (or Mn) and H_2O are formed and O is evolved.—20 A fairly conc solution of bleaching powder is heated with addition of a small quantity of a salt of Co, Co_2O_3 is formed (by the action of the CaO on the Co salt), and this is probably oxidised to CoO_2 , which is again reduced to Co_2O_3 with evolution of O (Winkler, *J pr* 98, 340, Fleitmann, *A* 134, 64, Reinsch, *Z* [2] 2, 31, Böttger, *J pr* 95, 309, 375, Stolba, *J pr* 97, 309). A modification of this process consists in passing Cl into warm thick milk of lime containing a little Co_2NO_2 (W, *lc*).

Preparation.—1 Pure KClO_3 , which has been fused, is well mixed with c half its weight of pure dry MnO_2 , and the mixture is heated in a flask of hard glass to $c\ 210^\circ\text{--}220^\circ$. As the O thus obtained contains a little Cl , the gas is passed through NaOH or milk of lime. It is then dried by H_2SO_4 , and passed in a rapid stream through a glass tube heated to redness. After a time the stream of O may be allowed to slacken, and the glass tube need not be kept hot. The rapid stream of O, passing through the hot tube, washes out the last trace of air, which very obstinately adheres to glass at ordinary temperatures (Houzeau, *C R* 70, 89). Any

ozone present is also decomposed. The small quantity of air which adheres to the glass may also be removed, according to Illosva (*Bl* [3] 2, 734), by passing the O over Pt foil at $280^\circ\text{--}250^\circ$, or Pt black at $c\ 250^\circ$, whereby combination of O with the N present occurs, with formation of nitrites and nitrates, which may be absorbed in NaOH Aq. The Pt loses the power of inducing the combination of N and O after a time, but regains this power when heated in a stream of H at $c\ 250^\circ$ (*I, lc*). If the O is required quite dry, it is passed through conc boiled H_2SO_4 , and then over a long layer of P_2O_5 . Regarding the part played by MnO_2 and other substances in causing O to be evolved from KClO_3 at temperatures lower than that at which this salt gives off O when heated alone, *v* Hornsby, *Ph* 15, 352, Witt, *ibid* 411, 503, Brown, *ibid* 469, Wiederhold, *P* 116, 171, 118, 186, Baudrimont, *J Ph* [4] 14, 81, 161, Krebs, *Z* 6, 243, Mills a Donald, *C J* 41, 18, Mills a Stevenson, *C J* 41, 23.—2 Pure MnO_2 is heated to full redness in a hard glass tube, the gas is passed through milk of lime to absorb CO_2 , then over Pt black at $c\ 250^\circ$ to cause combination of N (which is generally present) with O, then through NaOH Aq to absorb nitrites and nitrates produced, and the O is then dried by conc H_2SO_4 , CaCl_2 , and P_2O_5 . It is difficult to obtain O by this method quite free from every impurity.—3 Böttger (*J pr* 103, 316, 107, 48) says that pure O is obtained by gently heating KMnO_4 , or by the reaction of dilute HNO_3 Aq, S G c 1 064 with a mixture of PbO_2 and BaO .—4 Fairly pure O may be obtained in a regular stream, at the ordinary temperature, by compressing into small cylinders a mixture of two parts BaO , one part MnO_2 , and one part CaSO_4 , and acting on these, in a Kipp's apparatus, with HCl Aq, S G 12, diluted with an equal volume of water (Neumann, *B* 20, 3058).

Properties.—A colourless, odourless, tasteless gas. Condensed to a liquid at very low temperatures and great pressures. Liquid O is bright blue when viewed in layers 30–40 mm thick (Olszewski, *W* 42, 663). Pictet supposed that O was solidified when pressure was suddenly reduced, and the liquid was allowed to issue into the air (*C R* 86, 37), but, according to more recent experiments by Wroblewski a Olszewski (*C R* 100, 350, 979), solid O has not been obtained. Light appears bluish when observed through liquid O, or through a column of the gas very strongly compressed (Livinge a Dewar, *P M* [5] 26, 286). One litre O at 0° and 760 mm weighs 1.424488 grms at the latitude of 45° , multiplying mean result of Regnault (*Acad* 21, 158) and Jolly (*W* 6, 520) for weight of one litre H by mean S G of O, referred to H, obtained by Scott and Rayleigh (*v N* 37, 418). The atomic volume of O

($\frac{\text{At weight}}{\text{S G of liquid}}$) is c 14 (*v* Wroblewski, *C R* 102, 1010). The atomic volume of O in compounds varies according to the number of atoms with which the O is directly united, and perhaps to a small extent according to the nature of these atoms. Kopp gives 12.2 for O \cdot and 7.8 for O $\cdot\cdot$ (*v* SPECIFIC VOLUMES, vol iv p 498). The effect of the atom of O on the magnetic rotatory

power of O compounds is not yet determined with certainty (Perkin, *C J* 45, 558, v *PHYSCAL METHODS*, section *Optical*). O is absorbed by molten Ag or Pt, and given off again as the metal solidifies. O is also absorbed by charcoal. One vol cocoa nut charcoal absorbs c 18 vols O at 0° and 760 mm. (Hunter, *P M* [4] 29, 116, v also *Saussure, G A* 47, 113). According to Angus Smith (*Pr* 28, 322), the vol of O absorbed by C is eight times that of H under the same conditions. O absorbed by charcoal brings about oxidation of H_2S , PH_3 , C_2H_4O , &c, fairly rapidly (Calvert, *C J* [2] 5, 293, cf *CARBON*, vol 1 p 686).

O is the most negative of the elements except F. It combines directly with all elements except F, Cl, Br, I, Au, and Pt. At least one binary compound of O with each element, except F and Br, is known. Compounds containing O exhibit the same diverse properties. The oxides of the positive elements, as a class, are basic, and those of the negative elements are acidic (v *OXIDES*, p 658, *ANHYDRIDES*, vol 1 p 267, *BASE*, vol 1 p 445). O is a constituent of the greater number of acids, the compounds of O with H and non metallic elements are acids. When H is combined with one of the less positive metals and a relatively large quantity of O, the compound so produced is generally an acid (v *ACIDS*, vol 1 p 47). The process of combining O with another element or with a compound is called oxidation (v *OXIDATION*, p 657, cf *DEOXIDATION*, vol 1 p 377). When the process is attended with the production of so much heat that the products become self luminous, it is called combustion (v *COMBUSTION*, vol 1 p 241, cf *FLAME*, vol 1 p 549). O is more closely related chemically to S, Se, and Te than to the other elements. It belongs to Group VI, which contains the even series elements O, Cr, Mo, W, and U, and the odd series elements S, Se, and Te (v *CHROMIUM GROUP OF ELEMENTS*, vol 1 p 168, and *OXYGEN GROUP OF ELEMENTS*, this vol *infra*). O exhibits allotropy, the allotropic modification is called *Ozone* (v art *OZONE*).

Atomic weight of oxygen—The value 16.01 was obtained by Berzelius and Dulong in 1819, by burning H by CuO and weighing the water formed (4 *Ch* 15 386). In 1842 Dumas by the same method, arrived at the number 15.98 (4 *Ch* [3] 8, 189). In the same year, Lrdmann & Marchand conducted very carefully a series of five experiments by the same method, their mean result was 16.005. Ostwald (*Lehrbuch der Allgemeinen Chemie*, 1, 43) shows that the value 15.84 is obtained from Stas's experiments on the ppn of NH_4Cl by Ag. In 1805 Gay-Lussac & Humboldt (*G A* 20, 38) determined the ratio in which H and O combine, by volume, to be 2:1. Using the values for SG of these gases obtained by Biot & Arago, the atomic weight of O became 15.1. If Regnault's more accurate determinations of SG of H and O are made use of, and it is assumed that H and O combine in the ratio 2:1 by volume, then the atomic weight of O becomes 15.964. In recent years Scott has made a series of direct determinations of the relative densities of H and O (*Pr* 1897 898, *B A* 1887 668; *Pr* 42, 896, *N* 87, 439). Keiser (*B* 20, 2828, *Am* 10, 249) caused Pd to occlude H, and then drove out the

H over hot CuO, and weighed the water produced. In this way a direct weighing of the H burnt was obtained. Keiser's final value for O was 15.9492. Cooke & Richards (*Am* 10, 81) weighed H in a large balloon, and then burnt it to H_2O , they obtained $O=15.953$. Lord Rayleigh, in 1888, pointed out a source of error in all experiments wherein a gas is weighed by pumping the air out of a glass globe, and thus allowing the gas to enter (the error is due to the shrinkage of the globe when evacuated). From his own experiments, Rayleigh gave $O=15.884$ (*Pr* 43, 356), and applying his correction to Scott's most carefully determined values (*Pr* 42 396), he obtained the value $O=15.912$. Cooke & Richards applied Rayleigh's correction to their experimental data, and arrived at the conclusion that $O=15.869$ (*Am* 10, 191). The mean of the results obtained by directly weighing H and then burning it to H_2O is 15.888. The most probable value obtained by weighing H by occluding it in Pd, driving out by heat, and weighing Pd before and after, is 15.9492. The mean of the values obtained by burning H to H_2O , weighing O by loss of weight of CuO, and determining H by difference between weights of H_2O and O, is 15.992. For a criticism of the methods and results up to 1885, v Ostwald's *Lehrbuch der Allgemeinen Chemie*, 1, 41. As the atomic weights of many elements are determined by directly referring them to O, some chemists take the atomic weight of O as 16 (H thus becomes a little more than 1). Most chemists at present take $O=15.96$, which is the value used in this Dictionary.

Reactions and Combinations—The reactions and combinations of O are too many to be described here, it must suffice to indicate them in a general way. Reference should be made to the various elements and compounds for details. 1. All non metals, except F, Br, Cl, and I, combine directly with O, compounds of all, except F and Br, with O are known. 2. All metals, except Au and Pt, combine directly with O, compounds of all with O are known. 3. Many lower oxides are changed to higher oxides by heating in O, e.g. BaO , Bi_2O_3 , NO , P_2O_3 , PbO , FeO , many other lower oxides form higher oxides when O is produced in contact with the lower oxides (cf *OXIDATION*, p 657, and *OXIDES*, p 638). 4. Innumerable compounds are decomposed by O, with formation of more oxidised compounds, e.g. NH_4Aq , H_2S , PbS , and $Na_2S_2O_3Aq$, with many compounds O combines directly, e.g. P_2O_3 , HNO_3Aq , SO_2Aq , NO .

Certain oxidations do not occur when very dry O is heated with the substance to be oxidised after it has been thoroughly dried (v *CARBON*, vol 1 p 687, *Carbon monoxide*, vol 1 p 690, v also *PHOSPHORUS*, *SULPHUR*).

Detection of free oxygen—Uncombined O changes an alkaline pyrogallol solution to dark brown, white indigo becomes blue, NO combines with O to form red brown NO_2 , easily soluble in water, $CrCl_3Aq$ rapidly absorbs O, changing from blue to green (for preparation of $CrCl_3Aq$, v *NITROGEN*, *Preparation* No 1, p 557). M M P M

OXYGENATED WATER An older name for *HYDROGEN DIOXIDE* (q v, vol 1 p 722).

OXYGEN GROUP OF ELEMENTS *Oxygen, sulphur, selenium, and tellurium*. Oxygen was Z Z

prepared by Priestley in 1774; sulphur has been known from early times, selenium was discovered by Berzelius in 1817, tellurium was found by Klaproth, in 1798, in gold ore from the *Siebergberge*. The name oxygen was given by Lavoisier to express the fact that many compounds of this element are acidic, the word sulphur is supposed to be derived from *sal* = salt and *rip* = fire (burning salt or stone), the name tellurium is derived from *tellus* = the earth, selenium received its name from *σεληνη* = the moon, because of its association with and likeness to tellurium. The following table presents some of the chief properties of the four elements.—

The elements S, Se, and Te show very distinct resemblances in their chemical properties. All form hydrides MH_2 , which are gases at ordinary temperatures, SH_2 and SeH_2 in water reddens litmus, and react as very weak acids. TeH_2 is not acidic, these hydrides are decomposed by heat, stability in this respect decreasing as at w of M increases. These hydrides are produced by passing H over molten S, Se, or Te, also by reacting on various sulphides, selenides, and tellurides, with dilute acids. The hydride of O, viz OH_2 , is analogous in composition, and is formed similarly to those of S, Se, and Te, OH_2 is neutral H_2O , and H_2S , are also similar in composition and many pro-

	OXYGEN	SULPHUR	SELENIUM	TELLURIUM
<i>Atomic weights</i>	15.96	81.98	78.8	125
<i>Molecular weights</i>	81.92 (47.88 as ozone)	63.96 (? 191.88 at lower temps)	157.6 (? 236.4 at c 800°)	250
One or more compounds of each element have been gasified, specific heats of solid S, Se, and Te have been determined directly				
<i>Melting points (approx)</i>	(not solidified)	112°–117°	c 200°	455°
<i>Boiling points (approx)</i>	–181.4° at 760 mm	449°	680°	below 1400°
<i>Spec grav. (approx)</i>	1.12 (liquid at B.P.)	1.96–2.05	4.3–4.8	6.2
<i>Atom weight</i>	18.3	16	17.5	20
<i>Spec grav (approx)</i>				
<i>Occurrence and preparation</i>	Very large quantities of O occur uncombined in air, O is a constituent of almost all rocks and minerals, and also of all living organisms, it forms 8ths of water. By heating various compounds, chiefly oxides of metals, and in many other ways.	In large quantities in Sicily, Spain, &c., sulphides and other S compounds are common minerals, S compounds are found in many living organisms. Prepared by refining crude sulphur, also by roasting various sulphides in absence of air.	Uncombined in very small quantities, compounds of Se with S, Fe, Cd, Hg, Pb, &c., occur very sparingly. By passing SO_2 into H_2SeO_4 aq., also by adding HCl aq. to $KSeCN$ aq.	Uncombined, also in combination with Au, Pb, S, Sb, Bi, &c., but in very small quantities. From solution of a salt of H_2TeO_4 , by passing in SO_2 , or by Zn.
<i>Physical properties</i>	A colourless, odourless gas, bluish when much compressed, condensed to a colourless liquid, appearing blue in thick layers, at great pressures and low temperatures, has not been solidified.	A yellow, brittle crystalline, solid, also a plastic semi-fluid, amorphous mass. Very bad conductor of electricity.	A red-grey, lustrous crystalline, solid, also an amorphous, vitreous, grey-black solid. The amorphous form is a very bad conductor of electricity, the crystalline form conducts fairly, and conductivity increases when exposed to sunlight.	A lustrous, white, metal like, crystalline, brittle, solid, also as an amorphous powder. Bad conductor of electricity.

	OXYGEN	SULPHUR	SELENIUM	TELLURIUM
<i>Chemical properties</i>	Combines directly with all elements except F, Cl, Br, I, Au, and Pt, oxides are known of all elements except F and Br. Compounds exhibit very different properties. O is a constituent of most acids, and of all alkalis. Compound with H, OH ₂ , is a neutral oxide, H ₂ O ₂ is a peroxide.	Combines directly with most elements. Sulphides are basic, or acidic, or neutral, according to nature and relative quantity of element combined with S. S is a constituent of several acids. H ₂ S is feebly acidic.	Compounds with several elements are known, they resemble sulphides. H ₂ Se is very feebly acidic. H ₂ SeO ₃ and H ₂ SeO ₄ are much weaker acids than H ₂ SO ₃ and H ₂ SO ₄ .	Combines with several elements (compounds not studied very fully). Binary compounds resemble those of Se. H ₂ Te is not acidic. TeO ₂ and TeO ₃ do not form acids by acting with water, but these oxides are obtained by heating the acids H ₂ TeO ₃ and H ₂ TeO ₄ .

perties. Se₂H₂ and Te₂H₂ have not been prepared. Binary compounds of S, Se, and Te with metals are analogous in composition to the metallic oxides. Chlorides of the form MCl₂ are known, M = O, S, Se, Te, no compound of S, Se, or Te analogous to O₂Cl₂ is known, nor has a compound of O and Cl analogous to SCl₂, SeCl₂, and TeCl₂ been obtained. The stability of the chlorides towards heat increases in passing from O to Te. The oxides of S, Se, and Te—MO₂ and MO₃—are acidic (SeO₃ is not known), acidity decreasing rapidly as at w of M increases, TeO₃ is scarcely acidic, but a corresponding acid H₂TeO₄ exists, TeO₃ combines with some strong anhydrides to form compounds of the type of salts. The oxyacids H₂MO₃ and H₂MO₄ are analogous in composition, but the acids of S are very much stronger than those of Se or Te, those acids yield oxychlorides, MOCl₂ and MO₂Cl₂, by reacting with PCl₅. Many other oxyacids of S are known.

The elements of the O family exhibit allotropy, O and S in a very marked way, Se less markedly, and Te only slightly. The allotropic forms of O exist as gases, their molecular formulæ are O₂ and O₃, the allotropic forms of S, Se, and Te are known with certainty only as solids. It is doubtful whether molecules of S or Se, other than S₂ and Se₂, exist as gases through any considerable range of temperature, the only molecule of Te existing as a gas is Te₂. The mol w of S in solution is certainly greater than S₂, and probably varies according to the solvent.

The elements S, Se, and Te form the odd-series members of Group VI, of which group O forms the first even series member, the other even series members are Cr, Mo, W and U. All the members of Group VI are characterised by forming acidic oxides MO₃, as the at w increases these oxides become less acidic. Only the odd series members, and O, form hydrides. All form chlorides MCl₄, the stability of which towards heat increases from O to U, the higher members also form several other stable chlorides. Group VI falls into two families, (i) S, Se, Te, (ii) Cr, Mo, W, U, O is the typical element of the group (*cf.* CHEMISTRY GROUP OF ELEMENTS, vol ii p 188, v. also OXYGEN, SULPHUR, SELENIUM, TELLURIUM). M. M. F. M.

α-OXY-GLUTARIC ACID

CO₂H CH(OH) CH₂ CH₂ CO₂H *Glutaric acid* [73°] ? (M) Formed from amido glutaric acid and HNO₂ (Ritthausen, *J pr* 103, 239, Markownikoff, *A* 182, 348) Occurs in beetroot molasses (Lippmann, *B* 15, 1156) Readily yields the lactic acid—CaA''₂aq—MgA''₂aq—PbA''₂aq—ZnA''₂aq—AgA''₂aq

Lactic acid C₃H₅O₃, s.e.

CH₃—CO—CH₂—CH₂—CO₂H [50°] Very hygroscopic needles (Wolff, *A* 260, 126)—CaA''₂aq—BaA''₂aq—ZnA''₂aq Yields glutaric acid [98°] on reduction by HI.

β-Oxy-glutaric acid Methyl derivative CO₂H CH₂—CH(OMe)—CH₂—CO₂H Formed by oxidising di-allyl-carbinol with 5 p.c. solution of KMnO₄ (Ryabinin, *J pr* [2] 23, 274) Extracted with ether (*v* OXYTARTARIC ACID)

Salts—CaA'' (at 100°) Syrup, covered by crystalline crusts—BaA''—AgA'' Prisms

αβ-Di-oxy glutaric acid

CO₂H CH₂—CH(OH)—CH(OH) CO₂H [156°] Formed by boiling glutaric acid-di bromide with water and CaCO₃. Needles (from water), or six sided tables (from alcohol). V e sol. water, v sol alcohol. Its Ca salt is easily soluble (Kiliani, *B* 18, 2517)

γ-Di-oxy glutaric acid

CO₂H CH(OH) CH₂—CH(OH) CO₂H Formed by heating the di-oxy-propane tri-carboxylic acid, which is obtained by oxidising isosaccharic acid with HNO₃. Colourless prisms V e sol water, v sol alcohol, sl sol ether. Its Ca salt is sparingly soluble (Kiliani, *B* 18, 2516)

Tri-oxy-glutaric acid C₅H₄O₇ [127°] Formed from arabinose or sorbin by oxidation with HNO₃ (SG 12) (Kiliani, *B* 21, 3006, 3276)—KA'' monoclinic plates

OXYGUANIDINE Obtained by boiling cyanamide with alcoholic hydroxylamine hydrochloride (Prætorius a Seidler, *J pr* [2] 19, 399). —(ON₂H₂)₂·PtCl₄ ruby-red prisms

OXY-HÆMOGLOBIN *v* HÆMOGLOBIN.

OXYHALOID COMPOUNDS Compounds of F, Cl, Br, or I, with O and another element. Oxyhaloid compounds of non-metals are often formed by reacting with PCl₅, PBr₅, or PI₅ on oxyacids, e.g. SO₂Cl₂ from SO₂(OH)₂; they are also produced sometimes by heating together an

oxide and chloride, *eg* BOCl by heating B_2O_3 with BCl_3 , sometimes the halogen is directly combined with an oxide, *eg* COCl and NOCl by combining CO and NO with Cl , and BOCl_2 by heating Cl with B_2O_3 mixed with C . Metallic oxyhaloid compounds are formed by decomposing the haloid compounds of the metals by water or steam, by heating together oxides and haloid compounds, and in some cases by the incomplete *ppn* of haloid compounds in solution by alkali. Non metallic oxyhaloid compounds are decomposed by water to haloid acids and other oxides or oxyacids of the non metal present. Some metallic oxyhaloid compounds are also decomposed by hot water to haloid acids and metallic oxides, in this respect the oxyfluorides are the most stable, and the oxyiodides, as a class, are more stable than the oxybromides and oxychlorides. The oxyhaloid compounds of H are acids. All elements form one or more oxy haloid compounds. A metal which readily forms many oxyhaloid compounds generally forms one or more oxides with acidic reactions, and is otherwise more or less non metallic in its chemical properties. M M P M

DI-OXY-HENDECIOIC ACID $\text{C}_{11}\text{H}_{22}(\text{OH})_2\text{O}_2$ [85°] Formed by the oxidation, in alkaline solution, of hendecenoic acid derived from castor oil (Hazura a Grussner, *M* 9, 952). Needles, sol hot water.

OXY HENICOSOIC ACID
 $\text{C}_{10}\text{H}_{20}(\text{CH}_2\text{OH})(\text{CO}_2\text{H})$ [103.5°] Occurs in the soda soap from carnauba wax (*q v*). Crystalline powder (from petroleum), sl sol alcohol, m sol isobutyl alcohol (whence it separates as a jelly) (Starcke, *A* 223, 310). As its alcoholic solution is not *ppd* by acetate of Mg , Cu , or Pb , it is possibly a lactone $\text{C}_{10}\text{H}_{20}\langle\frac{\text{CH}_2}{\text{CO}}\rangle\text{O}$. Heated with soda lme it splits off H_2 , forming a dibasic acid $\text{C}_{10}\text{H}_{18}(\text{CO}_2\text{H})_2$ [90°].

PENTA OXY HEPTANE The only anhydride $\text{C}_6\text{H}_{10}(\text{OH})_5\text{O}$ is formed by the action of HOCl followed by KOH on di allyl carbinol (Reformatsky, *J R* 21, 295). The acetyl derivative $\text{C}_6\text{H}_{11}(\text{OAc})_5$ (169.5°) S G \pm 919 is described by Saytzeff (*A* 185, 129).

OXY-HEPTANE PHOSPHONIC ACID
 $\text{C}_6\text{H}_{11}\text{PO}_4$, *ie* $\text{C}_6\text{H}_{11}\text{CH}(\text{OH})\text{PO}(\text{OH})_2$ [185°] Formed from α -naphthol and PCl_5 followed by water (Fossek, *M* 7, 27). Monoclinic tables $a b c = 1.844 \ 1.957$, $\beta = 74^\circ$ — CaA'' .

The compound $\text{C}_6\text{H}_{11}\text{CH}(\text{OH})_2\text{PO}(\text{OH})_2$ [160°] is formed by heating α -naphthol with hypophosphorous acid (Ville, *C R* 109, 72). It yields the salts BaA'_3 , 3aq , PbA'_3 , 3aq , KA'_4 , 4aq , and $(\text{C}_6\text{H}_{11}\text{CH}(\text{OAc}))_2\text{PO}_2\text{H}$ [94°].

OXY-HEPTENOIC ACID
 $\text{CHMe}(\text{OH})\text{CH}(\text{C}_6\text{H}_5)\text{CO}_2\text{H}$ Formed from allyl-acetoacetic ether and sodium amalgam (Zeidler, *A* 187, 45). Syrup, miscible with water — BaA'_2 — $\text{Zn}(\text{OHA})'$.

Oxy-heptenoic acid Nitrile
 $\text{CHEt} \cdot \text{CMe} \cdot \text{CH}(\text{OH}) \cdot \text{CN}$ Formed from methyl-ethyl-acrolein and HCy (Johanny, *M* 11, 401). Converted by hydrochloric acid into the amide $\text{CHEt} \cdot \text{CMe} \cdot \text{CH}(\text{OH}) \cdot \text{CONH}_2$ [101°]. The nitrile yields an oily acetyl derivative (114° at 22 mm), γ sol ether.

Oxy-heptenoic lactone *v* MESITONIC ACID

OXYHEPTIC ACID $\text{C}_7\text{H}_{14}\text{O}_2$, aq ? [185°] **A** product of the action of bromine, followed by alcoholic potash on isobutyl acetoacetic ether (Demarcay, *C R* 86, 1135). Pearly scales (from water). Yields $\text{C}_7\text{H}_{14}\text{O}(\text{OEt})\text{NH}_2$? [87°].

α -OXY-HEPTOIC ACID $\text{C}_7\text{H}_{14}\text{O}_3$, *ie* $\text{Pr} \cdot \text{CH}_2 \cdot \text{CH}_2 \cdot \text{CH}(\text{OH}) \cdot \text{CO}_2\text{H}$ [60°] (Helms, *B* 8, 1167), [65°] (Lay, *J R* 9, 141). Formed by heating aqueous potassium bromo heptoate at 140° Prisms — MeA' [160°–165°].

Amide [147°] Hexagonal tables
 α Oxy heptoic acid $\text{C}_7\text{H}_{14}\text{CH}(\text{OH})\text{CO}_2\text{H}$ [60.5°] Formed from its ether, EtA' (203°), which is produced, as well as the ethyl derivative thereof $\text{C}_7\text{H}_{11}\text{CH}(\text{OEt})\text{CO}_2\text{Et}$ (c 224°) by the action on oxalic ether of zinc and isoamyl iodide at 100°, followed by water (Frankland a Duppa, *Pr* 14, 191). Scales — BaA'_2 — CuA'_2 .

β Oxy-heptoic acid
 $\text{CHMe} \cdot \text{CH}(\text{OH}) \cdot \text{CMe} \cdot \text{CO}_2\text{H}$ [112°] The chief product of the action of sodium on isobutyric ether (Hantzsch, *A* 249, 60). Silky needles, m sol water, volatile with steam — BaA'_4 , 4aq .

Ethyl derivative of the ethyl ether
 $\text{Pr} \cdot \text{CH}(\text{OEt}) \cdot \text{CMe}_2 \cdot \text{CO}_2\text{Et}$ (181° cor) Oil.

β Oxy heptoic acid $\text{CHMe}(\text{OH})\text{CMe} \cdot \text{EtCO}_2\text{H}$
Formed by reducing $\text{CH}_2 \cdot \text{CO} \cdot \text{CMe} \cdot \text{CO}_2\text{Et}$ with sodium amalgam (Saur, *A* 188, 266). Syrup, γ sol water — $\text{CuC}_6\text{H}_{11}\text{O}_2$ — $\text{AgC}_6\text{H}_{11}\text{O}_2$ plates, m sol hot water.

β Oxy-heptoic acid $\text{CMePr}(\text{OH})\text{CH} \cdot \text{CO}_2\text{H}$
Formed by oxidising $\text{CMePr}(\text{OH})\text{C}_2\text{H}_5$ with KMnO_4 (Semljantzin, *J pr* [2] 23, 267). Syrup — CaA' (at 100°) — BaA'_2 — AgA' prisms.

β Oxy-heptoic acid $\text{CHEt}_2(\text{OH})\text{CH}_2 \cdot \text{CO}_2\text{H}$ [39°] Formed by oxidising $\text{CHEt}_2(\text{OH})\text{C}_2\text{H}_5$ with KMnO_4 (Schrokooff, *J pr* [2] 23, 201). Thin needles — $\text{LiA}' \cdot \text{aq}$ — $\text{CaA}'_2 \cdot \text{aq}$ — $\text{BaA}'_2 \cdot 2\text{aq}$ — $\text{PbA}'_2 \cdot 2\text{aq}$ S 6 at 19° — $\text{CuA}'_2 \cdot 5\text{aq}$ — AgA' .

γ Oxy-heptoic acid
 $\text{CHPr}(\text{OH})\text{CH}_2 \cdot \text{CH}_2 \cdot \text{CO}_2\text{H}$ Syrup — BaA'_2 — AgA' curdy *pp*.

Lactone $\text{Pr} \cdot \text{CH} \cdot \langle \frac{\text{CH}_2}{\text{CO}} \rangle$ (235° γ V)

Formed by the action of HI and P on the hexa oxy heptoic acid obtained from dextrose (Kiliani, *B* 18, 3066, 19, 1128). Obtained also from propyl paraconic acid $\text{Pr} \cdot \text{CH} \cdot \langle \frac{\text{CH}(\text{CO}_2\text{H})}{\text{O} \cdot \text{CO}} \rangle$, CH_2

by distilling, converting the resulting $\text{CHPr} \cdot \text{CH} \cdot \text{CH}_2 \cdot \text{CO}_2\text{H}$ into γ bromo heptoic acid, and boiling this with water (Fittig, *B* 20, 3180, *A* 255, 76).

An isomeric lactone (220° cor) is obtained in like manner from levulose.

γ Oxy-isheptoic acid
 $\text{CHPr}(\text{OH})\text{CH}_2 \cdot \text{CH}_2 \cdot \text{CO}_2\text{H}$ Very unstable — BaA'_2 — AgA' .

Lactone $\text{C}_6\text{H}_{11}\text{O}_2$ (225° uncor) S 3 in the cold. Formed from γ bromo isheptoic acid and also by distilling isopropyl paraconic acid (Fittig a Zanner, *A* 255, 94). Liquid, volatile with steam.

γ Oxy-heptoic acid — BaA'_2 — AgA'
Lactone $\text{C}_6\text{H}_{11}\text{O}_2$ [11°] (220° γ V) S 83 at 0°. Formed from tetrachloric acid and cold $\text{HBr} \cdot \text{aq}$ (Fittig a Kraft, *A* 208, 86). Liquid.

γ -Oxy-heptoic acid Lactone
 $\text{CHMe} \cdot \langle \frac{\text{CH}_2}{\text{O} \cdot \text{CO}} \rangle \cdot \text{CMe}_2$ [52°] (86° at 15 mm).

Formed by reducing β -acetyl valeric acid with

$\text{CH}_3\text{CH}(\text{OH})\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_2(\text{OH})$ *Heptylene*
δ-glycol (235° at 710 mm) S G at 0° = 9809
Prepared by reduction of aceto butyl alcohol
 $\text{CH}_3\text{COCH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{OH}$ with sodium-

Iso-oxy-hexic acid [187°] Formed in like manner from isopropyl-acetoacetic ether (D). Reduced by Zn and H_2SO_4 to $C_7H_{15}O_4$ (?) [113°]

Yields $C_6H_{11}O_2(NH_2)_2$ (?) [240°] and another amide $C_6H_9O_2(NH_2)(OEt)_2$ (?) [95°]

DI-OXY-HEXINENE or DI-OXY-HEXYLENE

α -OXY-HEXIC ACID $C_6H_{10}O_3$ *Leucic acid* Mol w 182 [73°] Formed by the action of nitrous acid on leucine (Strecker, *A* 68, 55, Thudichum, *C J* 14, 807, Waage, *A* 118, 297) Needles, v sol water, alcohol, and ether — BaA'_2 , silky laminae (from hot alcohol) — CaA'_2 , — CoA'_2 , — CuA'_2 , — ZnA'_2 , aq S 33 at 16°, 5 at 100° — ZnA'_2 , 2aq (Körner, *G* 13, 356) — AgA' crystalline pp

An isomeric or identical oxy hexoic acid [62°] is formed by the action of Na_2CO_3 on bromohexoic acid got from fermentation hexoic acid (Jehssoff, *J R* 12, 367, *Bn* 1, 523) It yields the salts CaA'_2 , BaA'_2 , MgA'_2 , 2aq, ZnA'_2 , 2aq (S 14 at 16°, 21 at 100°), CuA'_2 , and AgA' It forms a crystalline amide [142°]

α -Oxy-hexoic acid $CEt_2(OH)CO_2H$ [80°] S. 35 at 17.5° Formed from its ether which is got from oxalic ether by the action of $ZnEt_2$, followed by water (Frankland, *Pr* 12, 896, Frankland & Duppa, *Pr* 13, 140, *A* 135, 26, Geuther, *Z* 1867, 705, Fittig, *A* 200, 21) Trichmic crystals, v sol water, alcohol, and ether Yields di-ethyl ketone on oxidation — NH_4A' — BaA'_2 — BaA'_2 , 5aq — ZnA'_2 S 33 at 16° — CuA'_2 — AgA' aq needles

Methyl ether MeA' (165°) SG 165 987

Ethyl ether EtA' (175°) VD 5 24 (calc 553) SG 187 961 Oil Formed from $COClCO_2Et$ and $ZnEt_2$ (Henry, *B* 5, 949)

Isoamyl ether (225°) SG 13 983

Nitrile $CEt_2(OH)CN$ Formed from $COEt_2$ and HCy (Tiemann & Friedlander, *B* 14, 1974) Oil

α -Oxy-hexoic acid $CH_2PrCH(OH)CO_2H$ [56°] Got by saponification of its nitrile, which is formed from isovaleric aldehyde and HCy (Erlenmeyer & Sigel, *B* 7, 1109, Ley, *B* 10, 231) Formed also by heating isobutyl tartaric acid at 180° (Guthzeit, *A* 209, 239) Plates — ZnA'_2 , 2aq S 12 at 16°

Nitrile Oil, lighter than water

β Oxy-hexoic acid $CHMe(OH)CHEtCO_2H$ Formed from methyl acetoacetic ether and sodium-amalgam (Waldschmidt, *A* 188, 240) Syrup — NaA' — CuA'_2 — AgA' plates

α Oxy-hexoic acid $CMe_2CH(OH)CO_2H$ [68°] Formed by reducing tri methyl pyruvic acid with sodium amalgam (Glucksmann, *M* 10, 780) Monoclinic crystals — AgA'

β Oxy-hexoic acid $CHEt(OH)CHMeCO_2H$ Formed by reducing propionyl propionic acid (Hantzsch & Wohlbruck, *B* 20, 1320) — NaA'

γ -Oxy-*n*-hexoic acid

$CHEt(OH)CH_2CH_2CO_2H$ The salts are formed by the action of bases on the lactone The free acid quickly changes to lactone — CaA'_2 — BaA'_2 — AgA'

Lactone $C_6H_{10}O_3$ *ee* $CHEt \begin{smallmatrix} \diagup CH_2 \\ \diagdown O-CO \end{smallmatrix}$

(220°) **Formation** — 1 By boiling with water the bromo-hexoic acid formed by the union of hydrosorbic acid with HBr (Fittig, *B* 13, 955, *A* 208, 67) — 2 By heating glutamic acid with HI and P (Kiliani & Kleiman, *B* 17, 1800) — 3 By reduction of metasaccharin with HI

(Kiliani, *B* 18, 642) — 4 By warming hydrosorbic acid with dilute (1:1) H_2SO_4 (Fittig, *A* 256, 184)

Properties — Liquid, m sol water Volatile with steam Its aqueous solution becomes turbid when heated to 40°, but clear again at 80° K_2CO_3 separates it from the solution HI and P reduce it to *n*-hexoic acid

Amide $CHEt(OH)CH_2CH_2CONH_2$ [74°] Formed by heating the lactone with alcoholic NH_3 at 100° Prisms, v sol water and alcohol, sl sol ether

γ -Oxy-isohexoic acid

$OMe_2(OH)CH_2CH_2CO_2H$ Crystalline, but very unstable — BaA'_2 (at 100°) — AgA'

Lactone $C_6H_{10}O_3$ (207° i V) Formed from terebic acid by distillation, by successive treatment with HBr and water, or by boiling with dilute H_2SO_4 (Fittig & Bredt, *A* 200, 58, 259, Geisler, *A* 208, 43, Erdmann, *A* 228, 181) Formed also by oxidising isohexoic acid with $KMnO_4$ (Bredt, *A* 208, 59) Liquid, v sol water Neutral in reaction Its aqueous solution becomes turbid at 35°, clear again at 80° Yields on oxidation with HNO_3 the acid $C_6H_9O_4$ [68°], whence $CaC_6H_9O_4$, 7aq and $AgC_6H_9O_4$ Boiling $NaOEt$ forms an anhydride $C_6H_8O_3$ [106°], whence $HClAg$ forms $C_6H_8O_3$ (209°)

γ Oxy-hexoic acid $CH_2(OH)CH_2CHEtCO_2H$ Formed from $CH_2COCEt(CH_2CH(OH)CO_2Et$ by boiling with baryta water (Chanlaroff, *A* 226, 335) Thick liquid, changing to the lactone on boiling its solution $Ba(C_6H_9O_3)_2$ crystals (from alcohol) — CaA'_2 — AgA' , needles (from water)

Lactone $CH_2 \begin{smallmatrix} \diagup CH_2CHEt \\ \diagdown O-CO \end{smallmatrix}$ (215°) SG

12 1035 Liquid, m sol cold water

γ Oxy hexoic acid

$CHMe(OH)CH_2CHMeCO_2H$ Formed by reduction with sodium amalgam from β acetyl isobutyric acid derived from a bromo propionic ether and sodium acetoacetic ether (Fittig & Gottstein, *A* 216, 30) Formed also by reducing saccharin or isosaccharin with HI and P (Liebermann & Scheibler, *B* 16, 1821, Kiliani, *B* 18, 635) — BaA'_2

Lactone (206°) Liquid, sol 20-25 volumes of water An isomeric anhydride [187°] is also formed by reducing isosaccharin

γ Oxy hexoic acid

$CHMe(OH)CHMeCH_2CO_2H$ Formed by reducing β -acetyl *n* butyric acid (Gottstein, *A* 216, 36) Yields a lactone (210°)

δ Oxy-hexoic acid

$CHMe(OH)CH_2CH_2CH_2CO_2H$ Obtained by reducing γ -acetyl *n* butyric acid with sodium amalgam at 30° (Fittig & Wolff, *A* 216, 138) When boiled with water it is partly converted into the lactone When the lactone is boiled with water it is partly converted into the acid Equilibrium occurs with 65 p c lactone and 35 p c acid — AgA'

Lactone [19°] (c 231° i V) Colourless liquid, solidifying below 0° Miscible with water but separated by K_2CO_3

Oxy hexoic acid $C_6H_{10}O_3$ [108°] Formed by oxidising oil of millet with $KMnO_4$ (Kassner, *As Ph* [8] 25, 1081) Plates, v sol water. Yields an acetyl derivative [71°]

Di-oxy-hexoic acid [141°] Formed from ethyl crotonic acid by successive treatment with

bromine and with water at 100° (Fitting & Howe, *A* 200, 39) Trimetric prisms, $a \cdot b \cdot c = 96 \cdot 1 \cdot 33$ V sol water and alcohol.— CaA' , 8 $\frac{1}{2}$ aq. bright-blue needles

Di-oxy-hexoic acid

$\text{CHMe}(\text{OH})\text{CH}(\text{OH})\text{CO}_2\text{H}$ [96°] Formed by oxidising ethyl crotonic acid with KMnO_4 (Fitting, *B* 21, 920)

Di-oxy-hexoic acid

$\text{CHEt}(\text{OH})\text{CMe}(\text{OH})\text{CO}_2\text{H}$ [152° cor] Formed by oxidation of methyl ethyl acrolein (Lieben & Zeissel, *M* 4, 65) Needles or prisms.— CaA' , 3 aq crystalline m ss

Hexa-oxy-hexoic acid $\text{C}_6\text{H}_{10}\text{O}_7$, *Arabinose-carboxylic acid* Formed from arabinose by the action of aqueous HCl , followed by saponification (Kiliani, *B* 29, 3033, 20, 339) When liberated from its salts it immediately forms the lactone.— CaA' , amorphous

Amide Minute needles

Lactone $\text{C}_6\text{H}_{10}\text{O}_6$ [145°–150°] $[\alpha]_D^{20} = -54.8$ Needles or prisms Yields metasaccharic acid dilactone $\text{C}_6\text{H}_8\text{O}_6$ on oxidation by HNO_3 and n hexoic acid on reduction by HI and P

DI OXY HEXYLENE $\text{C}_6\text{H}_{10}\text{O}_2$, *Hexylene glycol* (218°–225°) Formed from epichlorohydrin and Na (Hubner & Miller, *A* 159, 186) Is perhaps di oxy hexylene (Claus, *B* 10, 556)

OXY HIPPURIC ACID $\text{C}_8\text{H}_7\text{NO}_4$, *se* $\text{C}_8\text{H}_7(\text{OH})\text{CO NH CH}_2\text{CO}_2\text{H}$ Formed from m amido hippuric acid by the diazo reaction (Griess, *B* 1, 190, Conrad, *J pr* [2] 15, 259) Needles, v sol hot water

OXY HYDRASTININE *v* **HYDRASTINE**

OXYHYDROANTHRANOL *v* **OXYANTHRANOL**

HYDRIDI

TRI OXY HYDROBENZAMIDE *v* **OXY BEN**

ZOIC ALDEHYDE

OXYHYDROBENZOIC ACID $\text{C}_8\text{H}_6\text{O}_3$ [275° cor] Formed from oxy urvic acid and aqueous KMnO_4 (Oppenheim & Emmerling, *B* 9, 327) Needles Yields benzoic acid by potash fusion.— CaA' , 3 aq.— AgA' , v sl sol water

OXY HYDRO COUMARILIC ACID *v* **COUMA**

RILIC ACID

OXY HYDRONAPHTHOQUINONE *v* **HYDRO**

JUGLONE

OXY HYDROQUINOLINE *v* **OXY QUINOLINE**

HYDRIDE

OXY HYDROQUINONE *v* **TRI OXY BENZENE**
OXY-DIHYDROQUINOXALINES *v* **OXY**

QUINOXALINE DIHYDRIDE

OXYHYPOGAEIC ACID $\text{C}_8\text{H}_8\text{O}_4$ [34°] Formed from di bromo palmic acid and Ag_2O (Schroder, *A* 143, 36) White mass

OXY-DIIMIDODIAMIDODIASTIN *v* **ISATIN**

OXY IMIDO METHYL PYRIMIDINE **DI**

HYDRIDE $\text{C}(\text{NH})\langle\text{NH CMe}\rangle\text{CH}$ *Imido-*

methyl uracil [270°] Formed from guanidine, alcohol, and acetoacetic ether (Jaeger, *A* 262, 365) Prisms, v sol hot water Yields a dibromide [160°] MeI forms $\text{C}_4\text{H}_5\text{MeN}_3\text{O}$ [312°], whence $\text{B}'\text{HI}$ [212°], $\text{B}'\text{HClaq}$, and $\text{B}'\text{H}_2\text{SO}_4$ [270°]

Salts— $\text{B}'\text{HCl}$ [295°]— $\text{B}'\text{H}_2\text{SO}_4$ [180°].— $\text{B}'\text{HNO}_3$, needles

Oxy imido-di-methyl-pyrimidine dihydride

$\text{C}(\text{NH})\langle\text{NH CMe}\rangle\text{CMe}$ [320°] Formed

from guanidine and methyl-acetoacetic ether— $\text{B}'\text{H}_2\text{SO}_4$ [265°]— $\text{B}'\text{HNO}_3$ [200°]. Plates

OXY-IMIDO-PHENYL-PYRIMIDINE **DI**

HYDRIDE $\text{C}(\text{NH})\langle\text{NH OPh}\rangle\text{CH}$ *Imido*

phenyl-uracil [294°] Formed from guanidine

and benzoyl acetic ether (Jaeger, *A* 262, 372)

Amorphous powder, insol water and alcohol

OXY IMIDO-DIPHENYL SULPHIDE

$\text{NH}\langle\text{C}_6\text{H}_5\rangle\text{S}$. Formed from oxydi

phenylamine by heating with S (Bernthsen, *A*

230, 182) Colourless mass, v sol alcohol and

ether FeCl_3 forms a chocolate pp of

DI-OXY-IMIDO-DIPHENYL SULPHIDE

$\text{NH}\langle\text{C}_6\text{H}_4(\text{OH})\rangle\text{S}$ Formed by reducing

thionol $\text{N}\langle\text{C}_6\text{H}_4(\text{OH})\rangle\text{S}$ which is itself got

by the action of conc H_2SO_4 on imido diphenyl

sulphide (Bernthsen, *A* 230, 188) Colourless

needles, v sl sol water Yields a tri acetyl

derivative [156°]

DI-OXY-IMIDO-PYRIDINE DIHYDRIDE *v*

DI OXY AMIDO PYRIDINE

DI OXY DI IMIDO QUINONE

$\text{C}_8(\text{NH})_2(\text{OH})_2\text{O}_2$ Formed by oxidising tetra

oxy di amido benzene by FeCl_3 (Nietzki, *B* 16,

2094, 18, 503) Plates, v sl sol alcohol

OXY INDAZOLE $\text{C}_8\text{H}_7(\text{OH})\text{N}_2$ [215°–266°](?)

Formed by boiling diazo indazole with water

(Witt, Noeltling, & Grandmougin, *B* 23, 3642)

Needles, sl sol cold water

OXY-INDOLE *v* **OXINDOLE**

(a) **OXY INDONAPHTHENE** $\text{C}_{12}\text{H}_8\text{O}$ *se*

$\text{C}_6\text{H}_4\langle\text{CO}\rangle\text{CH}_2$ (a) *Hydrindone* [40°] [244°]

Formed by heating o cyano benzyl acetic ether

with conc HClAq (Gabriel & Hausmann, *B* 22,

2018) Colourless tables Conc HClAq at 100°

forms $(\text{O}_2\text{H}_2)_n$, not melted at 280°

Oxim $\text{C}_6\text{H}_5(\text{NOH})$ [146°] White needles

(from alcohol), sol alkalis

Phenyl hydrazide $\text{C}_6\text{H}_5(\text{N}_2\text{HPh})$ [c

120°] White prisms, converted by conc HClAq

into $\text{C}_6\text{H}_5\text{N}$ [235°]

References—**DI BROMO** and **CHLORO DI-OXY**

INDONAPHTHENE

OXY-ITACONIC ACID $\text{C}_8\text{H}_6\text{O}_4$ Formed by

boiling acetic acid with baryta water (Meilly,

A 171, 153) Oil— BaA'' (at 160°)— $\text{Ag}_2\text{A}''$

flocculent pp

OXY-JUGLONE *v* **JUGLONE**

OXY-LEPIDINE *v* **OXY METHYL QUINOLINE**

OXY LEUCOTIN *v* **COTOIN**

OXY-LUTIDINE *v* **OXY DI METHYL PYRIDINE**

OXY-MALONIC ACID *v* **TARTRONIC ACID**

OXY-MARGARIC ACID $\text{C}_{17}\text{H}_{32}\text{O}_4$ [80°].

Occurs in adipocere (Ebert, *B* 8, 775)— $\text{Ag}_2\text{A}'$

amorphous

OXY-TRIMELLITIC ACID $\text{C}_6\text{H}_4\text{O}_4$ *se*

$\text{C}_6\text{H}_4(\text{OH})(\text{CO}_2\text{H})_2$ [5421] [c 245°] Formed

by fusing sulpho trimellitic acid with potash

(Jacobsen & Meyer, *B* 16, 192) Prisms (con

taining 2 aq) With HClAq at 240° it yields m -

oxy-benzoic acid— $\text{Ba}_2\text{A}'''$, 5 aq small prisma.

crystalline $C_6H_4(OH)(NMe_2)CO_2Me - B'HCl - B'HI - B'H_2PtCl_4$ 4sq small yellow prisms

Oxy-methylamido-benzoic acid *Methyl derivative* $C_6H_4(OMe)(NHMe)CO_2H$ [above 200°] Formed from potassium amido anisate and MeI (Griess, *B* 5, 1042, 6, 588) Slender needles, v sl sol hot water Yields $B'HClAq$ Further treatment with MeI and KOH forms $C_6H_4 \begin{smallmatrix} NMe_2 \\ \diagup \quad \diagdown \\ CO \end{smallmatrix} > O$ 6sq which gives the salts $B'H_2PtCl_4$ and $B'HI$ aq, and on distillation yields $C_6H_4(NMe_2)(OMe)CO_2H$ (288°)

β OXY METHYL A AMIDO BUTYRIC ACID $CHMe(OH)CH(NHMe)CO_2H$ *S* 56 at 12° Formed from β methyl glycidic acid and methylamine at 100° (Selnasky, *Bl* [2] 43, 247) Crystals

α OXY-TETRA-METHYL-DI- p -AMIDO-TRI-PHENYL CARBINOL $C_{24}H_{28}N_2O_7$ *4e* $C_6H_4(OH)C(OH)(C_6H_4NMe_2)_2$ Formed by oxidising the leuco-base obtained by condensation of salicylic aldehyde with dimethylaniline (O Fischer, *B* 14, 2522) Dyes yellowish green

α OXY-TETRA-METHYL DI p AMIDO TRI-PHENYL METHANE $C_{24}H_{28}N_2O_7$ *4e* $C_6H_4(OH)CH(C_6H_4NMe_2)_2$ Leuco base of salicylic aldehyde green [128°] Prepared by heating a mixture of dimethylaniline (24 pts), salicylic aldehyde (10 pts), and $ZnCl_2$ (20 pts) to 100° for 7 or 8 hours, the yield being nearly theoretical (Fischer, *B* 14, 2522) Colourless rosettes Sol hot alcohol and benzene, nearly insol water Combines with both acids and bases On gentle oxidation it gives a green of yellow shade

Acetyl derivative $C_{24}H_{26}N_2(OAc)$ Iridescent plates [144°]

The isomeride [163°] from p oxybenzoic aldehyde yields a green dye and forms an acetyl derivative [146°] crystallising in prisms

Di-oxy tetra methyl di amido tri phenyl methane *Methyl derivative* $C_6H_4(OH)(OMe)CH(C_6H_4NMe_2)_2$ [136°] Formed from vanillin, $PhNMe_2$, and $ZnCl_2$ (O Fischer a Schmidt, *B* 17, 1895) Crystals, v sol alcohol **OXY TETRA METHYL-AMMONIUM HYDROXIDE** $CH_2(OH)NMe_2OH$ Formed from CH_3NMe_2 and moist Ag_2O (Hofmann, *J* 1859, 377) Yields $(CH_2OH)NMe_2Cl$, $PtCl_4$ crystallising in octahedra

DI OXY METHYL AMYL KETONE *Di-methyl derivative* $CH(OMe)_2COCH_2Et$ (134°) *SG* 15 886 Formed as one of the products of the action of $NaOMe$ upon $CHCl_3COCH_2Et$, CO_2Et (James, *A* 231, 243, *C J* 49, 57) Oil Miscible with alcohol and with ether, burns with pale flame Does not combine with $NaHSO_4$ or react with AcO

OXY-METHYL AMYL PYRROLE CARB OXYLIC ETHER $C_6H_4N(CMe_2)COCH_2Et$ [52°] $COCH_2Et > COCH_2Et$

(188° at 16 mm) Formed from acetosuccinic ether, amylamine, and alcohol in the cold (Emery, *A* 260, 150) White plates (from CS_2)

DI OXY DI METHYL ANILINE v TETRA METHYL DI AMIDO DI PHENYL DI-OXIDE

OXY METHYL ANTHRANOL *Acetyl derivative* $C_6H_4 \begin{smallmatrix} COAc \\ \diagup \quad \diagdown \\ CO \end{smallmatrix} > C_6H_4Me$ [217°] Formed from methyl anthraquinone [177°], Ac_2O , $NaOAc$, and zinc-dust (Liebmann, *B* 21, 1172) Plates

OXY METHYL-ANTHRAQUINONE $C_{14}H_8O_3$ *4e* $C_6H_4 \begin{smallmatrix} CO \\ \diagup \quad \diagdown \\ CO \end{smallmatrix} > C_6H_4Me(OH)$ $\begin{bmatrix} 1 & 4 & 3 \\ 6 & & \end{bmatrix}$

[262°] Formed by heating *o* cresol phthalein or a mixture of phthalic anhydride, *o* cresol, and H_2SO_4 at 160° (Baeyer a Fraude, *B* 12, 241, *A* 202, 163) Yellow leaflets

Bromo- derivative [205°]

Oxy methyl-anthraquinone

$C_6H_4C_6O_2C_6H_4Me(OH)$ [1625] [170°] Formed in like manner from *p* cresol (Drewson, *A* 212, 346, Bruckoff, *B* 20, 2069) Orange needles (by sublimation)

Acetyl derivative [180°] Needles

Oxy methyl anthraquinone $C_{14}H_8Me_2O_3(OH)$ [178°] Formed from amido methyl anthraquinone [202°] by the diazo reaction (Römer a Lank, *B* 16, 699) Yellow needles (by sublimation)

Acetyl derivative [177°] Plates

Di oxy methyl anthraquinone

$C_6H_4(OH)C_6O_2C_6H_4Me(OH)$ [162°] Mol w 254 *S* (boiling 86 pc alcohol) 45 The yellow colouring matter of rhubarb, the wall lichen (*Parmelia parietina*), and of the root of *Rumex obtusifolius* (Rochleder a Heldt, *A* 48, 12, Döpping a Schlossberger, *A* 50, 215, De la Rue a Muller, *C J* 10, 298, Thann, *A* 107, 324) It occurs also in the root of *Rheum pyramidalis*, *Rumex palustris*, and of other varieties of *Rumex* (Grothe, *P* 113, 190) and in senna leaves (Batk, *C C* 1864, 622) Formed by passing air through an alkaline solution of chrysarobin (Liebmann a Seidler, *A* 212, 36) Golden plates (from alcohol), forming a cherry red solution in $NaOH$ aq Insol Na_2CO_3 aq Conc H_2SO_4 forms a red solution Conc HNO_3 forms a tetra nitro derivative Yields methyl anthracene on distilling with zinc dust Does not dye mordants Zinc, $HOAc$, and HCl aq form $C_{14}H_8O_3$ [200°-206°] whence $C_{14}H_8O_3Ac_2O$ [231°] (Liebmann, *B* 21, 437)

Di acetyl derivative [200°] Plates

Di benzoyl derivative [c 201°]

Di oxy methyl anthraquinone

$C_6H_4C_6O_2C_6HMe(OH)_2$ [16325] *Methyl-quinizarin* [160°] Formed from hydrotoluquinone, phthalic anhydride, and H_2SO_4 at 140° (Nietzki, *B* 10, 2011) Red needles (from alcohol) Yields methyl-anthracene when distilled with zinc dust

Acetyl derivative [185°] Needles

Di oxy methyl anthraquinone

$C_6H_4C_6O_2C_6HMe(OH)_2$ [16432] *Methyl-alizarin* [252°] Formed by potash fusion from bromo or oxy methyl anthraquinone (O Fischer, *B* 8, 675, Fraude, *B* 12, 241) Orange needles, which may be sublimed Dyes mordants like alizarin

Alkannin (vol 1 p 125) is probably a di oxy-methyl-anthraquinone as it yields methyl-anthracene on distillation with zinc dust (Liebmann a Römer, *B* 20, 2428)

Di-oxy-di-methyl-anthraquinone

$C_6H_4Me(OH)C_6O_2C_6H_4Me(OH)$ *Di-methyl-anthraruflin* [300°] Got, together with the two following isomerides, by the action of H_2SO_4 on *s*-oxy-toluic acid (Kostanecki a Nismentowski, *B* 18, 256, 2140, *A* 240, 376) Yellow needles

(from benzene) Its alkaline solutions are yellow Does not dye mordants

Di-acetyl derivative [237°] Tables

Di-oxy-di-methyl-anthraquinone

$C_{12}H_8Me_2(OH)_2O_2$, *Di-methyl-anthraflavic acid* (by needles or small yellow plates (by sublimation), not solid at 360° Does not dye mordants

Di-acetyl derivative [223°] Needles

Di-oxy-di-methyl anthraquinone (benz) [213°]. Yellow needles, yielding $C_{12}H_{10}Ac_2O$, [188°]

Tri-oxy-methyl-anthraquinone $C_{12}H_{10}O_3$

Emodin [254°] Occurs in rhubarb root (Warren de la Rue a Hugo Müller, *C J* 10, 304), in the bark and berries of *Rhamnus frangula* (Liebermann a Waldstein, *B* 8, 970, 9, 1775, Schwabe, *Ar Ph* [3] 26, 569), and in the lichen *Nephroma Lusitanica* (Bachmann, *C* 8 1888, 47) Orange red monoclinc prisms (containing aq) Yields methyl anthracene on distilling with zinc dust Its alkaline solution is dark cherry red

Mono-acetyl derivative [180°]

Tri-acetyl derivative [190°]

Tri oxy methyl anthraquinone

$C_{12}H_8Me_2C_2O_2C_6H_5(OH)_3$ [1 6 2 3 4] *Methyl-anthragallol* [c 375°] Made by heating gallic acid with *p* toluic acid at 130° for 15 hours (Cahn, *B* 19, 2335, *A* 240, 284) Orange red needles (by sublimation) Its solution in conc KOHAq is green, becoming violet on dilution Hot NH_3 aq forms a blue solution Conc H_2SO_4 forms a red liquid turned green by a trace of HNO_3 , decolourised by more HNO_3

Tri-acetyl derivative [204°]

Tri oxy methyl anthraquinone

[2' 1' 6'] $C_{12}H_8Me_2C_2O_2C_6H_5(OH)_3$ [1 6 2 3 4] *Methyl anthragallol* [298°] Formed by heating gallic acid with *o* toluic acid (C) Minute yellow needles Forms a green solution in conc KOHAq turned violet on dilution Dyes like anthragallol Conc H_2SO_4 gives a red solution turned green by HNO_3

Tri-acetyl derivative [210°] Tables

(2, 3, 4) **Tri oxy 3' and 5' methyl anthraquinones (methyl anthragallols)** These two isomerides are formed simultaneously by heating gallic acid with *m* toluic acid at 130°-135° The one melts at [312°] and gives an acetyl derivative which forms needles melting at [190°] The other isomeride melts at [235°-240°], and its acetyl derivative, which crystallises in small prisms, melts at 218° Their other properties are almost the same as those of the other methyl-anthragallols (Cahn, *B* 19, 2336)

Tri-oxy-methyl anthraquinone

[3 2 1 6] $C_6H_2(OH)_3C_2O_2C_6H_5Me(OH)$ [1' 6' 2' 5'] *Methyl-oxy alizarin* Formed by saponifying its di-methyl ether which is produced by heating hemipic acid with *p* cresol and H_2SO_4 (Liebermann a Kostanecki, *A* 240, 303) Brownish-yellow flakes, v sol alcohol Dyes like alizarin

Di-methyl ether $C_{12}H_{10}O_2$ Flakes

Tri-oxy-di-methyl-anthraquinone

[4' 2' 1' 6'] $C_{12}H_8Me_2C_2O_2CH(OH)_3$ [1 6 2 3 4] Formed by heating (4,2,1)-di-methyl benzoic acid with gallic acid and H_2SO_4 (Birukoff, *B* 20, 871, *A* 240, 287) Yellowish-red needles Forms a red solution in conc H_2SO_4 Yields di-methyl anthracene [224°], when distilled with zinc-dust

Tri-oxy-tri-methyl-anthraquinone

[4' 3' 2' 1' 6'] $C_{12}H_8Me_3C_2O_2C_6H_5(OH)_3$ [1 6 2 3 4] *Tri-methyl anthragallol* [244°] Formed from tri methyl benzoic (durylo) acid, gallic acid, and H_2SO_4 (Wende, *B* 20, 867) Brown needles Yields tri methyl anthracene [236°]

Acetyl derivative [174°] Plates

Tetra-oxy-di-methyl-anthraquinone

[2' 4' 5' 1' 6'] $C_{12}H_8Me_2(OH)_2C_2O_2C_6H_5Me(OH)_3$ [1 6 2 3 4] *Di methyl anthrachrysone* [above 360°] Formed by heating di oxy-*o* toluic acid with H_2SO_4 (10 pts) at 100° (Cahn, *B* 19, 755) Reddish yellow plates (by sublimation) Does not dye mordants

Tetra acetyl derivative [234°]

OXY-METHYL-BENZOIC ACID v OXY TOLUIC ACID

Oxy di methyl benzoic acid

$C_6H_8Me_2(OH)CO_2H$ [6 3 2 1] *Oxy isoxyllylic acid* [142°] Formed by fusing ethyl *p* xylene sulphonic acid with potash (Stahl, *B* 23, 991) Needles Gives a bluish violet colour with $FeCl_3$

Oxy-di methyl benzoic acid

$C_6H_8Me_2(OH)CO_2H$ [5 4 2 1] *Oxy p xylylic acid* [199°] Formed from ψ cumenol by potash fusion (Reuter, *B* 11, 30, Jacobsen, *B* 12, 436) Needles, volatile with steam Gives a bluish violet colour with $FeCl_3$, HCl aq at 220° gives xylene [61°] —BaA', *S* 11 at 0°

Oxy di methyl-benzoic acid

$C_6H_8Me_2(OH)CO_2H$ [4 2 5 1] [170 5°] Formed by fusing $C_6H_5Me_2BrCO_2H$ (Gunter, *B* 17, 1608) Scarcely volatile with steam Gives no colour with $FeCl_3$

Oxy di methyl benzoic acid [137°] Formed

from *p* xylene, Na, and CO_2 at 180° (Oliver, *G* 12, 166) Needles, coloured violet by $FeCl_3$, —BaA', 4aq

Oxy-di methyl benzoic acid

$C_6H_8Me_2(OH)CO_2H$ [6 3 4 1] [153°] Got from $C_6H_5Me_2BrCO_2H$ and KOH Not coloured blue by $FeCl_3$

Oxy di methyl benzoic acid *Xyletic acid*

[155°] Formed from crude xylene, Na, and CO (Wroblewski, *Z* 1868, 233) —CaA', 2aq —BaA', aq needles

Di oxy di methyl-benzoic acid

$C_6H_8Me_2(OH)_2CO_2H$ [5 3 6 2 1] [196°] Formed from betorcin, $NaHCO_3$, and water at 130° (Kostanecki, *B* 19, 2323) Prisms Coloured blue by $FeCl_3$

Oxy tri-methyl-benzoic acid

$C_6H_8Me_3(OH)CO_2H$ [6 4 3 2 1] *Oxydurylic acid* [148°] Formed by fusing durenol with potash (Jacobsen a Schnappauff, *B* 18, 2844) Needles —CaA', aq prisms, m sol water

Di oxy-tetra methyl benzoic acid *Tetra*

hydride of the methylene derivative $CH_2O_2C_6H_4Me_2CO_2H$ *Piperhydraemic acid* [96°] Formed from (β)-hydropiperic acid and sodium amalgam (Buri, *A* 216, 178) —CaA', aq crystals (from water)

OXY TRI-METHYL-BENZOIC ALDEHYDE

$C_6H_8Me_3(OH)CHO$ [1 3 4 6 5] [106°] Formed by the action of chloroform on an alkaline solution of ψ cumenol (Auwers, *B* 17, 2976) Needles $FeCl_3$ gives a blue colour

OXY-METHYL-BUTYL-PYRROLE CARB-

OXYLIO ETHER $C_6H_5N < \begin{smallmatrix} OMe \\ CO \end{smallmatrix} (CO_2Et)$ [68°].

(175° at 15 mm) Formed from aceto succinic ether (10g) and isobutylamine (8.4g) in the cold (Emery, *A* 360, 149) Needles (from CS₂)

OXY METHYL CINNAMIC ACID *Anhydride* $\left[64\frac{1}{2}\right] \text{C}_6\text{H}_5\text{Me}(\text{OH}) \begin{smallmatrix} \text{CH CH} \\ \text{O CO} \end{smallmatrix} \quad [248^\circ]$

Hemoubelliferon Formed by heating orcin with malic acid and H₂SO₄ (Fechmann & Welsh, *B* 17, 1649) Tables, sol alcohol and aqueous alkalis Conc H₂SO₄ forms a solution with blue fluorescence Potash fusion gives orcyal aldehyde and KOAc

Acetyl derivative [127°] Formed by heating orcyal aldehyde with Ac₂O and NaOAc (Tiemann & Helkenberg, *B* 12, 1002) Needles

OXY METHYL COUMARILIC ACID C₁₀H₈O₄, $\left[4\frac{1}{2}\right] \text{C}_6\text{H}_3(\text{OH}) \begin{smallmatrix} \text{CMe} \\ \text{O} \end{smallmatrix} \text{C CO}_2\text{H} \quad [226^\circ]$

Formed by saponification of its ethyl ether, which is obtained by boiling chloroacetic ether (1 mol) with a conc alcoholic solution of mono sodium resorcin C₆H₃(OH)(ONa) Needles (containing $\frac{3}{4}$ aq) V sol hot water On distillation it loses carbonic acid, giving oxy methyl coumarone C₈H₆(OH) $\begin{smallmatrix} \text{CMe} \\ \text{O} \end{smallmatrix} \text{CH} \quad [97^\circ]$

Ethylether A'Et [178°], white needles, v sol ether, its dilute solutions have a blue fluorescence (Hantzsch, *B* 19, 2928)

Di-oxy-methyl-coumarilic acid C₁₀H₈O₄, $\left[64\frac{1}{2}\right] \text{C}_6\text{H}_3(\text{OH})_2 \begin{smallmatrix} \text{CMe} \\ \text{O} \end{smallmatrix} \text{C CO}_2\text{H} \quad [281^\circ]$

Formed by saponification of its ethyl ether, which is obtained by boiling chloroacetic ether (1 mol) with a conc alcoholic solution of mono sodium phloroglucin C₆H₃(OH)(ONa) (1 mol) Crystals (containing $\frac{1}{2}$ aq) The acid and its ether give an indigo blue colour with warm conc H₂SO₄

Ethyl ether A'Et [242°], small white needles, the alkaline solution is fluorescent (Lang, *B* 19, 2934)

OXY METHYL COUMARONE v COUMARONE

OXYMETHYLENE v FORMIC ALDEHYDE

OXY-TRIMETHYLENE-DIAMINE

CH(OH)(CH₂NH₂)₂ Formed by heating with HClAq the compound got by the action of epichlorhydrin or (a) dichlorhydrin on potassium phthalimide (Goedeckemeyer, *B* 21, 2689, Gabriel, *B* 22, 225) —B''H₂Cl₂ [184°] Hygroscopic mass —B''H₂PlCl₂ [240°] —B''H₂Br₂ [200°] Needles —B''C₆H₅(NO₂)₂OH yellow needles [230°]

OXY TRIMETHYLENE DIPHTHALAMIC ACID CH(OH)(CH₂NH COC₆H₄CO₂H)₂ [c 120°] Formed by boiling oxytrimethylene diphthalamide with potash (Goedeckemeyer, *B* 21, 2690) Hygroscopic needles Yields oxytrimethylene diamine on evaporating with HClAq

OXYMETHYLENE-PHTHALIDE? C₈H₆O₄, [146°] A crystalline body formed in the action of Br and HOAc on acetophenone carboxylic acid (Gabriel & Michael, *B* 11, 1010)

Phenyl derivative C₈H₅ $\begin{smallmatrix} \text{CH OPh} \\ \text{CO O} \end{smallmatrix}$

[144°] Made by heating phthalic anhydride with phenoxy acetic acid and NaOAc (Gabriel, *B* 14, 922) Needles The homologous *p*-tolyl derivative [174°] is also crystalline

OXYTRIMETHYLENE - DIPHTHALIMIDE

CH(OH)(CH₂NC₆H₄O)₂ [205°] Formed from potassium phthalimide and (a) dichlorhydrin at 150° (Goedeckemeyer, *B* 21, 2689, Gabriel, *B* 22, 224) Needles (from HOAc) Fuming HBr at 200° converts it into CHBr(CH₂NH₂)₂

OXY - METHYL - ETHYL - BENZOIC ACID

C₆H₅MeEt(OH)CO₂H [149°] Made by fusing *s* di methyl ethyl benzene sulphonic acid with potash (Jacobsen, *A* 195, 284) Long needles from dilute alcohol Turned blue by FeCl₃

OXY METHYL-ETHYL KETONE *Ethyl derivative* CH₃(OEt)COEt (100°-105°)

(Formed from CH₃(OEt)COCHMeCO₂Et and alcoholic KOH at 120° (Isbert, *A* 234, 196)

OXY - METHYL - ETHYL - PYRIMIDINE

C₈H₈N₂O $\begin{smallmatrix} \text{N CMe} \\ \text{N C(OH)} \end{smallmatrix} \text{CH} \quad [160^\circ]$

Formed by the action of a 10 p.c. solution of NaOH (1 mol) upon a mixture of propionamide hydrochloride (1 mol) and acetic ether (1 mol) (Pinner, *B* 18, 2847, *B* 22, 1619) Fine white needles V sol water and alcohol Reduced by zinc dust to methyl ethyl pyrimidine

Salts —B''HCl [240°-246°], very soluble thick prisms —B''H₂Cl₂PlCl₂ [236°], thick yellow prisms

Oxy methyl di ethyl pyrimidine

$\begin{smallmatrix} \text{N CMe} \\ \text{N C(OH)} \end{smallmatrix} \text{CET} \quad [135^\circ]$ Formed from propionamide hydrochloride, ethyl acetoacetic ether, and dilute (10 p.c.) NaOH aq (Pinner, *B* 22, 1621) Needles, v sol water

Oxy methyl di ethyl pyrimidine C₈H₈N₂O

$\begin{smallmatrix} \text{N CET} \\ \text{N C(OH)} \end{smallmatrix} \text{CMe} \quad [155^\circ]$ Formed from propionamide hydrochloride, propionyl propionic ether, and KOH (Meyer, *J pr* [2] 39, 264) When warmed with a solution of nitrous acid in HOAc it yields C₈H₈N₄O₂ and a little C₈H₈N₂O₂ [205°]

Oxy di methyl ethyl pyrimidine

$\begin{smallmatrix} \text{N CMe} \\ \text{N C(OH)} \end{smallmatrix} \text{CET} \quad [146^\circ]$ Formed from acetamide and ethyl acetoacetic ether (Pinner, *B* 22, 1618) Needles, v sol water

Oxy di methyl-ethyl pyrimidine

$\begin{smallmatrix} \text{N CMe} \\ \text{N C(OH)} \end{smallmatrix} \text{CMe} \quad [165^\circ]$ Formed from propionamide and methyl acetoacetic ether (P) Needles, v sol water and alcohol

Oxy di methyl-ethyl pyrimidine

$\begin{smallmatrix} \text{N CET} \\ \text{N C(OH)} \end{smallmatrix} \text{CMe} \quad [168^\circ]$ Formed from acetamide and propionyl propionic ether (E von Meyer, *J pr* [2] 40, 304)

Di-oxy-methyl-ethyl-pyrimidine

C₈H₈MeEt(OH)N₂ $\begin{smallmatrix} \text{CO} \text{NH CMe} \\ \text{Net CO} \end{smallmatrix} \text{CH}$

Methyl ethyl uracil [195°] Formed, together with di-oxy methyl di ethyl pyrimidine [58°], by heating potassium methyl uracil with EtBr at 155° (Hoffmann, *A* 253, 68) Prisms (from EtBr) or needles (from alcohol)

OXY-METHYL-ETHYL-PYRROLE CARB

OXYLIC ETHER $\begin{smallmatrix} \text{Net CMe} \\ \text{CO CH} \end{smallmatrix} \text{C CO}_2\text{Et} \quad [76^\circ]$

(165° at 14 mm) Formed from acetyl-succinic ether and cold aqueous ethylamine (Emery, *A* 260, 148) Needles

Oxy-methyl-ethyl-pyrrole dihydride carb-oxylie acid $\text{CO NEt} \begin{array}{c} \text{CH}_2 \\ \text{CH}_2 \end{array} \text{CMe CO}_2\text{H}$ [123°] Obtained, as nitrile, by heating acetyl propionic (levulic) ether with HCl and alcoholic ethyl amine (Kühling, *B* 23, 709) Needles, v sol water and alcohol

Nitrile $\text{C}_6\text{H}_5\text{N}_2\text{O}$ Syrup

Amide $\text{C}_6\text{H}_5\text{N}_2\text{O}_2$ [183°] Needles

Thioamide [176°] Formed from the nitrile and alcoholic H_2S

Amidoxim [c 160°] Formed from the thioamide by boiling with hydroxylamine hydrochloride

OXY-(B 2 Py 2)-DI-METHYL-(Py 3)-ETHYL QUINOLINE $\text{C}_6\text{H}_4(\text{OH})\text{N}$ [45°] (312°-316° at 720 mm) Colourless crystals Formed by fusing di-methyl ethyl quinoline sulphonic acid with KOH—A'K* easily soluble colourless plates (Harz, *B* 18, 3390)

Di-oxy-methyl ethyl quinoline

$\text{C}_6\text{H}_4\text{Me} \begin{array}{c} \text{C(OH) CEt} \\ \text{N}=\text{C OH} \end{array}$ [c 220°] Formed from chloro ethyl oxy o toluquinoline and dilute HCl at 165° (Rugheimer a Schramm, *B* 20, 1235, 21, 302) Needles (from HOAc), insol water

Reference — CHLORO OXY METHYL ETHYL-QUINOLINE

α-OXY-α-METHYL GLUTARIC ACID

$\text{C}_6\text{H}_4\text{O}_2$ *ie* $\text{CO}_2\text{H CMe(OH) CH}_2\text{CH}_2\text{CO}_2\text{H}$ Formed from the lactone of oxy isohexoxic acid by oxidation with dilute nitric acid (Fittig a Bredt, *A* 208, 63, 236, 225) Made also by saponification of the nitrile formed by combination of β acetyl-propionic (levulic) acid with HCl (Kreckeler a Tollens, *B* 18, 2018, *A* 238, 287) The free acid at once changes to the lactonic acid BaA'' 4aq — SrA'' 4aq — CaA'' 7aq — ZnA'' — AgA'' white flocculent pp

Lactonic acid $\text{CMe(CO}_2\text{H)} \begin{array}{c} \text{CH}_2\text{CH}_2 \\ \text{O}=\text{CO} \end{array}$ [70°] Deliquescent prisms, v sol ether Converted by hot H_2SO_4 into CO_2 and levulic acid CaA'' 44aq — MeA'' (252°) — EtA'' (262°)

Amide of the lactonic acid [c 124°]

Nitrile of the lactonic acid [c 33°]

β-Oxy β methyl-glutaric acid

$\text{CMe(OH)(CH}_2\text{CO}_2\text{H)}_2$ Formed by oxidising methyl-di ethyl carbinol with KMnO_4 (Sorokin, *J pr* [2] 23, 278) — CaA'' amorphous — ZnA'' (dried at 110°) — CuA'' (OH)₂ 5aq small tablets AgA''

Oxy-tri methyl glutaric acid Lactonic acid $\text{CMe}_2 \begin{array}{c} \text{CH}_2\text{CMe CO}_2\text{H} \\ \text{CO O} \end{array}$ [104°] Formed

from a bromo tri methyl glutaric anhydride and KOHAq at 0° (Auwers a V Meyer, *B* 23, 307) Crystals, v sol water — AgA'' crystalline powder

Di-oxy-di-methyl glutaric acid?

$\text{CH}_2(\text{CMe(OH)CO}_2\text{H})_2$ [96°] Formed by dissolving di a bromo-di a methyl glutaric anhydride in cold 10 p.c caustic soda solution (Auwers a Jackson, *B* 23, 1614) Six sided plates, v sol cold water

OXY-METHYL-INDONAPHTHENE

$\text{C}_6\text{H}_4\text{Me} \begin{array}{c} \text{CH}_2 \\ \text{C(OH)} \end{array} \text{CH}$ [59°] Formed by condensing *m* tolyl propionic acid by means of hot H_2SO_4 (Von Miller, *B* 23, 1899) White needles with characteristic odour Volatile with steam

The isomeride from *p* tolyl propionic acid melts at 63°

Oxy methyl-indonaphthene

$\text{C}_6\text{H}_4 \begin{array}{c} \text{CH}_2 \\ \text{C(OH)} \end{array} \text{CMe}$ (246° at 719 mm).

Formed from β phenyl isobutyric acid and H_2SO_4 at 150° (Von Miller, *B* 23, 1888) Heavy oil smelling of peppermint, volatile with steam Yields phthalic acid on oxidation Gives a phenyl hydrazide [116°]

OXY METHYL INDOLE Ethyl derivative

$\text{C}_6\text{H}_4 \begin{array}{c} \text{NH} \\ \text{C(OEt)} \end{array} \text{CMe}$ or $\text{C}_6\text{H}_4 \begin{array}{c} \text{NH} \\ \text{CH} \end{array} \text{C CH}_2\text{OEt}$

[142°] Formed by heating at 90° the phenyl-hydrazide of $\text{CH}_2\text{CO CH}_2\text{OEt}$ (Fittig, *B* 21, 2649) Crystals (from alcohol)

OXY-METHYL-JULOLE DIHYDRIDE

$\text{CH}_2 \begin{array}{c} \text{CH}_2\text{CH}_2 \\ \text{CH} \end{array} \begin{array}{c} \text{N CO} \\ \text{C}=\text{C} \\ \text{CH CH} \end{array} \text{CMe} \text{CH}$ [130° cor]

Formed by boiling quinoline tetrahydride with acetoacetic ether (Reissert, *B* 24, 845) White needles, v sol alcohol, insol alkalis, sol conc HClAq — B'HCl 14aq — B'_2H₂PtCl₄ orange needles

OXY METHYL MALONAMIC ACID

$\text{C}_6\text{H}_4\text{NO}_2$ *ie* $\text{CMe(OH)(CONH}_2\text{)(CO}_2\text{H)}$ Formed from pyruvic acid and KCl by heating with conc HClAq (Böttger, *B* 14, 88) Syrup — ZnA'' 2aq crystalline, v sol water

DI OXY DI METHYL MALONIC ACID Di methyl derivative $(\text{CH}_3\text{O CH}_2)_2\text{C(CO H)}$ [138°] Formed from sodium malonic ether, chloro di methyl oxide and NaOEt (Kleber, *A* 246, 111) Prisms, v sol water Its salts are very hygroscopic

Di ethylether $\text{C}_6\text{H}_4\text{O}_2(\text{CO}_2\text{Et})_2$ (c 240°)

OXY-DI-METHYL PENTAMETHENYL

HYDRIDE $\text{CO} \begin{array}{c} \text{CH CMe} \\ \text{CH}_2\text{CHMe} \end{array}$ or

$\text{CO} \begin{array}{c} \text{CH}=\text{CMe} \\ \text{CHMe CH}_2 \end{array}$ 'Di methyl ketopentene' (119°) Made from methylulvic acid by prolonged boiling (Dietzel, *A* 250, 195) Oil, lighter than water

Oxy methyl heptamethenyl trihydride?

$\text{C}_6\text{H}_4\text{O}_2$ *ie* $\text{CH} \begin{array}{c} \text{CMe CH}_2\text{CH}_2 \\ \text{CO CH}_2\text{CH}_2 \end{array}$ (189°) An oil, smelling like camphor, formed by heating its dicarboxylic ether which is formed by distilling $\text{CO}_2\text{Et CHAc CH}_2\text{CH}_2\text{CHAc CO}_2\text{Et}$ (Perkin a Obrembsky, *B* 19, 2051) Reacts with phenyl hydrazine

DI-OXY-DI-METHYL-HEPTAMETHYLENE

$\text{C}_6\text{H}_4\text{O}_2$ *ie* $\text{CH}_2 \begin{array}{c} \text{CH}_2\text{CH}_2\text{CMe(OH)} \\ \text{CH}_2\text{CH}_2\text{CMe(OH)} \end{array}$ (201° at 180 mm) Formed by reducing

$\text{CH}_2(\text{CH}_2\text{CH}_2\text{Ac})_2$ with sodium (Kipping a W H Perkin, jun, *C J* 59, 220) Thick oil with burning taste, smelling of thyme Does not form an oxim or a phenyl hydrazide Yields deliquescent $\text{C}_6\text{H}_4\text{NaO}$ aq Forms an oily product of condensation $\text{C}_{16}\text{H}_{24}\text{O}_2$ (305°-310° at 200 mm)

Di-acetyl derivative $\text{C}_6\text{H}_4\text{AcO}_2$ (201° at 65 mm) Liquid, insol cold water

OXY METHYL-(α)-NAPHTHOQUINOLINE

$\text{C}_{10}\text{H}_7\text{NO}$ *ie* $\text{C}_{10}\text{H}_6 \begin{array}{c} \text{C(OH) CH} \\ \text{N}=\text{CMe} \end{array}$ [292°] (K), [over 800°] (C & L) Formed by boiling with

HCl a naphthyl β imido-butyric ether, which is formed by condensation of acetoacetic ether with (a)-naphthylamine (Knorr, *B* 17, 543, Conrad a. Lumpach, *B* 21, 631) Needles (from alcohol) —B'H₂PtCl₆ v sl sol hot water

The isomeride derived from (8) naphthylamine crystallises in flat needles [286°] (K.) or [above 300°] (C & L) decomposed by distillation

OXY-DI-METHYL (8)-NAPHTHOQUINOLINE DISULPHONIC ACID

C₁₂H₁₁(OH)(SO₃H)N 1 $\frac{1}{2}$ aq Formed by potash fusion from the disulphonic acid of di methyl-(8) naphthoquinoline (Reed, *J pr* [2] 35, 309) small needles, insol water and alcohol

DI-OXY-METHYL-NAPHTHOQUINONE (?)
C₁₁H₈O₂ [170°] Extracted from tubers of *Drosera Whitakeri* (Rennie, *C J* 51, 376) Forms a deep red solution in KOHAq

Tri oxy methyl-naphthoquinone (?) C₁₁H₆O₃ [c 193°] A substance of this empirical composition may be extracted from tubers of *Drosera Whitakeri* (Rennie, *C J* 51, 371) Red plates (from alcohol or glacial HOAc) v sl sol water Gives a violet solution in KOHAq or NH₄Aq Reduced by SnCl₂ to C₁₁H₈O₂ [217°]

OXY-METHYL-ISO OXAZOLE

CMe < CH₂ CO
N—O [170°] Formed from acetoacetic ether, hydroxylamine hydrochloride, and NaOH at 50° (Hantzsch, *B* 24, 497) Satiny needles, sl sol cold water, sol HClAq —Ac O forms on heating a compound [136°] possibly N—O > C < CMe N —NH₄A' [207°] —CMe CH > C < C(OAc) O —BaA' 2aq —CuA'. —AgA' gelatinous

Di oxy di methyl di iso oxazyl
Q CO < CH CH < CO Q
N CMe > CH CH < CMe N Formed by warming isocarbopvrotitanic ether with hydroxylamine and HOAc on the water bath (Knorr, *B* 22, 161) Hair like needles which explode at 190°

OXY-TETRA-METHYL-PHENYL ACETIC ACID [6 5 3 2 1] C₆HMe, CH(OH) CO H [146°] Formed from tetra methyl phenyl glyoxylic acid by reduction with sodium amalgam (Claus a. Foecking, *B* 20, 3102) Nodules —BaA', 2aq —CaA', 8aq needles

Oxy tetra-methyl-phenyl acetic acid
[6 5 4 3 1] C₆HMe, CH(OH) CO₂H [160°] Formed in like manner (Claus, *J pr* [2] 38, 283) Hexahedra, m sol hot water

Oxy tetra methyl phenyl acetic acid
[6 4 3 2 1] C₆HMe, CH(OH) CO₂H [156°] Prisms, v sol alcohol and ether —NaA' 1 $\frac{1}{2}$ aq —CaA' 8aq —BaA', 3aq small crystals

OXY-TRI-METHYL-PHENYL METHYL-PYRAZOLE C₆H₃N₂O

C₆H₃Me, N < CO CH₂
N=CMe [155°] Formed by heating the ψ cumyl hydrazide of acetoacetic ether at 140° for two hours (Haller, *B* 18, 706) Glistening crystals, v sol alcohol Yields a nitrosamine C₆H₃N₂O₂ [156°] and a methyl derivative C₆H₃N₂O [166°]

OXY-TETRA-METHYL-PHENYL-DI-METHYL-PYRIDINE CARBOXYLIC ACID

C₆HMe, N < CMe C(CO₂H)
CMe CH CO > [145°] Made by

heating tetramethyl phenyl amido crotonic ether at 280° (Conrad a. Lumpach, *B* 21, 1655)

OXY-METHYL-ISOPHTHALIC ACID
C₆H₄Me(OH)(CO₂H)₂ [4 5 3 1] [c 270°] Formed by heating C₆H₄Me(SO₃H)(CO₂H)₂ with conc HClAq at 220° (Jacobsen, *B* 14, 2115) Needles (from hot water) Not coloured by FeCl₃

OXY-DI-METHYL-PROPIONAMIDINE
CH, CH(OH) C(NHMe) NMe The crystalline hydrochloride B'HCl [215°], formed by the action of methylamine on the hydrochloride of CH, CH(OH) C(OC₂H₅) NH, is v sol water and m sol alcohol (Pinner, *B* 23, 2948)

OXY-METHYL-PROPYL-BENZOIC ACID
Methyl derivative C₆H₄MePr(OMe)CO₂H [3 6 1 4] [139°] Formed from its amide [149°], which is made by the action of ClCONH₂ on C₆H₄MePr(OMe) in CS₂ in presence of AlCl₃ (Gattermann, *A* 244, 68) The ethyl derivative [159°] and its amide C₆H₄MePr(OEt) CONH₂ [127°] are got in like manner

Oxy-methyl isopropyl-benzoic acid
C₆H₄MePr(OH)CO₂H [3 5 2 1] *Cymenotic acid* [147°] Formed from (4,2,1) cymenol, sodium, and CO₂ (Jesurun, *B* 19, 1414) Slender needles (from water) —BaA', 4aq —AgA' —MeA' [148°]

Isomeric acids v CARVACROTIC ACID and THYMOTIC ACID

OXY-METHYL-PROPYL-CINNAMIC ACID
[1 4 3 6] C₆H₄(CH₂)₃(C₆H₄)₂(OH) C₆H₄ CO₂H *Thymo acrylic acid* [280°] Prepared by heating *p* thymotic aldehyde with acetic anhydride and sodium acetate (Kobek, *B* 16, 2104) White microscopic crystals

Methyl derivative
C₆H₄(CH₂)₃(C₆H₄)₂(OMe) C₆H₄ CO₂H [141°] Needles, v sol alcohol, sl sol water

OXY-METHYL-PROPYL-CINNAMIC ANHYDRIDE v METHYL-PROPYL-COUMARIN

OXY-METHYL-PROPYL-KETONE *Ethyl ether* EtO CH₂ CO Pr (112°-115°) Formed from EtO CH₂ CO CHET CO Et and alcoholic KOH at 120° (Isbert, *A* 234, 195) Lighter than water, smells like a ketone

DI-OXY-METHYL-ISOPROPYL-PYRIMIDINE CMe₂(OH) C < N CMe > CH [98°]

Formed from oxy-isobutyramidine hydrochloride, acetoacetic ether, and NaOH in equivalent proportions (Pinner, *B* 22, 2625) Needles (from acetone), v sol most solvents

OXY-METHYL-PROPYL-PYRROLE CARBOXYLIC ETHER C₆H₄N—CMe > CO CO₂Et [50°]

(172° at 15 mm) Formed from aceto succinic ether and propylamine in the cold (Emery, *A* 260, 148) Needles

OXY-METHYL-PURIN C₆H₂Me(OH)N₄ a.s. CO < NH—C CH N > CH [233°] Made by heating di chloro oxy methyl purin with HI (Fischer, *B* 17, 332) Prisms, v sol water Alkaline in reaction —B'HI —B'H₂PtCl₆

Oxy-di-methyl-purin C₆H₄Me(OMe)N₄ [112°] Formed from di chloro oxy di methyl-purin and HIAq (Fischer, *B* 17, 334) Slender needles, v sol water, insol alkalis

Di-oxy-di methyl purin C₆H₂N₄O₂ Formed by reducing the ethyl derivative of chloro-di-oxy di methyl purin with HI Sparingly soluble crystals

Di-ethyl derivative $C_{14}H_{18}N_4O_4$ ϵ
 $G(OEt) \begin{smallmatrix} N \\ \diagup \end{smallmatrix} C(OEt) \begin{smallmatrix} C \\ \diagdown \end{smallmatrix} NMe \begin{smallmatrix} CO \\ \diagup \end{smallmatrix} ? [127^\circ]$ Formed from di-chloro oxy-di methyl purin and alcoholic NaOH. Fine plates, sol HCl aq, insol alkalis.

Tri-oxy-methyl-purin v METHYL URIC ACID

OXY-METHYL-PYRAZOLE $C_4H_6N_2O$ ϵ

$CMc \begin{smallmatrix} CH_2 \\ \diagup \end{smallmatrix} CO \\ \diagdown \end{smallmatrix} NH$ [215°] Formed by warming acetoacetic ether with hydrazine and water (Curtius, *J pr* [2] 39, 52). Prisms (from water), sl sol hot alcohol. Has a sweet taste.

OXY-DI-METHYL-PYRIDINE *Ethyl derivative* $C_8H_{10}NO$ ϵ $N \begin{smallmatrix} CMc \\ \diagup \end{smallmatrix} CH \\ \diagdown \end{smallmatrix} COEt$ CH $\begin{smallmatrix} CO \\ \diagup \end{smallmatrix} Me$ (246°) Formed from acetoacetic ether and ammonia zinc chloride (Canzoneri & Spica, *G* 16, 449). Oil.

Oxy-di methyl-pyridine *Methyl derivative* $N \begin{smallmatrix} CMc CH \\ \diagup \end{smallmatrix} CO Me$. (208°) SG $\frac{2}{13}$

1011 Formed from chloro-di methyl pyridine, NaOMe, and MeOH at 160° (Conrad & Eckhardt, *B* 22, 81). Oil— $B'H_2PtCl_6$ — $B'MeI$ [204°] Prisms (from water).

Ethyl derivative C_8H_8EtNO (215°)— $B'MeI$ [196°] Crystals (from water).

Oxy di methyl-pyridine C_8H_8NO ϵ
 $NH \begin{smallmatrix} CMc CH \\ \diagup \end{smallmatrix} CO \begin{smallmatrix} CH \\ \diagdown \end{smallmatrix} CO Me$, ψ *Lutidostyryl* [176°] (304°) Formed by heating oxy tri methyl pyridine (methyl ψ lutidostyryl) in a current of HCl (Hantzsch, *B* 17, 2904). Formed also by distilling its carboxylic acids (Collie, *B* 20, 446, Nieme & Pechmann, *A* 261, 205), and by passing dry NH_3 over mesitene lactone at 160° (Anschutz, *A* 259, 169). Needles (from alcohol)— $B'HCl2aq$ prisms— $B'H_2PtCl_6$ — C_6H_5KNO silvery spangles, sl sol KOH aq.

Oxy di methyl-pyridine C_8H_8NO ϵ
 $CO \begin{smallmatrix} CH CMc \\ \diagup \end{smallmatrix} CH CMc \begin{smallmatrix} NH \\ \diagdown \end{smallmatrix}$ *Lutidone* [231°] (Collie, *C J* 59, 177) (350°) Formed by heating its carboxylic or dicarboxylic acid at 280°, and also by heating dehydracetic acid with NH_3 at 100° (Hantzing, *B* 18, 452, Conrad & Guthzeit, *B* 20, 156). Formed also from di acetyl acetone and NH_3 aq (Feist, *B* 22, 1571). Monoclinic pyramids (containing 8aq) PCl_5 gives chloro-di-methyl-pyridine (178°). $FeCl_3$ gives a brownish-red colour— $B'H_2PtCl_6$ [at 100°] [231° cor]— $B'H_2Cr_2O_7$, [125°]— $B'C_6H_5(OH)(NO_2)$, [220°]

Oxy tri methyl-pyridine C_8H_8NO ϵ
 $CO \begin{smallmatrix} CH CMc \\ \diagup \end{smallmatrix} CH CMc \begin{smallmatrix} NMe \\ \diagdown \end{smallmatrix}$ *Methyl-lutidone* [245°] Formed by heating its dicarboxylic acid (Conrad & Guthzeit, *B* 20, 159). Needles (containing 8aq) [111°], v sol water— $B'HI$ [242°] Formed by heating lutidone with MeI and MeOH at 140° (Conrad & Eckhardt, *B* 22, 80). Crystals.

Oxy-tri-methyl pyridine C_8H_8NO ϵ
 $NMe \begin{smallmatrix} CMc CH \\ \diagup \end{smallmatrix} CO \begin{smallmatrix} CH \\ \diagdown \end{smallmatrix} Me$ *Methyl ψ -lutidostyryl* [92°], (292°) Formed by heating 'dicarbo-collidylum dehydride' with H_2SO_4 or HCl at 150°–180° (Hantzsch, *B* 17, 1025, 2908). Formed also by methylation of ψ -lutidostyryl. Very hygroscopic crystals, not volatile with steam. V. sol. water, v. sl. sol. ether.— $B'HCl \frac{1}{2} aq$ —

$B'HI$ — $B'H_2PtCl_6 2aq$ — $B'H_2PtCl_6 2EtOH$ — $B'H_2PtCl_6$, long yellowish red needles.

Di-oxy-tri-methyl-pyridine *Di-ethyl derivative* $N \begin{smallmatrix} CMc C(OEt) \\ \diagup \end{smallmatrix} CMc C(OEt) \begin{smallmatrix} CO \\ \diagdown \end{smallmatrix} Me$ (218°) at 726 mm. Formed from di bromo-collidine and NaOEt (Pfeiffer, *B* 20, 1850). Oil— $B'H_2PtCl_6$.

OXY-DI-METHYL-PYRIDINE CARB-

OXYLIC ACID $CMc NH CMc$ *Lutidone carboxylic acid* $CH-CO C CO_2H$ [258° cor] Got by saponifying its ether. Crystals (containing aq)— BaA' — CuA' — AgA' amorphous. *Ethyl ether* [164° cor] (240°–250°) Formed in small quantity in distilling β amido crotonic ether under reduced pressure (Collie, *A* 226, 310, *C J* 59, 174). Needles, sol water, sl sol alcohol. Does not react with Ac_2O or phenyl hydrazine. Yields a bromo derivative C_8H_8BrNO , [250°] PCl_5 forms C_8H_8ClNO , (264° cor) which yields a chloro-di methyl-pyridine (177°–180°) on heating.

Oxy-di-methyl pyridine carboxylic acid

$CO \begin{smallmatrix} CH CMc \\ \diagup \end{smallmatrix} NH CMc \begin{smallmatrix} CO_2H \\ \diagdown \end{smallmatrix}$ [258°] Formed from isodehydracetic acid and ammonia (Nieme & Pechmann, *A* 261, 206).

Oxy methyl-pyridine dicarboxylic acid v METHYL CHELIDAMIC ACID, vol 1 p 729

Oxy-di-methyl pyridine dicarboxylic acid
 $CO \begin{smallmatrix} CH C(CH_2 CO_2H) \\ \diagup \end{smallmatrix} NH CMc C(CO_2H) \begin{smallmatrix} CO_2H \\ \diagdown \end{smallmatrix}$ ψ *Lutidostyryl di carboxylic acid* [201°] Formed from citra cumalic acid and NH_3 aq (N & P). Needles, v sl sol cold water and alcohol.

Oxy di-methyl pyridine carboxylic acid

$CO \begin{smallmatrix} CH CMc \\ \diagup \end{smallmatrix} NH CMc \begin{smallmatrix} CO_2H \\ \diagdown \end{smallmatrix}$ *Ethyl ether EtA'*. ψ *Lutidostyryl carboxylic ether* [187°] Formed by passing dry NH_3 through mesitene carboxylic acid lactone at 160° (Anschutz, *A* 259, 173). It is also a product of the condensation of amido acetoacetic ether and of β amido crotonic ether (Collie, *B* 20, 445). Needles (from hot water). The corresponding acid is an insoluble crystalline powder [300°].

Oxy-di methyl-pyridine dicarboxylic acid

$CO \begin{smallmatrix} C(CO_2H) CMc \\ \diagup \end{smallmatrix} C(CO_2H) CMc \begin{smallmatrix} NH \\ \diagdown \end{smallmatrix}$ [267°] Made by saponifying its ether. Prisms (from water)— KA' — CaA' 2aq— CuA' 1 $\frac{1}{2}$ aq (dried at 100°). *Ethyl ether EtA'* [221°] *S* (alcohol) 1 at 20° Formed by adding NH_3 aq to an alcoholic solution of $CO \begin{smallmatrix} C(CO_2Et) CMc \\ \diagup \end{smallmatrix} C(CO_2Et) CMc \begin{smallmatrix} O \\ \diagdown \end{smallmatrix}$ obtained from cupric acetoacetic ether and $COCl_2$ (Conrad & Guthzeit, *B* 19, 24, 20, 154). Forms an acetyl derivative C_8H_8NO , [65°]— $B'H PtCl_6$, [190°] Orange pp.

Oxy-tri methyl-pyridine dicarboxylic acid

$CO \begin{smallmatrix} C(CO_2H) CMc \\ \diagup \end{smallmatrix} C(CO_2H) CMc \begin{smallmatrix} NMe \\ \diagdown \end{smallmatrix}$ [245°] Formed by saponifying its ether (C & G)— NaA' crystalline, v sol water— EtA' [193°] Formed by the action of methylamine on dimethyl pyrone dicarboxylic ether, which is obtained from cupric acetoacetic ether and $COCl_2$. Needles (Gerichten, *B* 19, 25, Conrad & Eckhardt, *B* 22, 80).

OXY-METHYL-PYRIDYL-PROPIONIC ACID v. ECGONINE.

OXY-DI-METHYL PYRIMIDINE

$\text{CMe} \begin{smallmatrix} \text{NH CMe} \\ \text{N C(OH)} \end{smallmatrix} \text{CH}$ [192°] Formed from acetamide hydrochloride, acetoacetic ether, and dilute (10 p c) NaOHAq (Pinner, B 18, 2845, 2°, 1616). Needles, v e sol ordinary solvents

Ethyl derivative [55°] (259°) Prisms

Oxy-tri-methyl pyrimidine

$\text{CMe} \begin{smallmatrix} \text{NH CMe} \\ \text{N C(OH)} \end{smallmatrix} \text{CMe}$ Formed from acetamide hydrochloride, methylacetoacetic ether, and dilute NaOHAq (Pinner, B 22, 1617) Needles, v sol water

Di-oxy-methyl-pyrimidine

$\text{CO} \begin{smallmatrix} \text{NH CMe} \\ \text{NH CO} \end{smallmatrix} \text{CH}$ *Methyl uracil* Formed by boiling β uramido crotonic acid with acids (Behrend, A 229, 8, 231, 256) Needles (from alcohol) Decomposes at 270°-280° By heating with potash it is converted into $\text{C}_6\text{H}_2\text{I}_2\text{N}_2\text{O}_2$ Potash and Mel at 140° forms a dihydride $\text{C}_6\text{H}_4\text{N}_2\text{O}_2$ [219°] and di-oxy-tri-methyl pyrimidine A mixture of PCl_5 and POCl_3 at 125° forms oily $\text{C}_6\text{H}_4\text{Cl}_2\text{N}_2$ (246°), SG $\frac{218}{226}$ 1.6273 Yields deep violet coffin-like crystals of di iodide $\text{C}_6\text{H}_2\text{N}_2\text{O}_2\text{I}_2$ (Hoffmann, A 253, 74)

Di-oxy tri-methyl-pyrimidine

$\text{CO} \begin{smallmatrix} \text{NMe CMe} \\ \text{NMe CO} \end{smallmatrix} \text{CH}$ [109°] Formed by methylation of the preceding body (Behrend, Hagen, A 244, 2) Plates, v sol water and alcohol, sl sol ether Yields methylamine on heating with baryta water at 200° Dry Br forms $\text{C}_6\text{H}_4\text{NO}_2\text{Br}_2$, aqueous bromine yields $\text{CO} \begin{smallmatrix} \text{NMe CMe(OH)} \\ \text{NMe CO} \end{smallmatrix} \text{CBr}_2$ [163°], whence boil

ing alcohol gives bromo oxy tri-methyl pyrimidine [126°], converted by conc aqueous ammonia into oxy-amido tri-methyl pyrimidine $\text{CO} \begin{smallmatrix} \text{NMe CMe} \\ \text{NMe CO} \end{smallmatrix} \text{CNH}_2$ [167°], a body that is changed by potassium cyanate and HCl into crystalline $\text{CO} \begin{smallmatrix} \text{NMe CMe} \\ \text{NMe CO} \end{smallmatrix} \text{C NH CO NH}_2$

OXY-METHYL-PYRROLE Dihydride

$\text{CH}_2 \text{CO} \begin{smallmatrix} \text{CH}_2 \text{CHMe} \end{smallmatrix} \text{NH}$ γ Amido valeric lactam [37°] Formed by heating γ amido valeric acid and by the action of sodium amalgam and HOAc at 280° on the phenyl-hydrazide of levulic acid (Tafel, B 20, 250, 22, 1862) Yields an oily nitrosamine, which produces valerolactone on distillation -B'HCl [110°] Needles, v sol water -B', H_2PtCl_6

ν Oxy-di-methyl pyrrole $\text{C}_6\text{H}_4\text{NO}$ 4.6

$\text{CH CMe} \begin{smallmatrix} \text{C(OH) N} \\ \text{C(OH) N} \end{smallmatrix} \text{NOH}$ Made by heating its carb-oxylic acid (Knorr, A 236, 302) Reduces silver solution in the cold and Fehling's solution on boiling Gives a red substance with acids, and exhibits the pine-wood reaction

Oxy-tri-methyl-pyrrole ? $\text{C}_6\text{H}_4\text{NO}$ *Tri-methyl-pyrrolone* (175°) SG $\frac{2}{2}$ 945 Formed from sodio ethyl cyanide and Mel (Hanriot A. Bouveault, Bl. [8] 1, 175)

Di-oxy-tri-methyl pyrrole v. ACETONAMINES, vol. i. p. 27.

OXY-METHYL-PYRROLE CARBOXYLIC

ACID $\text{CO} \begin{smallmatrix} \text{NH CMe} \\ \text{CH}_2 \text{C CO}_2\text{H} \end{smallmatrix}$ *Ethyl ether EtA'*.

[134°] Formed by heating α -amido-ethylidene-succinic ether at 150° (Emery, A 260, 144) Needles, v sol alcohol Yields an acetyl derivative [142°]

Oxy-di-methyl-pyrrole carboxylic acid

Ethyl ether $\text{CO} \begin{smallmatrix} \text{NH CMe} \\ \text{CHMe C CO}_2\text{Et} \end{smallmatrix}$ [127°]

Formed from methyl acetosuccinic ether and alcoholic NH_3 at 0° (Emery, A 260, 151) Small white prisms (from HOAc)

Oxy-di-methyl-pyrrole carboxylic acid

Ethyl ether $\text{CO} \begin{smallmatrix} \text{NMe CMe} \\ \text{CH}_2 \text{C CO}_2\text{Et} \end{smallmatrix}$ [42°]

(160° at 11 mm) Formed from acetosuccinic ether and alcoholic methylamine (E) Bunches of needles, v sol ether

Oxy-di-methyl-pyrrole carboxylic acid

$\text{C}_6\text{H}_4\text{NO}_2$ 1.6 $\begin{smallmatrix} \text{N(OH) CMe} \\ \text{CMe CH} \end{smallmatrix} \text{C CO}_2\text{H}$ Formed by boiling oxy di-methyl pyrrole dicarboxylic ether with NaOHAq (Knorr, A. 236, 301) Slender needles, v sol alcohol Gives off CO_2 at 138°

Oxy-di-methyl-pyrrole dicarboxylic acid

$\text{N(OH)} \begin{smallmatrix} \text{CMe C CO}_2\text{H} \\ \text{CMe C CO}_2\text{H} \end{smallmatrix}$

Mono-ethyl ether EtHA' Formed by boiling the di-ethyl ether with alcoholic potash (Knorr, A 236, 299) Crystals (from alcohol) Decomposes at 185° into CO_2 and oxy-dimethyl pyrrole carboxylic ether

Di-ethyl ether EtA'' [99°] Formed by heating di-acetyl succinic ether with hydroxyl amine hydrochloride, NaOAc, and HOAc (Knorr) Yields the salt $\text{C}_{12}\text{H}_{16}\text{KNO}_3$

OXY METHYL-PYRROLE DIHYDRIDE CARBOXYLIC ACID Nitrile

$\text{CH}_2 \text{CH}_2 \begin{smallmatrix} \text{CH}_2 \text{CH}_2 \end{smallmatrix} \text{CMe CN}$ [141°] Formed by heating $\text{CH}_2 \text{CO CH}_2 \text{CH}_2 \text{CO}_2\text{Et}$ with HCy and ammonia in a closed tube (Kuhling, B 22, 2369, 23, 708) Octahedra (from hot alcohol) Successive treatment with cold H_2SO_4 and water converts it into the corresponding amide [161°], Hydroxylamine yields the amidoxim [156°]

OXY-METHYL-PYRROLE DIHYDRIDE THIOCARBOXYLIC AMIDE

$\text{CH}_2 \text{CH}_2 \begin{smallmatrix} \text{CH}_2 \text{CH}_2 \end{smallmatrix} \text{CMe CS NH}_2$ [220°] Formed by passing H_2S through an ammoniacal solution of the nitrile of oxy-methyl pyrrole dihydride carb-oxylic acid (Kuhling, B 22, 2370) Prisms (from hot water), almost insol alcohol

OXY-METHYL-QUINAZOLINE

$\text{C}_6\text{H}_4 \begin{smallmatrix} \text{C(OH) N} \\ \text{N=CMe} \end{smallmatrix}$ *Anhydro-acetyl o-amido benzamide* [238°] Formed from acetyl o-amido benzamide by the action of heat, alkalis, or boiling water (Weddige, J pr [2] 36, 143)

Silky needles (containing aq), v sol hot alcohol Mel and KOH yield the methyl derivative $\text{C}_6\text{H}_4\text{N}_2(\text{OMe})$ [109°] -B'HCl -B', H_2PtCl_6

Oxy-methyl-quinazoline

$\text{CH CH C(OH) N} \begin{smallmatrix} \text{C} \\ \text{C} \end{smallmatrix} \text{CH}$ [238°] Formed from o-amido-p-toluic amide and formic acid (Niemen towski, J pr [2] 40, 12). Crystals, insol water.

Oxy-di-methyl-quinazoline $C_{10}H_{10}N_2O$ *see* $C_6H_4Me < \begin{smallmatrix} CO NH \\ N CMe \end{smallmatrix}$ [255°] Formed by boiling *o* amido-*p* toluic amide with Ac_2O (Niementowski, *B* 21, 1534, *J pr* [2] 40, 13) Needles (from boiling water), sol acids and alkalis

Oxy-di-methyl-quinazoline $C_6H_4 < \begin{smallmatrix} CO-N \\ NMe CMe \end{smallmatrix}$ [199°] Formed by heating acetyl *o* amido benzamide above its melting point (Weddige, *J pr* [2] 36, 154) Crystals (containing 3aq), v e sol alcohol— $B'HCl$

Di-oxy-methyl quinazoline $\begin{smallmatrix} CH CH C CO NH \\ CH CMe C NH CO \end{smallmatrix}$ (?) Formed by heating amido *m*-toluic acid with urea at 180° (Niementowski, *J pr* [2] 40, 21) White needles (from amyl alcohol), insol benzene Not melted at 300° HI and P yield *o* toluidine

Di oxy methyl-quinazoline $C_6H_4 < \begin{smallmatrix} CO NMe \\ NH CO \end{smallmatrix}$ [284°] Formed by heating $C_6H_4(NH_2)CO NHMe$ with urea at 200° (Abt, *J pr* [2] 39, 147) Long white needles, sl sol hot water

Di-oxy-methyl quinazoline $C_6H_4 < \begin{smallmatrix} CO NH \\ NMe CO \end{smallmatrix}$ [148°] Formed by heating $C_6H_4(NHMe)CO NHCO_2H$ with urea (Abt)

Di-oxy-di-methyl quinazoline $C_6H_4 < \begin{smallmatrix} CO NMe \\ NMe CO \end{smallmatrix}$ [151°] Formed by the action of MeI and NaOH on either of the two preceding bodies or on di oxy quinazoline (Abt, *J pr* [2] 39, 145) White needles (from water), v sol alcohol

Oxy methyl-quinazoline dihydride $C_6H_4 < \begin{smallmatrix} CH_2 NMe \\ NH CO \end{smallmatrix}$ [120°] Formed by boiling *o*-oxy tolyl methyl thio urea with yellow HgO Soderbaum a Widman, *B* 22, 2936) Needles, v e sol methyl alcohol— $B'_2H_2PtCl_6$ [203°] $B'HAuCl_4$ [185°] Yellow prisms

OXY-METHYL QUINIZINE v OXY PHENYL-METHYL-PYRAZOLE

(*B* 1, 4)-**OXY METHYL-QUINOLINE** $\begin{smallmatrix} CH C(OH) \\ CH CMe \end{smallmatrix} > C_5H_4N$ [263°] Formed from the amido compound and HNO_2 (Noelting a Trautmann, *Bl* [3] 4, 244), and by fusing (*B* 4) methyl quinoline (*B* 1)-sulphonic acid with NaOH (Herzfeld, *B* 17, 905, 1551) Needles (from dilute alcohol) Yields a nitroso derivative $C_9H_8Me(NOH)(OH)N$ [200°] By KOH, MeI, and MeOH it is converted into the methyl derivative $C_{10}H_9(OMe)N$ (*c* 230°), whence $B'_2H_2PtCl_6$

(*B* 4, 1) **Oxy-methyl-quinoline** $\begin{smallmatrix} CH CMe \\ CH C(OH) \end{smallmatrix} > C_5H_4N$ [124°] Formed by heating amido *p*-cresol (10 g) with glycerin (24 g), H_2SO_4 (20 g), and picric acid (2 g) (*N* a T) Needles, v sl sol cold water Dyes fabrics mordanted with alumina, yellow

(*B* 4, 2)-**Oxy methyl-quinoline** $\begin{smallmatrix} CMe CH \\ CH C(OH) \end{smallmatrix} > C_5H_4N$ [96°] Formed from the amido- compound by the diazo- reaction (*N* a T) Formed also by fusing the sulphonic acid with NaOH (O Fischer a Willmack, *B* 17, 441, Herzfeld, *B*, 17, 1552) Needles (from chloro-

form) Smells like vanilla Its alcoholic solution is coloured green by $FeCl_3$ Yields a nitroso-derivative [200°]— $B'_2H_2PtCl_6$ 2aq orange needles

Methyl derivative $C_{11}H_{11}NO$ Oil— $B'_2H_2PtCl_6$ 2aq brown crystalline pp

Tetrahydride $C_9H_8Me(OH) < \begin{smallmatrix} CH_2 CH_2 \\ NH CH_2 \end{smallmatrix}$ Formed by reduction with tin and HCl Needles or plates, sl sol water Yields a nitrosamine $C_{10}H_9(NO)NO$ crystallising in small yellow needles

(*B* 4, 3) **Oxy-methyl-quinoline** $\begin{smallmatrix} CH CH \\ CMe C(OH) \end{smallmatrix} > C_5H_4N$ [74°] Formed from amido *o* cresol hydrochloride (10 g), glycerin (24 g), H_2SO_4 (20 g), and picric acid (2 g) (Noelting a Trautmann, *B* 23, 3663) Needles (from dilute alcohol), volatile with steam $FeCl_3$ gives a dark green colour Mixed with CuO it colours a flame green Yields a crystalline *p* nitroso derivative decomposing at 200° without melting

(*B* 2, 4)-**Oxy-methyl quinoline** $\begin{smallmatrix} C(OH) CH \\ CH CMe \end{smallmatrix} > C_5H_4N$ [200°] Formed by soda fusion from *o* toluquinoline sulphonic acid (Herzfeld, *B* 17, 903) Needles Not volatile with steam $FeCl_3$ colours its alcoholic solution brownish red

(*B* 1, 2) **Oxy-methyl quinoline** $\begin{smallmatrix} CMe C(OH) \\ CH CH \end{smallmatrix} > C_5H_4N$ *Ana oxy paratoluquinoline* [230°] Formed from the amido compound, and also from *p*-toluquinoline by sulphonation (with 25 p c SO_3 extra) at 90° and potash fusion (Noelting a Trautmann, *B* 23, 3658) Needles, v sl sol hot water Not volatile with steam

(*Py* 3, 1) **Oxy methyl quinoline** $C_6H_4 < \begin{smallmatrix} CMe CH \\ N = C(OH) \end{smallmatrix}$ [224° cor] (above 360°) Formed by heating the amide of acetoacetic acid with H_2SO_4 (Knorr, *A* 236, 83, *C J* 46, 334, Roos, *B* 21, 624, Reissert, *B* 24, 855) Small needles (from water) Reduced by sodium amalgam to $C_{10}H_9N_2O_2$ [280°] Sodium added to its alcoholic solution reduces it to a dihydride $C_{10}H_{11}NO$ [101°] and methyl quinoline tetrahydride $C_{11}H_{13}N$ (253°) $NaOEt$ and MeI form oxy-di-methyl quinoline [132°] and the methyl derivative $C_{10}H_9(OMe)N$ [276° cor], whence $B'_2H_2PtCl_6$ The ethyl derivative $C_{11}H_{11}(OEt)N$ [51°] (250°) is formed from chloro lepidine and KOEt

Salts— $B'HCl$ [187°]— $B'_2H_2PtCl_6$ 2aq— $B'_2H_2SO_4$ needles— $B'HNO_3$ prisms— $B'HI$ —Picrate [166°] Needles— $Ba(C_{10}H_9NO)_2$ 2aq

(*Py* 3, 4) **Oxy methyl quinoline** $C_6H_4 < \begin{smallmatrix} CH CH \\ NMe CO \end{smallmatrix}$ *Methyl- ψ carbostyryl* [72°]

Formed by digesting carbostyryl with MeOH and MeI, adding NaOH as required to neutralise the HI formed (Friedlander a Muller, *B* 20, 2010) Slender needles Weak base Sodium amalgam forms $C_{10}H_9NO$ [276°]— $B'_2H_2PtCl_6$ 2aq— $B'HgCl_2$ [189°] Small pyramids

Methyl- α -iodide $B'MeI$ Bronzed needles

(*Py* 1, 3) **Oxy methyl-quinoline** $C_6H_4 < \begin{smallmatrix} C(OH) CH \\ N = CMe \end{smallmatrix}$ [231°] (above 360°) S 1 in the cold, 10 at 100° Formed by heating

phenylamido crotonic ether rapidly to 240° (Conrad a Limpach, *B* 20, 947) Prisms (containing 2aq), v sol alcohol Tastes bitter FeCl₃ colours its solution yellowish-red KMnO₄ oxidises it to acetyl nitrilic acid Yields quinaldine on distillation with zinc dust —B'HCl —B'₂H₂PtCl₄ [215°] —B'₂H₂CrO₄ [108°] —B'₂O₂H₂(NO₃)₂OH [200°] Yellow needles

Methyl derivative C₁₀H₈(OMe)NO [82°] (296°) Formed from chloro methyl quinoline, MeOH, and NaOMe at 135° Needles, sl sol water

Methylo chloride C₁₀H₇NOMeCl aq [217°] Formed from the methylo iodide and AgCl (Conrad a Eckhardt, *B* 22, 74) —B'₂H₂PtCl₄ [240°] Yellow crystalline pp

Methylo iodide C₁₀H₇NOMeI aq [201°] Formed from the base and MeI at 100° Satiny needles (from hot water)

Sulphonic acid C₁₀H₈NO(SO₃H) [283°]. Long prisms (containing 2aq) —BaA₂ 4aq

(*B* 2)-Oxy-(*Py* 1) methyl quinoline
C(OH) CH C CMe CH [218°] Formed by fusing lepidine sulphonic acid with NaOH (Busch a Koenigs, *B* 23, 2684) Obtained also by boiling with HBrAq the methyl derivative which is produced by heating quinine sulphate (40 g) with KOH (95 g) and water (45 cc) at 220° in a current of superheated steam (Koenigs, *B* 23, 2674) Groups of slender needles, v sol warm alcohol and acetone Gives no colour with FeCl₃

Methyl derivative C₁₁H₁₁NO [52°] Formed as above, and also by heating quinine zinc chloride with water at 200° Slender needles (containing aq) Its solution fluoresces like quinine Gives a bluish green colour with ammonia and chlorine water —B'₂H₂PtCl₄ [237°] Orange powder

Oxy (Py 1) methyl-quinoline [141°] Formed from a sulphonic acid of lepidine prepared by mixing lepidine with H₂SO₄ and heating to 300° (Busch a Koenigs, *B* 23, 2686) Greenish needles, v sol benzene —B'₂H₂PtCl₄ 2aq orange-yellow needles

(*B* 2) Oxy (*Py* 3) methyl quinoline
C(OH) CH C CH CH *p* Oxy quinaldine
CH —CH C N CMe [213°] Formed by the action of paraldehyde and HCl upon *p* amido-phenol, and also by fusing (*Py* 3) methyl quinoline sulphonic acid with potash (Doebner a Miller, *B* 17, 1708) Crystals, not volatile with steam, v sl sol cold water —B'₂H₂PtCl₄ 2aq yellow needles

(*B* 2') Oxy (*Py* 3) methyl-quinoline
C₁₀H₇Me(OH)N [234°] Formed by fusing quinaldine (*B*) sulphonic acid with potash (*D* a *M*) Silvery plates, sol ether and hot alcohol, nearly insol hot water —B'HCl 2aq —B'₂H₂PtCl₄ 2aq small yellow needles

(*B* 4)-Oxy (*Py* 3) methyl-quinoline
CH CH — C CH CH
CH C(OH) C N CMe *o* Oxy quinaldine [74°] (266°) Formed by fusing (*Py* 3) methyl-quinoline (*B* 4)-sulphonic acid with potash, and also by the action of paraldehyde and HCl on *o* amido phenol (Doebner a Miller, *B* 17, 1705). Trimetric prisms, easily volatile with steam. —B'₂H₂PtCl₄ 2aq yellow needles

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Methyl derivative C₁₀H₇Me(OMe)N [125°] (282°) Formed from *o* anisidine, paraldehyde, and HCl —B'₂H₂PtCl₄ yellow needles

Tetrahydride C₁₀H₇Me(OH)N (280°) Formed by reducing *o* oxy quinaldine with tin and HCl (Doebner a Miller, *B* 17, 1706) Yields C₁₀H₇Me(OMe)N (270°) whence B'HCl and C₁₀H₇Me(OMe)NMe (261°), whence B'₂H₂PtCl₄ yellow needles

(*B* 2) Oxy (*Py* 4)-methyl quinoline tetrahydride **Methyl derivative** C₁₀H₇NMe(OMe) **Methyl thallium** (278°) Formed by methylation of thallium (Skraup, *M* 6, 776) Oil —B'₂H₂SO₄ —B'MeI aq [224°] —B'₂Me₂PtCl₄ Orange plates

(*B* 4) Oxy (*Py* 4) methyl quinoline
Tetrahydride C₁₀H₇NO 2e

CH CH — C CH₂ CH₂ [114°] Formed from CH C(OH) C NMe CH₂ (Fischer, *B* 16, 714) Trimetric tables (from ether) *abc* = 631.1.538 Its alcoholic solution is coloured brown by FeCl₃ —B'HCl aq *Kaurne*, a febrifuge —B'MeI [216°] Prisms (from MeOH) —B'C₂H₂Cl₄ (312°) Oil (Fischer a Kohn, *B* 19, 1040, *C* 7, 49, 503)

Methyl derivative C₁₀H₇(OMe)NMe (257°) Formed by means of MeI and MeOH Pale yellow oil Gives a crimson colour with NaNO₂ and a little acid —B'₂H₂PtCl₄ [199°] —B'₂H₂SO₄ prisms, v sol water

Methyl derivative of the methyloiodide C₁₀H₇(OMe)NMe I [175°] Prisms (from MeOH) Moist Ag₂O converts it into strongly alkaline crystalline C₁₀H₇(OMe)NMe₂OH, whence [C₁₀H₇(OMe)NMe Cl]₂ PtCl₄ [200°]

Ethyl derivative C₁₂H₁₃(OEt)NMe (270°) at 716 mm Oil

(*Py* 3) Oxy (*B* 2) methyl quinoline
CMe CH C CH CH
CH CH C N H CO **Methyl carbostyryl** [228°] Formed by boiling chloro methyl carbostyryl with NaOHAq (Einhorn a Lauch, *A* 243, 359) Crystalline, v sol alcohol

(*B* 1,2,4) Oxy di methyl quinoline
CMe C(OH) C CH CH [198°] Formed by the action of nitrous acid on (*B* 1,2,4) amido di methyl quinoline (Noeltling a Trautmann, *B* 23, 3683) Plates (from chloroform), v sol alcohol

(*Py* 3,1,4) Oxy-di methyl quinoline C₁₁H₁₁NO
CMe CH
C₁₀H₇ NMe CO **Methyl lepidone Di-**

methyl-*q* carbostyryl [132°] (290° at 250 mm) Formed by heating (*Py* 3,1) oxy methyl quinoline with KOH and MeI, or by heating the isomeric methoxy lepidine above its boiling point (290°) Formed also by heating methylaniline with acetoacetic ether, and treating the product with H₂SO₄ (Knorr, *B* 17, 2876, 19, 3301, *A* 236, 104) Needles, sl sol water and ether, v sol alcohol, insol alkalis Sodium amalgam reduces it to (C₁₁H₁₁NO)₂ [268°] —B'₂H₂PtCl₄ 3aq [214°] Slender needles

(*Py* 1,3,4) Oxy di-methyl-quinoline
C₁₀H₇ < CO CH
NMe CMe [176°] Formed from methoxy methyl quinoline by heating at 315° in a sealed tube, or by heating oxy quinaldine methylo iodide with NaHCO₃ (Conrad, *B* 20, 956, 22, 75) Needles, v sol water —B'HHgCl₄

3 A

[187°] Needles — $B'_2H_2PtCl_4$ [240°]. — $B'MeI$ [210°] Sl sol cold water

(*Py* 3, 2, 1) Oxy di methyl quinoline

$C_8H_7 \begin{smallmatrix} \text{CMe CMe} \\ \text{N}=\text{COH} \end{smallmatrix}$ [262°] Formed from methyl-acetoacetic anilide and H_2SO_4 in the cold (Knorr, *A* 245, 357) Yields a sulphonic acid, which forms a crystalline Ba salt $Ba(C_{11}H_{10}NSO_3)_2$. — $B'HCl$ Long silky needles

(*B* 2)-Oxy (*Py* 1, 3) di methyl quinoline

$C(OH)CH \begin{smallmatrix} CMe CH \\ CN=O \end{smallmatrix}$ [214°] (above 360°)

Formed by heating *p* amido phenol hydrochloride (1 vol) with acetone (3 vols) for some days at 175° (Engler a Bauer, *B* 22, 213) Prisms or tables, v sl sol water $FeCl_3$ colours its alcoholic solution brown — $B'HCl$. — $B'_2H_2PtCl_4$ 2aq — $B'_2H_2SO_4$ — $B'_2H_2Cr_2O_7$. — $B'_2C_2H_2(NO_2)_2OH$ [225°] Yellow plates or prisms

(*B* 4)-Oxy (*Py* 1, 3) di methyl-quinoline

$CH \begin{smallmatrix} CH \\ CN=O \end{smallmatrix}$ [65°] (281° uncor)

Formed in like manner from *o* amido phenol Prepared by saturating a mixture of acetone (3 mols) and paraldehyde (8 mols) with dry HCl , adding, after 3 days, *o* amido phenol (2 mols) dissolved in conc HCl aq, and heating on the water bath Crystals, v sol alcohol, ether, and benzene $FeCl_3$ colours its alcoholic solution green — $B'_2H_2SO_4$. — $B'HCl$. — $B'_2H_2PtCl_4$ 2aq — $B'_2H_2Cr_2O_7$. — $B'_2C_2H_2N_2O_3$ [207°] Plates or prisms, sl sol hot alcohol

(*Py* 3)-Oxy-(*B* 2 *Py* 1)-di-methyl-quinoline

$CMe \begin{smallmatrix} CH \\ CN=O \end{smallmatrix}$ [250°] Formed from $CMe \begin{smallmatrix} CH \\ CN=O \end{smallmatrix}$ by successive treatment with *p* toluidine and H_2SO_4 (Knorr, *A* 245, 365) Flat prisms, sl sol hot water, dilute acids, and alkalis Yields (*B* 2, *Py* 1)-di methyl-quinoline on distillation with zinc dust

(*Py* 3) Oxy (*B* 3, *Py* 1) di-methyl-quinoline

$CH \begin{smallmatrix} CH \\ CN=O \end{smallmatrix}$ [220°] Formed in like manner from *m* toluidine (K) V sl sol hot water Yields (*B* 3, *Py* 1)-di methyl quinoline on distillation with zinc dust — Platinochloride [234°] The hydrochloride is crystalline, and is decomposed by water

(*Py* 3)-Oxy-(*B* 4; *Py* 1) dimethyl quinoline

$CH \begin{smallmatrix} CH \\ CN=O \end{smallmatrix}$ Formed in like manner from *o* toluidine (Knorr, *A* 245, 368) Slender needles (from water) Yields (*B* 4, *Py* 1) di-methyl-quinoline on distillation — $B'_2H_2PtCl_4$ 2aq [220°] — $NaC_{11}H_{10}NO$ Plates

(*Py* 1) Oxy (*B* 2, *Py* 3) di-methyl quinoline

$CMe \begin{smallmatrix} CH \\ CN=O \end{smallmatrix}$ [275°] Formed by heating *p* tolyl amido crotonic ether $C_8H_7NH \begin{smallmatrix} CMe CH \\ CO_2Et \end{smallmatrix}$ at 250° (Conrad a Limpach, *B* 21, 525, cf Knorr, *B* 17, 542) Needles (containing aq) — $B'HCl$ Needles (from hot water) — $B'_2H_2PtCl_4$ [228°] Prisms (from hot water)

(*Py* 1)-Oxy-(*B* 4, *Py* 3)-di methyl quinoline

$CH \begin{smallmatrix} CH \\ CN=O \end{smallmatrix}$ [261°] Formed by distilling *o*-tolyl amido-crotonic ether (C a L) Plates (containing aq) — $B'_2H_2PtCl_4$ Needles

Oxy-(*Py* 1, 3)-di-methyl-quinoline

$C_{11}H_{11}N(OH)$ [44°]. Made from (*Py* 1, 3) di-

methyl-quinoline by conversion into the sulphonic acid and fusing this with $NaOH$ (Beyer, *J pr* [2] 33, 409) — $(B'HCl)_2PtCl_4$ 2aq

Oxy-tri-methyl quinoline

$CMe \begin{smallmatrix} CH \\ CN=O \end{smallmatrix}$ [264°] Formed by heating *m* xylyl β amido crotonic ether (Conrad a Limpach, *B* 21, 526) Needles (containing aq) — Platinochloride [282°] Needles

Oxy tetra methyl-quinoline $C_{11}H_{11}NO$ etc

$C_8H_7 \begin{smallmatrix} C(OH)CH \\ CN=O \end{smallmatrix}$ Formed by quickly heating ψ -cumyl amido crotonic ether at 250° (Conrad a Limpach, *B* 21, 529) Prisms (from alcohol) Sublimes at 285°, without previous fusion — $B'_2H_2PtCl_4$ Prisms (from alcohol)

(*Py* 1, 3, 4) Di oxy-methyl-quinoline

$C_8H_7 \begin{smallmatrix} C(OH)CH \\ NMe CO \end{smallmatrix}$ [260°] Formed by heating its methyl derivative with HCl aq at 120° Small needles, sol alkalis Yields a nitrosamine $C_{10}H_9NO_2(NO)$ crystallising in red needles [c 188°]

Methyl derivative $C_8H_7 \begin{smallmatrix} C(O)CH \\ NMe CO \end{smallmatrix}$

[68°] Formed by heating (*Py* 1, 3, 4) chloro oxy methyl quinoline with $NaOMe$ (Friedlander a Muller, *B* 20, 2014) Slender white needles, v sol alcohol — $B'_2H_2PtCl_4$ pyramids

Ethyl derivative $C_{10}H_9(OEt)NO$ [87.5°] Slender white needles

(*B* 4, *Py* 1) Di oxy-(*Py* 3) methyl quinoline

Methyl derivative $CH \begin{smallmatrix} CH \\ CN=O \end{smallmatrix}$ [229°] Formed by heating *o* methoxy phenyl amido crotonic ether at 260° (Conrad a Limpach, *B* 21, 1654) Long silky needles (containing aq) — $B'_2H_2PtCl_4$ [239°] Tables

(*B* 2, *Py* 1)-Di oxy (*Py* 3) methyl-quinoline

Methyl derivative $C(O)CH \begin{smallmatrix} CH \\ CN=O \end{smallmatrix}$ [290°] Formed by the action of heat on *p* methoxy phenyl amido crotonic ether which is made from *p* anisidine and acetoacetic ether (C a L) — $B'HCl$ — $B'_2H_2PtCl_4$ — $B'_2H_2SO_4$ S 6 in the cold — $B'MeCl$ [251°] Needles — $B'_2Me.PtCl_4$ 4aq — $B'MeI$ Converted by moist Ag_2O into $C_{11}H_{10}NO$ [149°]

Di-methyl derivative $C_8H_7Me(O)Me_2N$ [94°] Formed from the methyl derivative, $NaOMe$, and MeI Needles or prisms

(*B* 2, 3)-Di oxy-(*Py* 3) methyl quinoline

Methylene derivative $CH_2 \begin{smallmatrix} CCH \\ CCH \end{smallmatrix}$ [152°] Formed on reduction of

CH_2O_2 , $C_8H_7(NO_2)CH \begin{smallmatrix} CH \\ CN=O \end{smallmatrix}$ (Haber, *B* 24, 623) Silvery needles, m sol ether — $B'_2H_2PtCl_4$ — $B'_2H_2CrO_3$ Detonates at 210° — Picrate [175°] Yellow crystals

(*Py* 1, 2, 3, 4) - Tri - oxy - methyl - quinoline

$C_8H_7 \begin{smallmatrix} C(OH)CH \\ NMe CO \end{smallmatrix}$ Di-oxy-methyl-pseudo carbostyryl Formed by warming the nitroso-derivative of (*Py* 1, 3, 4)-di-oxy methyl-quinoline with $SnCl_4$ White needles, sl sol water, benzene, and $CHCl_3$ By $FeCl_3$ it is oxidised to methyl-pseudo-quinisatin

$C_6H_4 \begin{matrix} \diagup CO \\ \diagdown CO \\ | \\ NMe CO \end{matrix}$ [122°] (Friedlander & Muller, *B* 20, 2015)

References — Bromo, Chloro, and Di-chloro di nitro- oxy methyl quinolines

(*B* 4) OXY (*Py* 4) METHYL QUINOLINE CARBOXYLIC ACID *Tetrahydride*

$C_6H_4Me(OH)(CO_2H)N$ [211°] (S & E), [216°] (K & N) Formed by heating oxy quinoline carboxylic acid tetrahydride with MeI and MeOH at 120° (Schmitt & Engelmann, *B* 20, 1219, Krokowsky & Nencki, *M* 9, 208) Prisms (containing 2aq), v sol hot water and alcohol Has no toxic action After administration to dogs the urine contains the acid $C_6H_4Me(OH)_2(CO_2H)N$ [255], insol water

(*B* 4) Oxy- (*Py* 3) methyl quinoline carboxylic acid $NO_2C_6H_4Me(OH)(CO_2H)$ [207°] Formed by heating potassium oxy quinaldine with liquid CO_2 at 190° (König, *B* 21, 883) Yellow needles (containing aq), sl sol cold water Gives a cherry red colour with $FeCl_3$

(*Py* 1) Oxy- (*Py* 3) methyl quinoline (*Py* 2)-carboxylic acid $C_6H_4 \begin{matrix} \diagup C(OH) \\ \diagdown C(CO_2H) \\ | \\ N=CMe \end{matrix}$ [245°]

Formed by oxidising the corresponding aldehyde with alkaline $KMnO_4$ (Conrad & Lumpach, *B* 21, 1975) Crystals (from alcohol) — MgA_2 (dried at 100°) Crystalline pp

(*Py* 3, 1)-Oxy-methyl quinoline (*B* 4) carboxylic acid [312°] Formed by oxidising (a) oxy (γ) methyl julole dihydride with $KMnO_4$ in acid solution (Reissert, *B* 24, 853) Needles, m sol alcohol, sl sol hot water

(*Py* 1) OXY (*Py* 3) METHYL QUINOLINE (*Py* 2) CARBOXYLIC ALDEHYDE

$C_6H_4 \begin{matrix} \diagup C(OH) \\ \diagdown CCHO \\ | \\ N=CMe \end{matrix}$ [273°] Formed, together with $C_6H_4N_2O_3$ [192°], by boiling (*Py* 1, 3) oxy methyl quinoline with chloroform and KOHAq (Conrad & Lumpach, *B* 21, 1972) Yellow plates (from alcohol), sl acids and alkalis — $B'HCl$ — $B'H \cdot H \cdot PtCl$, [215°-220°]

Phenyl hydrazide $C_6H_4H_2N_2O$ — $B'HCl$ yellow needles, sl sol hot water

(*Py* 1) Oxy (*B* 1, 2, 4, *Py* 3) tetra methyl quinoline (*Py* 2) carboxylic aldehyde $CMe CMe C(OH) CCHO$ Formed by the action of caustic soda and chloroform on the corresponding oxy tri methyl quinoline (C & L) Yields, with phenyl hydrazine hydrochloride, crystalline $C_6H_4H_2N_2O \cdot OHCl$

OXY METHYL QUINOXALINE

$CH CH CN CH$ *Oxytoluquinoxaline*
 $CMc CH ON COH$

Formed by oxidation of its dihydride by the air or by ammoniacal $AgNO_3$ (Hinsberg, *B* 19, 483, *A* 248, 75) Feebly basic plates PCl , yields chloro methyl quinoxaline [77°]. — NaA' aq plates, sol water.

Methyl derivatives [71°].

Ethyl derivatives [67°] From the chloro methyl quinoxaline and $NaOEt$ Satiny needles *Dihydride* $C_6H_4N_2O$ [95°-124°] Formed by reducing o nitro tolyl amido acetic acid with tin and HCl (Plöchl, *B* 19, 10, Leuckart & Hermann, *B* 20, 27) Formed also from chloro acetic ether and tolylene o-diamine (Hinsberg, *B* 18, 2870) Very oxidisable.

Oxy di methyl quinoxaline

$C_6H_4Me \begin{matrix} \diagup N CMe \\ \diagdown N COH \end{matrix}$ [238°] Formed by passing air through an alkaline solution of the dihydride, which is obtained by condensing tolylene o diamine with a bromo propionic or pyruvic acid (Hinsberg, *A* 237, 351, 248, 78) Thin crystals (from alcohol), v sl sol water Is perhaps a mixture of isomerides

Dihydride $C_6H_4Me \begin{matrix} \diagup NH CMe \\ \diagdown NH CO \end{matrix}$ [o 135°]

Crystallises from alcohol.

Oxy-tri methyl quinoxaline *Dihydride*

$C_6H_4Me \begin{matrix} \diagup NH CMe_2 \\ \diagdown NH CO \end{matrix}$ [227°] Formed by heating tolylene o diamine with bromo isobutyric ether (Hinsberg, *A* 248, 79) Plates, v sl sol water Yields an acetyl derivative [206°], a nitrosamine [154°], and a di-nitro compound [above 280°]

Di oxy-di methyl-quinoxaline *Di-ethyl derivative* $CH C(OEt) C N CMe$ [127°]
 $CH C(OEt) C N CMe$

Formed by the action of di methyl-diketone (diacetyl) on the hydrochloride of the di ethyl derivative of (a) di amido-hydroquinone (Nietzki & Reehberg, *B* 23, 1212) Long yellow needles

Di-oxy-di-methyl-di-quinoxaline, so called, $CMc N C CH C N CMe$ Formed by the action of $NaOAc$ and pyruvic acid in excess on a solution of the hydrochloride of tetra amido benzene (Nietzki & Muller, *B* 22, 445) Dissolves in alkalis with greenish yellow fluorescence

OXY METHYL QUINOXALINE CARBOXYLIC

LIC ACID $C_6H_4H_2N_2O_2$ *te* $C_6H_4(CO_2H) \begin{matrix} \diagup N CMe \\ \diagdown N COH \end{matrix}$ Formed by adding pyruvic acid in slight excess to an aqueous solution of di amido benzoic acid (Zehra, *B* 23, 3629) Pale yellow needles, m sol hot alcohol Blackens above 300° without melting — BaA' 3aq yellowish white needles

Oxy-methyl-quinoxaline carboxylic acid $C_6H_4Me \begin{matrix} \diagup N C CO_2H \\ \diagdown N COH \end{matrix}$ Formed by the action of boiling potash on $C_6H_4H_2N_2O$, [258°], which is made by mixing aqueous solutions of alloxan and tolylene o diamine (Hinsberg, *A* 237, 356) Yellow needles (from dilute alcohol) Gives off CO_2 at 214°

OXY-METHYL SUCCINIC ACID v OXYTARTARIC ACID

Oxy di methyl succinic acid v Oxy adipic acid

Di oxy di methyl succinic acid $CO_2H CMe(OH) CMe(OH) CO_2H$ *Di methyl-racemic acid* [179°] Formed by saponifying its nitrile with HCl aq Crystals (containing aq) — KHA'' Plates, m sol hot water — CaA'' 1½aq — BaA'' 2aq small needles

Nitrile $CN CMe(OH) CMe(OH) CN$ [o 110°] Formed from di methyl di ketone (diacetyl) and HCy (Fittig, *A* 249, 208) Very hygroscopic needles, sl sol chloroform

Isomeride — Di-methyl tartaric acid

OXY METHYL-TEREPHTHALIC ACID $C_6H_4Me(OH)(CO_2H)_2$ [254° 1] [285°-290°] Formed by fusing $C_6H_4Me(SO_2NH_2)(CO_2H)_2$ with potash (Jacobsen & H Meyer, *B* 16, 191). Minute prisms (from very dilute alcohol), decomposed on fusion.

OXY - METHYL-THIAZOLE $\begin{smallmatrix} S C(OH) \\ CH CMe \end{smallmatrix} > N$.

Thiocyanacetone [102°] Formed by heating the carboxylic acid (Wohmann, *A* 259, 298), or by heating with dilute HCl aq the compound $CH_3CO CH_2SCy$, formed from chloro acetone and a sulphocyanide (Tscherniak a Norton, *C J* 44, 568, Hantzsch, *B* 20, 3127, 3337, 21, 941, *A* 249, 20) Needles Yields methyl thiazole on distilling with zinc-dust With phenylene *m* diamine it forms $C_6H_4(NH C_6H_4SN)_2$ [152°] Reacts with hydroxylamine

Di - oxy - methyl - thiazole $\begin{smallmatrix} S - CO \\ CH_2 CO \end{smallmatrix} > NMe$

Formed from di oxy thiazole, NaOMe, and MeI (Arapides, *A* 249, 28) Liquid, v e sol water Decomposed by boiling NaOH aq into methyl amine and thioglycollic acid PCl_5 yields $C_4H_4Cl_2NOS$ [161°]

OXY-METHYL THIAZOLE CARBOXYLIC

ACID $\begin{smallmatrix} S C(OH) N \\ C(CO_2H) CMe \end{smallmatrix}$ [222°] Formed from its ether, and also by heating chloro methyl thiazole carboxylic acid with conc KOH aq (Zurcher, *A* 250, 286, Wohmann, *A* 259, 296) Plates or needles (containing aq), v sol water — NH_4A' baq Prisms, decomposing at 138°

Ethyl ether EtA' [129°] Formed together with $(SC_2Me(CO_2Et)N)_2O$ [142°] from chloro-acetoacetic ether and ammonium or barium sulphocyanide (Hantzsch a Weber, *B* 20, 3132) Silky plates Hydroxylamine forms $C_{10}H_{10}O_4N_2S_2$ [α 217°]

OXY METHYL-THIOPHENE $\begin{smallmatrix} CMe CH \\ C(OH) CH \end{smallmatrix} <$

Formed from levulic acid and P_2S_5 (Kues, *B* 19, 555) Oil, sl sol water — $B'HOAc$ (210°) Yellow oil

OXY METHYL UREA *Di-ethyl derivative of the glycollyl derivative* $CH_2(OEt)NH CO NH CO CH_2OEt$ [80°]

Formed from $CH_2(OEt)CONH_2$, bromine, and potash (Hofmann, *B* 18, 2736) Needles (from water)

OXY-METHYL XANTHINE $C_6H_5N_4O_3$ *S* 16 at 16° Formed by the action of potassium cyanate on the hydrochloride of di oxy amido-methyl pyrimidine $CO < \begin{smallmatrix} NMe CH \\ NH CO \end{smallmatrix} > C NH_2$ (Behrend, *A* 231, 252, Lehmann, *A* 253, 80) Needles or prisms (containing 2aq), sl sol water Does not reduce alkaline $AgNO_3$

OXY-MYRISTIC ACID $C_{14}H_{28}O_3$ [51°] Occurs in the essential oil from the fruit of *Angelica archangelica* (Müller, *B* 14, 2480) Pearly plates (from alcohol) — KA' aq — CuA' — AgA' bulky pp *Benzoyl derivative* [68°] — AgA'

(1,2) DI OXY NAPHTHALENE

$C_{10}H_6(OH)_2$ [1 2] [60°] v (β) HYDRONAPHTHOQUINONE, vol n p 728

(1,4) Di-oxy-naphthalene [176°] v (α) HYDRONAPHTHOQUINONE

(1,4') Di-oxy-naphthalene [260°] Formed by potash fusion from (α) naphthol sulphonic acid and from naphthalene (1,4')-disulphonic acid (Erdmann, *A* 247, 356, Bernthsen, *B* 20, 938, Armstrong a Wynne, *C J Proc* 3, 43) Scales, v sol hot alcohol, sl sol water Yields crystalline $C_{10}H_6(OAc)_2$ [160°]

Pers di oxy-naphthalene $C_{10}H_6(OH)_2$ [1 17] [140°] Formed by potash fusion from the anhydride got by boiling naphthylamine *persulphonic* acid with water (Erdmann, *A* 247, 357) Needles or plates, sl sol water Yields $C_{10}H_6(OAc)_2$ [148°] crystallising in white plates

Di-oxy-naphthalene Made by reducing *pers* naphthoquinone with HOAc and zinc-dust (Mel dola a Hughes, *C J* 57, 631) Minute needles which blacken at 205° Yields $C_{10}H_6(OAc)_2$ [227°] According to theory, this body should be identical with the preceding

(1,2')-Dioxynaphthalene [178°] Formed by fusing (β) naphthol (α) sulphonic acid (Bayer, *B* 15, 1851) with potash (Emmert, *A* 241, 371) Needles, v e sol alcohol, m sol water Its alkaline solution blackens quickly $FeCl_3$ gives a blue pp Yields $C_{10}H_6(OAc)_2$ [108°] and $C_{10}H_6(OEt)_2$ [67°]

(1,3') Di-oxy naphthalene [135°] Made by potash fusion from naphthalene (1,3') di sulphonic acid (Ewer a Pick, *G P* 45,229 [1887]) or (β) naphthal (α) sulphonic acid (Claus, *J pr* [2] 39, 315) White prisms (from benzene) Turns red in air $FeCl_3$ gives a blue colour Yields $C_{10}H_6(OAc)_2$ [78°]

(2,3') Di-oxy naphthalene [216°] Formed by potash fusion from naphthalene (2,3') di sulphonic acid (Dusart, *Bl* [2] 8, 200, Darmstadter a Wichelhaus, *A* 152, 306) and from (β) naphthol (β) sulphonic acid (Schaffer, *A* 152, 298, Armstrong a Graham, *C J* 39, 140, Emmert, *A* 241, 369) Thin plates, v sol alcohol, sl sol cold water Gives a yellowish white pp with $FeCl_3$ Yields $C_{10}H_6(OAc)_2$ [175°] and $C_{10}H_6(OEt)_2$ [162°] Forms a disulphonic acid which gives $BaC_{10}H_6S_2O_6$ 2aq (Griess, *B* 13, 1959)

(2,2') Di oxy naphthalene [190°] Formed by potash fusion from naphthalene (2,2') di sulphonic acid (Ebert a Merz, *B* 9, 609, Weber, *B* 10, 1233, 14, 2206, Clausius, *B* 23, 519) Needles (from hot water) Its alkaline solution blackens in air Yields $C_{10}H_6(OAc)_2$ [136°], $C_{10}H_6(OBz)_2$ [139°], $C_{10}H_6(OMe)_2$ [134], and $C_{10}H_6(OEt)_2$ [104°] (Liebermann, *B* 15, 1428)

Dioxynaphthalene [161°] Got by fusing (β) naphthol (β) disulphonic acid (1 pt) with NaOH (4 pts) at 310°, or by heating with dilute (25 p c) H_2SO_4 at 200° (*E P* 15,803) Crystals (from water) $FeCl_3$ gives an intense blue colour Combines with diazo compounds

Tri oxy naphthalene $C_{10}H_3O_3$ Got by reducing oxy (α) naphthoquinone (Graebe, *A* 154, 324) Yellow needles (from ether) Very readily oxidised

DI-OXY-NAPHTHALENE DICARBOXYLIC ACID $C_{10}H_6(OH)_2(CO_2H)_2$ [162°] Formed by heating narcotic acid (Claus a Meixner, *J pr* [3] 37, 1) White needles, v sol ether Yields $C_{10}H_6(CO_2H)_2$ [253°] on reduction — NaA' 6aq — $NaHA'$ 5½aq, small needles — BaA' 2aq — AgA'

DIOXYNAPHTHALIC ACID $C_{10}H_6O_4$ [126°] Made by boiling chloro oxy naphthalic acid (v NAPHTHALENE) with baryta (Hermann, *A* 151, 67) Prisms — KHA' aq — BaA' 3aq — $Ba(HA')_2$ — $Ba(NH_4A')_2$ 2aq — $Ca(NH_4A')_2$ — $Cu(NH_4A')_2$ blue prisms

DI-OXY-NAPHTHAZINE $C_{10}H_6N_4O_2$ [α , 300°] Formed from di oxy naphthoquinone and naphthylene-(1, 2) diamine in alcohol (Nietzke

a. Hasterlik, B 24, 1839). Reddish brown needles, sl sol alcohol

(a) OXY (β)-NAPHTHOIC ACID

$C_{10}H_7(OH)CO_2H$ [182°] *Naphthol carboxylic acid* [187°], S 558 at 17° Formed by passing CO_2 over sodium α -naphthol at 100° (Eller, A 152, 277, Schaffer, A 152, 291, Nietzki & Gietermann, B 20, 1274), or by heating sodium- α -naphthol with liquid CO_2 at 130° (Schmitt & Burkard, B 20, 2699) Stellate groups of needles (from alcohol) $FeCl_3$ colours its solution blue PCl_5 forms $C_{10}H_7(OPOCl_2)Cl_2$ [115°], whence moist air forms $C_{10}H_7(OPO(OH)_2)CO_2H$, while alcohol yields $C_{10}H_7(OPO(OEt)_2)CO_2H$ [63°] (Wolfenstein, B 21, 1186) — NaA' 3aq — NaA' — NH_4A' — PhA' [96°]

Acetyl derivative $C_{10}H_7(OAc)CO_2H$ [158°]

Methyl ether MeA' [78°] Converted by hydroxylamine into $C_{10}H_7(OH)CONH(OH)$ [174°]

Ethylether EtA' [49°]

Phenyl ether PhA' [96°]

Methyl derivative of the anilide $C_{10}H_7(OMe)CONHPh$ [218°] Made from methyl α -naphthol, phenyl cyanate, and $AlCl_3$ (Leuchart & Schmitt, B 20, 2340) Prisms

(β) Oxy (α) naphthoic acid

$C_{10}H_7(OH)(CO_2H)$ [21] [157°] Formed, in like manner, from β -naphthol (S a B, cf Kauffmann, B 15, 804) Slender needles, decomposing at 125° when slowly heated Completely decomposed by boiling water into CO_2 and β -naphthol $FeCl_3$ gives a blue colour PCl_5 yields $C_{10}H_7(OPOCl_2)COCl$ [38°], whence moist air forms $C_{10}H_7(OPO(OH)_2)CO_2H$ [156°], while dilute alcohol produces $C_{10}H_7(OPO(OEt)_2)CO_2H$ [113°] (Rabe, B 22, 392) — NH_4A' yellow needles, m sol cold water

Methyl ether MeA' [76°] Converted by hydroxylamine into amorphous $C_{10}H_7(OH)CONH(OH)$ [178°] (Jeaurenaud, B 22, 1277)

Ethylether EtA' [55°]

Methyl derivative of the amide $C_{10}H_7(OMe)CONH_2$ [186°] Form from methyl β -naphthol, $AlCl_3$, and $ClCONH_2$ (Gietermann, A 244, 75) Crystals (from alcohol)

Ethyl derivative of the amide $C_{10}H_7(OEt)CONH_2$ [161°] Plates (from alcohol)

Methyl derivative of the anilide $C_{10}H_7(OMe)CONHPh$ [169°] Made from methyl β -naphthol and phenyl cyanate in presence of $AlCl_3$ (L a S) Needles

(β) Oxy-naphthoic acid

$C_{10}H_7(OH)CO_2H$ [23°] [216°] Formed in small quantity from sodium β -naphthol and CO_2 at 290° (Schmitt & Burkard, B 20, 2702) Plates, v sol alcohol Very stable $FeCl_3$ colours its solution blue Its methyl ether does not react with hydroxylamine

Pers-Oxy-naphthoic acid

$C_{10}H_7(OH)CO_2H$ [11°] [169°] Made by heating its internal anhydride for a long time with dilute KOHAq (Ekstrand, B 19, 1138, J pr [2] 38, 278) Small needles (from ether), v sol water Solutions of its salts are coloured violet by $FeCl_3$. — CaA' , 8½aq decomposed on boiling

Anhydride $C_{10}H_6\begin{smallmatrix} CO \\ \diagup \diagdown \\ O \end{smallmatrix}$ *Naphtholactone.*

[108°] Made from amido α -naphthoic acid by the diazo reaction Needles (from alcohol) or tables (from ether), insol cold aqueous alkalis

(a)-Oxy-naphthoic acid

$C_{10}H_7(OH)CO_2H$ [14°]

Ethyl derivative $C_{10}H_7(OEt)CO_2H$ [214°] Formed by the action of alcoholic potash on the amide $C_{10}H_7(OEt)CONH_2$ [244°], which is made from $C_{10}H_7OEt$ and $ClCONH_2$ in presence of $AlCl_3$ (Gietermann, B 23, 1198, A 244, 73) Needles, m sol alcohol — NaA' 3aq. plates — CaA' , 3aq needles, sl sol water

Methyl derivative $C_{10}H_7(OMe)CO_2H$ [232°] Formed, in like manner, from $C_{10}H_7(OMe)CONH_2$ [234°] which is prepared from $C_{10}H_7OMe$

(a)-Oxy (α) naphthoic acid [234°–237°]

Formed by fusing α -sulpho α -naphthoic acid [235°] with potash (Battershall, A 168, 114) Needles (from water), v sol alcohol Aqueous solutions of its salts blacken on boiling $FeCl_3$ gives a dirty violet pp

(a)-Oxy-(β) naphthoic acid [213°] Made by potash fusion from sulpho β -naphthoic acid (Needles from boiling water) Its salts are unstable $FeCl_3$ gives a dirty red pp

(β) Oxy (α) naphthoic acid [247°] Made by potash fusion from the sulpho naphthoic acid [218°–222°] (Stumpf, A 188, 6) Mass of needles (from water) $FeCl_3$ has no effect in the cold, but gives a brown pp on warming

(β) Oxy (α) naphthoic acid [187°] Made by potash fusion from sulpho-naphthoic acid [182°–185°] (Stumpf) Branching needles (from water) Gives a chocolate colour with $FeCl_3$. Yields β -naphthol on distilling with lime

(β) OXY NAPHTHOIC ALDEHYDE

$C_{10}H_7(OH)CHO$ *Aldehyde naphthol* [76°] Formed from β -naphthol, chloroform, and aqueous $NaOH$ (Kauffmann, B 15, 805, 16, 683). Prisms (from alcohol), almost insol water $FeCl_3$ colours its solution brown Reduces warm ammoniacal $AgNO_3$, forming a mirror With Ac_2O and $NaOAc$ it forms $C_{10}H_7(OAc)CH(OAc)$, [124°] Yields, on oxidation, oxy-naphthoic acid [150°] — $C_{10}H_7(ONa)CHO$ yellow plates

(a) OXY (α)-NAPHTHOPHENAZINE

$C_{18}H_{10}NO$ *α-Naphthophenazine* [43°] Formed by heating α -amido or diethylamido- α -naphthophenazine with conc $HClAq$ for 5 hours at 180° (Fischer & Hepp, B 23, 845, Eicker, B 23, 3805) Reddish yellow needles (from alcohol) The hydrochloride forms red prisms with green lustre

OXY-(β) NAPHTHOQUINOLINE $C_{18}H_{10}NO$

Formed by fusing β -naphthoquinoline sulphonic acid with potash (Gentil, B 18, 202) Does not melt below 250°

(a) OXY-(α)-NAPHTHOQUINONE $C_{18}H_{10}O_2$

Naphthalic acid [190°] Formed by heating amido- or oxy-naphthoquinonamide or oximido-naphthol with $HClAq$ or alkalis (Martius & Gries, A 134, 377, Graebe & Ludwig, A 154, 321, B 4, 970, Merz & Diehl, B 11, 1314, Kehrman, B 23, 2453) Formed also in like manner from phenyl-amido α -naphthoquinone and from phenyl amido- β -naphthoquinone (Baltzer, B 14, 1900, Zincke, B 14, 1496, Lie-

bermann & Jacobson, *A* 211, 80) Yellow needles, v sl sol cold water. May be sublimed

Reactions—1 Reduced by tin and HCl to tri oxy-naphthalene—2 The Na salt heated with *o*-phenylene-diamine forms (a) naphthoeurhodol (Kehrmann, *B* 23, 2453)—3 *Benzoin aldehyde* forms $\text{CHPh}(\text{C}_6\text{H}_4(\text{OH})\text{O})_2$ [211°–214°] (Zincke, *B* 21, 2203)—4 *Phenyl-hydrazine* yields $\text{C}_{10}\text{H}_7(\text{OH})\text{O}(\text{N}_2\text{HPh})$ [230°] which forms the salts CAa' , 4aq and BAa' , 10aq, the ethers MeA' [175°] and EtA' [173°], and the acetyl-derivative $\text{C}_{10}\text{H}_7\text{AcN}_2\text{O}_2$ [179°]. It reacts with aldehyde, benzoic aldehyde, and acetone, forming $\text{C}_{10}\text{H}_{12}\text{N}_2\text{O}_4$ [c 258°], $\text{C}_{10}\text{H}_{12}\text{N}_2\text{O}_4$, and $\text{C}_{10}\text{H}_{12}\text{N}_2\text{O}_4$ [245°–250°] respectively (Zincke & Thelen, *B* 17, 1812, 21, 2205)— $\text{AgC}_{10}\text{H}_7\text{O}_2$ scarlet needles (from hot water)

Ethyl ether $\text{EtC}_{10}\text{H}_7\text{O}_2$ [127°] Needles

Imide v AMIDO NAPHTHOQUINONE

Amide v PHENYL AMIDO NAPHTHOQUINONE

Oxim $\text{C}_{10}\text{H}_7(\text{NOH})(\text{OH})\text{O}$ [1 2 4] *Nitroso naphtharesorcin* Formed from oxy-naphthoquinone and hydroxylamine (Kostanecki, *B* 22, 1843) Needles, decomposing at 180° Gives with ferrous salts a green, and with ferric salts a dark-brown lake The hydrochloride forms yellow needles Nitrous acid forms $\text{C}_{10}\text{H}_7(\text{NOH})_2\text{O}_2$ crystallising in plates (containing aq), decomposing at 165°

Oxy-(β)-naphthoquinone $\text{C}_{10}\text{H}_6(\text{OH})\text{O}_2$ [2' 1 2] Formed by oxidising its oxim with FeCl_3 . Reddish brown, amorphous pp, v sol alcohol

Oxim $\text{C}_{10}\text{H}_7(\text{OH})(\text{NOH})\text{O}$ [2' 1 2] [235°] Formed from (2,2') di oxy naphthalene and nitrous acid (Clausius, *B* 23, 521) Brownish-yellow needles

Di-oxy-naphthoquinone $\text{C}_{10}\text{H}_6(\text{OH})_2\text{O}_2$ *Naphthazarin* Formed by heating di nitro naphthalene [214°] with zinc and H_2SO_4 at 200° (Roussin, De Aguiar & Baeyer, *B* 4, 251, 438, Liebermann, *B* 3, 905, *A* 162, 330) Reddish brown needles with green lustre, v sl sol boiling water, v sol alcohol Its alkaline solution is blue Its solution in H_2SO_4 is crimson Baryta- and lime- waters give violet blue pps

Isomeride v Oxy juglone, p 101

Di oxy-naphthoquinone $\text{C}_{10}\text{H}_6(\text{OH})_2\text{O}_2$ Formed by heating oxy amido naphthoquinone with HClAq at 175° (Merz & Diehl, *B* 11, 1322) Reddish-brown needles (from alcohol) Its alkaline solution is brownish-violet Dyes violet with alumina mordants and dark blue with iron mordants Dyes silk brownish violet with metallic lustre It forms a crystalline di acetyl derivative— BAa'' — PbA'' — $\text{Ag}_2\text{A}''$. greenish-blue pp

Tri-oxy-naphthoquinone A black by product in preparing naphthazarin (Aguar, *B* 4, 439)

References—Bromo- and Chloro- OXY NAPHTHOQUINONE

OXY-NAPHTHOQUINONE SULPHONIC

ACID $\text{C}_{10}\text{H}_6(\text{OH})\text{O}_2\text{SO}_3\text{H}$ Formed by the action of alkalis on the acid $\text{C}_{10}\text{H}_6(\text{OH})(\text{SO}_3\text{H})(\text{OSO}_3\text{H})_2$ got by dissolving dichloro naphthoquinone in aqueous KHSO_4 (Graebe, *A* 149, 9)— $\text{K}_2\text{A}'$ (dried at 140°) orange crystals

References—Bromo- and Chloro- OXY NAPHTHOQUINONE SULPHONIC ACID

(β)-OXY- α NAPHTHOYL-BENZOIC ACID $\text{C}_{10}\text{H}_6(\text{OH})\text{CO C}_6\text{H}_4\text{CO}_2\text{H}$ *Phenyl oxy-naphthoyl-ketone-o-carboxylic acid* [256°] Fre-

pared by oxidising an alkaline solution of ($\beta\beta$ -dioxo-dinaphthyl with KMnO_4 (Walder, *B* 16, 299) Silky prisms Sol alcohol, ether, and acetic acid, nearly insol water

Reactions—1 On fusion with KOH gives (β) naphthol and phthalic acid—2 Heated with HI it is reduced to (β -oxy ω naphthyl *o*-toluic acid ($\text{C}_6\text{H}_4(\text{OH})\text{CH}_2\text{C}_6\text{H}_4\text{CO}_2\text{H}$)—3 Heated with ZnCl_2 to 220° it yields a substance $\text{C}_{10}\text{H}_6\text{O}$, which forms colourless crystals [146°], easily soluble in alcohol—4 Heated with resorcin it gives a red colouring matter, and with dimethylaniline and ZnCl_2 a green

Salts— $\text{A}'\text{Na}$ white crystalline powder, sl sol cold water and alcohol— $\text{A}'\text{Ag}$ insoluble pp— $\text{A}'\text{Ba}$ 2aq slightly soluble white pp

Methyl ether $\text{A}'\text{Me}$ [199°], prisms

Ethyl ether $\text{A}'\text{Et}$ [206°], needles

Acetyl derivative $\text{C}_{10}\text{H}_6(\text{OAc})$ [170°]

DI (α) OXY DINAPHTHYL

$\text{C}_{10}\text{H}_6(\text{OH})\text{C}_6\text{H}_4(\text{OH})$ *Dinaphthol* [300°] Formed by adding FeCl_3 to a conc solution of (α) naphthol (Dianin, *B* 6, 1252, 7, 125, 187, Julius, *Chem Ind* 10, 97) Silvery plates (from alcohol), insol water Its alcoholic solution is coloured red by FeCl_3

Benzoyl derivative $\text{C}_{20}\text{H}_{12}(\text{OBz})_2$ [253°]

Di-methylether $\text{C}_{20}\text{H}_{12}(\text{OMe})_2$ [251°]

Di-ethyl ether $\text{C}_{20}\text{H}_{12}(\text{OEt})_2$ [211°] Pearly plates (Osteimayer & Rosenhek, *B* 17, 2453)

Di-(β)-oxy (aa) dinaphthyl [218° cor] *V D* 9 52 (obs) Prepared by oxidation of an ethereal solution of (β) naphthol with FeCl_3 , the yield being 60 p c of the theoretical amount (Dianin, Walder, *B* 15, 2166) Needles or plates Gives with FeCl_3 a greenish colour, becoming red on heating On heating with ZnCl_2 it gives (β) dinaphthylene oxide H_2SO_4 forms $\text{C}_{20}\text{H}_{12}(\text{OH})_2(\text{SO}_3\text{H})_2$, whence BAa'' 6aq, $\text{C}_{20}\text{H}_{12}(\text{OH})_2(\text{SO}_3\text{H})_2$ is also formed ZnCl_2 and NH_3 at 330° form $\text{C}_{20}\text{H}_{12}\text{N}$ [157°], which forms a picrate [217°] and an acetyl derivative [144°]—*Picrate* $\text{C}_{20}\text{H}_{12}\text{O}_2\text{C}_6\text{H}_4(\text{NO}_2)_3\text{OH}$ [174°] White crystals, sol alcohol

Benzoyl derivative $\text{C}_{30}\text{H}_{18}(\text{OH})(\text{OBz})$ [204°]

Di benzoyl derivative $\text{C}_{20}\text{H}_{12}(\text{OBz})_2$ [160°]

Di methylether $\text{C}_{20}\text{H}_{12}(\text{OMe})_2$ [190°]

Di-ethylether $\text{C}_{20}\text{H}_{12}(\text{OEt})_2$ [90°]

Di-oxy dinaphthyl [195°] A product of the action of melted potash on (β) naphthoaldehyde (Kauffmann, *B* 15, 807) Silky needles (from alcohol)

Tetra oxy-dinaphthyl $\text{C}_{20}\text{H}_{10}(\text{OH})_4$ '*Di-naphthyl dihydroquinone*' [178°] Formed by reduction of the quinone $\text{C}_{20}\text{H}_{10}\text{O}_4$ (Stenhouse & Groves, *C J* 33, 415), and by the action of tin and HCl on (β) naphthoquinone (Korn, *B* 17, 3024) Colourless needles, darkening by oxidation in air, insol water, v sol HOAc The quinone $\text{C}_{20}\text{H}_{10}\text{O}_4$ is oxidised in alkaline solution by the air to [4 3 2 1] $\text{C}_{10}\text{H}_6\text{O}_2(\text{OH})\text{C}_{10}\text{H}_6\text{O}_2(\text{OH})$ [1 4 3 2] [245°–250°]

Acetyl derivative $\text{C}_{20}\text{H}_{10}(\text{OAc})_4$ [166°]

α OXY-(α)-NAPHTHYL-ACETIC ACID

$\text{C}_{11}\text{H}_{10}\text{O}_2$, *c c* $\text{C}_{11}\text{H}_9\text{CH}(\text{OH})\text{CO}_2\text{H}$ (α)-*Naphthylglycolic acid* [98°] Formed by reducing (α)-naphthyl glyoxylic acid with sodium amalgam (Boessneck, *B* 16, 640) Formed also by saponifying its nitrile (Brandis, *B* 22, 2158), and

from (a) naphthyl methyl ketone by successive treatment with bromine and potash (Schweizer, *B* 24, 549) Rosettes of needles (from water) Yields (a) naphthoic aldehyde on oxidation with dilute HNO_3 . — BaA' . — AgA' pulverulent pp

Methyl ether MeA' [79°] Needles

Nitrile $\text{C}_{10}\text{H}_7\text{CH}(\text{OH})\text{CN}$ Formed from (a)-naphthoic aldehyde, KC_2H_5 , and cold HClAc Oil, v sol alcohol

a-Oxy-(β) naphthyl-acetic acid [158°] (S), [c 176°] (C & T) Prepared in the same way as its isomeride (Claus & Tersteegen, *J pr* [2] 42, 518, Schweizer, *B* 24, 547) Small needles (from water) Reduced by HIAc to $\text{C}_{10}\text{H}_7\text{CH}_2\text{CO}_2\text{H}$ [c 142°]

Methyl ether MeA' [75°] Needles

Ethyl ether EtA' [87°] Needles

Acetyl derivative $\text{C}_{10}\text{H}_7\text{CH}(\text{OAc})\text{CO}_2\text{H}$ [150°]

Amide $\text{C}_{10}\text{H}_7\text{CH}(\text{OH})\text{CONH}_2$ [228°]

Oxy di-naphthyl acetic acid $\text{C}_{10}\text{H}_7\text{O}_2$ & $\text{C}_{10}\text{H}_7\text{O}_2$ $\text{C}(\text{OH})\text{CO}_2\text{H}$ Yellowish green tables, decomposing about 100°

Anhydride $\text{C}_{20}\text{H}_{12}\text{O}_2$ Formed by heating (a) naphthol with oxalic acid and H_2SO_4 (Hoenig, *M* 1, 251) Pale red plates, sl sol chloroform

(β) OXY-NAPHTHYL-ACRYLIC ACID
 $\text{C}_{10}\text{H}_7(\text{OH})\text{CH}=\text{CHCO}_2\text{H}$ [170°] Formed by heating its anhydride with aqueous potash at 170° Crystalline powder, sol alcohol

Anhydride $\text{C}_{10}\text{H}_7\text{C}(\text{CH}=\text{CH})\text{CO}$ (β)-Naphtho-coumarin [118°] Prepared by heating (β)-oxy-naphthoic aldehyde (2 pts) with Ac_2O (10 pts) and NaOAc (2 pts) at 180° (Kauffmann, *B* 16, 685) Needles, v sol alcohol, sl sol hot water

An isomeride of this anhydride [141°] is got by heating (β) naphthol with malic acid and H_2SO_4 (Pechmann & Welsh, *B* 17, 1651)

OXY-DINAPHTHYLAMINE v. Imido di-NAPHTHYL OXIDE

Tri oxy (β) naphthylamine

$\text{C}_{10}\text{H}_7(\text{OH})_3(\text{NH}_2)_3$ [1342] Formed from nitro oxy quinone, SnCl_4 , and HCl (Kehrmann & Weichardt, *J pr* [2] 40, 181) Brownish violet needles — B'HCl monoclinic prisms

Tetra-acetyl derivative [145°]

(β) OXY-NAPHTHYL-CROTONIC ACID

Anhydride $\text{C}_{10}\text{H}_7\text{C}(\text{CMe}=\text{CH})\text{CO}$ [162°] Formed by the action of H_2SO_4 on a mixture of acetoacetic ether and (β) naphthol (Pechmann & Cohen, *B* 17, 2190) Needles, sol alcohol Its solution in H_2SO_4 exhibits green fluorescence

DI-OXY-NAPHTHYLENE-DIAMINE

$\text{C}_{10}\text{H}_4(\text{OH})_2(\text{NH}_2)_2$ [1824] Formed by the action of SnCl_4 and HCl on the oxim of oxy-amido naphthoquinone (Kehrmann & Weichardt, *J pr* [2] 40, 186) Its alkaline solutions become crimson on oxidation, and then blue on boiling — B'HCl white needles

TRI-OXY-TRI-NAPHTHYL ETHANE

$\text{C}_{10}\text{H}_7(\text{OH})\text{CH}_2\text{CH}(\text{C}_{10}\text{H}_7\text{OH})_2$ Formed from di-chloro di-ethyl oxide and (a) naphthol (Wislicenus, *A* 243, 165) White amorphous powder, insol water, sol alcohol

(a) OXY-NAPHTHYL ETHYL KETONE

$\text{C}_{10}\text{H}_7(\text{OH})\text{CO}\text{C}_2\text{H}_5$ [81°] Formed by heating (a)-naphthol with propionic acid and ZnCl_2 at 178° Pale yellow plates (from ether or alcohol).

Phenyl hydrazide [128°].

Methyl ether [58°] Formed from propionyl chloride, $\text{C}_{10}\text{H}_7\text{OMe}$, and AlCl_3 (Gattermann, *B* 23, 1209) Prisms (from ether). Yields an oxim $\text{C}_{10}\text{H}_7(\text{OMe})\text{C}(\text{NOH})\text{C}_2\text{H}_5$ [172°].

OXY-NAPHTHYL METHYL KETONE

[18] $\text{C}_{10}\text{H}_7(\text{OH})\text{CO}\text{CH}_3$, 'Ketonaephthol' [168°] Formed by the rapid distillation of phenyl aceto isocrotonic acid (Erdmann, *A* 254, 197) Crystals (from dilute alcohol)

Acetyl derivative $\text{C}_{10}\text{H}_7\text{O}_2$ [109°]

Oxim $\text{C}_{10}\text{H}_7\text{NO}_2$ [174°]

Oxy naphthyl methyl ketone [103°] Formed by heating (a)-naphthol with HOAc and ZnCl_2 (Witt, *B* 21, 321) Pale-green six sided prisms (from benzene) Its alkaline salts are yellow Alcoholic NH_3 at 200° forms

$\text{C}_{10}\text{H}_7(\text{OH})\text{C}(\text{NH}_2)\text{CH}_3$ [203°]

Oxim $\text{C}_{10}\text{H}_7(\text{OH})\text{C}(\text{NOH})\text{CH}_3$ [170°]

(a) **Oxy-naphthyl methyl ketone** Probably identical with the preceding isomeride *Methyl ether* $\text{C}_{10}\text{H}_7(\text{OMe})\text{CO}\text{CH}_3$ [72°] (above 350°) Formed from $\text{C}_{10}\text{H}_7(\text{OMe})$, AcCl , and AlCl_3 (Gattermann, *B* 23, 1208) Six sided tables

Ethyl ether $\text{C}_{10}\text{H}_7(\text{OMe})\text{Ac}$ [79°]

(β) **Oxy naphthyl methyl ketone** *Methyl ether* $\text{C}_{10}\text{H}_7(\text{OMe})\text{CO}\text{CH}_3$ [58°] Formed from $\text{C}_{10}\text{H}_7\text{OMe}$, AcCl , and AlCl_3 (G) Slender needles

Ethyl ether [63°] Compact tables

OXY (a) NAPHTHYL METHYL PYRAZOLE

$\text{C}_{10}\text{H}_{12}\text{N}_2\text{O} \rightleftharpoons \text{C}_{10}\text{H}_7\text{N}(\text{C}(\text{OH})\text{CH}=\text{CMe})_2$ [c 190°] Formed by heating (a) naphthyl hydrazine with acetoacetic ether at 130° (Knorr, *B* 17, 551)

Yields a methyl derivative [129°] (a) Naphthyl hydrazine and thio acetoacetic ether form $\text{C}_{10}\text{H}_{12}\text{N}_2\text{O}$ [220°] (Sprague, *C J* 59, 343)

Oxy-(β) naphthyl methyl-pyrazole [190°]

Formed in the same way Yields a methyl derivative [129°]

(a) **OXY-NAPHTHYL PROPYL KETONE**

$\text{C}_{10}\text{H}_7(\text{OH})\text{CO}\text{C}_3\text{H}_7$ [78°] Formed by heating butyric acid with (a) naphthol and ZnCl_2 (Goldzweig, *J pr* [2] 43, 97) Silky needles (from ether), m sol hot water

DI-OXY-DI-(a)-NAPHTHYL-PYRAZINE HEXAHYDRIDE

$\text{C}_{10}\text{H}_7\text{N}(\text{C}(\text{OH})\text{CH}=\text{C}(\text{OH})\text{CH}_2)_2\text{NC}_{10}\text{H}_7$ [275°]

Formed by boiling the acetyl derivative of chloro (a) naphthylamine with alcoholic potash (Abenius, *J pr* [2] 40, 437) Formed also by heating (a)-naphthylamido acetic acid with Ac_2O at 200° (Bischoff, *B* 22, 1807, 23, 2003) Plates (from HOAc), insol alkalis

Di-oxy-di (β) naphthyl-pyrazine hexahydride [above 360°] Formed by heating (β)-naphthyl amido acetic acid at 220° in hydrogen, and by the action of chloro acetic ether on the naphthalide of (β) naphthylamido acetic acid (Bischoff) Plates, sol aniline Converted by boiling alcoholic potash into

$\text{C}_{10}\text{H}_7\text{NHCH}_2\text{CO}\text{N}(\text{C}_{10}\text{H}_7)\text{CH}_2\text{CO}_2\text{H}$ [105°]

DI-OXY-DI (β) NAPHTHYL SULPHIDE

$\text{S}(\text{C}_{10}\text{H}_7\text{OH})_2$, *Naphthol sulphide* [215°] Formed, together with the di- and tri-sulphides, by the action of sulphur at 180° in presence of PbO , or of S_2Cl_2 in benzene, upon (β)-naphthol (Tassinari, *G* 17, 94, Onufrovitch, *B* 21, 3559, 23, 3355). Prisms, insol water, ether, and

benzene, m sol hot alcohol Yields di oxy di-naphthyl on heating with powdered copper NH_4Aq forms, on heating, (8) naphthylamine Alcohol and ammoniacal AgNO_3 forms $\text{C}_{20}\text{H}_{12}\text{SO}_2$ [164°] —NaA' 6aq needles

Acetyl derivative $\text{SC}_{20}\text{H}_{12}(\text{OAc})_2$ [154°]

Benzoyl derivative [208°] Plates

Ethyl ether $\text{SC}_{20}\text{H}_{12}(\text{OEt})_2$ [189°] Yields a di-nitro derivative [235°]

Di-oxy di (8) naphthyl disulphide

$\text{S}_2(\text{C}_{10}\text{H}_7\text{OH})_2$ [169°] Formed as above Thin yellow needles, m sol benzene Yields di oxy dinaphthyl on heating with Cu at 240°

Di-acetyl derivative $\text{S}_2\text{C}_{20}\text{H}_{12}(\text{OAc})_2$

[c 140°] Yellowish crystalline mass

Di benzoyl derivative [187°] Prisms

Di-ethyl ether [158 5°] Needles

Di oxy di-naphthyl disulphide $\text{S}_2(\text{C}_{10}\text{H}_7\text{OH})_2$ [210°] Formed by heating (8) naphthol with NaOHAq and S (Lange, B 21, 260) Needles, insol water, v sol HOAc It is accompanied by the preceding isomeride [170°]

Di-oxy di-(8)-naphthyl trisulphide

$\text{S}_3(\text{C}_{10}\text{H}_7\text{OH})_2$ Formed as above (Onufrovitch) Powder Yields a dibenzoyl derivative [194°]

ν OXY-(α)-NAPHTHYL-THIO UREA

$\text{C}_{10}\text{H}_7\text{NH CS NH}(\text{OH})$ [116°] Formed from (α) naphthyl thiocarbimide and hydroxylamine (Tiemann, B 22, 1939, Voltmer, B 24, 382) Pearly plates (from alcohol), insol water Hot alcohol converts it into (α) naphthyl cyanamide

Benzyl ether $\text{C}_{10}\text{H}_7\text{NH CS NH}(\text{OCH}_2\text{Ph})$ [133°] Formed by using benzyl hydroxylamine

(8)-OXY ω NAPHTHYL o TOLUIC ACID

$\text{C}_{10}\text{H}_7(\text{OH})-\text{CH}_2-\text{C}_6\text{H}_4\text{CO}_2\text{H}$ [261°] Formed by reduction of oxy naphthoyl benzoic acid by heating with HI and P to 200° (Waldner, B 16, 304) Small colourless prisms Nearly insol water, sl sol alcohol and ether —A'Ag flocculent pp

OXY-NITRO v NITRO OXY

DI-OXY-OCTANE $\text{C}_8\text{H}_{16}\text{O}_2$ 2e

$\text{CPrH}(\text{OH})\text{CPrH}(\text{OH})$ Mol w 146 [51 5°] (228°) Formed by the action of alcoholic potash on isobutyric aldehyde (Fossek, M 4, 664, 11, 397) Monoclinic tables, v sol alcohol Combines with CaCl_2 Yields isobutyric and oxalic acids on oxidation by HNO_3 Boiling dilute H_2SO_4 forms a (8) pinacolin $\text{C}_{10}\text{H}_{18}\text{O}_2$ (162°) Conc H_2SO_4 in the cold forms an (α)-pinacolin $\text{C}_8\text{H}_{16}\text{O}$ (122°) smelling like camphor

Acetyl derivative $\text{C}_8\text{H}_{16}(\text{OAc})_2$ (235°)

Di oxy-octane $\text{C}_8\text{H}_{16}(\text{OH})_2$ *Octylene glycol* (235°–240°) SG ^a 932 Obtained from octylene bromide by successive treatment with AgOAc and solid KOH (De Clermont, C R 59, 80) Oil, sol alcohol

Acetyl derivative $\text{C}_8\text{H}_{16}(\text{OAc})_2$ (245°–250°)

Di-oxy-octane $\text{CMeEt}(\text{OH})\text{CMeEt}(\text{OH})$

Methyl ethyl-pinacone [c 28°] (200°–205°) A product of the action of sodium amalgam on methyl ethyl ketone (Lavrinovitch, A 185, 124) Crystalline mass, m sol water

Tri-oxy-octane

$\text{CMePr}(\text{OH})\text{CH}_2\text{CH}(\text{OH})\text{CH}_2\text{OH}$ (210° at 60 mm) Formed by the action of HOCl followed by KOH upon methyl allyl propyl carbinol (Reformatsky, J pr [2] 40, 412) Thick

liquid, v. sol water and alcohol, sl sol alcohol. Yields a liquid tri acetyl derivative

Tetra oxy-octane $\text{C}_8\text{H}_{16}(\text{OH})_4$ 2e

$\text{CH}_2(\text{OH})\text{CMe}(\text{OH})\text{CH}_2\text{CH}_2\text{CMe}(\text{OH})\text{CH}_2\text{OH}$ [154°] Formed by the action of water on the liquid $\text{C}_8\text{H}_{16}\text{O}_2$ (170°–180° at 125 mm) which is produced by the action of potash on $\text{C}_8\text{H}_{16}\text{ClO}(\text{OH})$, a body formed by the addition of oxygen and HOCl to $\text{CH}_2\text{CMeCH}_2\text{CH}_2\text{CMeCH}_2$ (Przybytek, B 20, 3242)

DI OXY OCTENOIC LACTONE

$\text{CH}_2\text{CHCH}_2\text{C}(\text{OH})\begin{smallmatrix} \text{CH}_2 \\ \text{CO} \end{smallmatrix}\begin{smallmatrix} \text{CHMe} \\ \text{O} \end{smallmatrix}$ Formed by the action of conc H_2SO_4 on $(\text{C}_6\text{H}_5)_3\text{C}(\text{OH})\text{CO}_2\text{H}$ [48°] (Bultsch, J pr [2] 39, 94) Yellow syrup Yields amorphous $\text{Ba}(\text{C}_8\text{H}_{13}\text{O}_4)_2$

OXY OCTINOIC ACID $\text{C}_8\text{H}_{13}\text{O}_4$ 2e

$\text{C}(\text{C}_6\text{H}_5)_3(\text{OH})\text{CO}_2\text{H}$ '*Diallyloxalic acid*' [48°] Formed by saponifying its ether, which is made by the action of zinc and allyl iodide on oxalic ether (Saytzeff, A 185, 183, Schatzky, J pr [2] 34, 485, Bl [2] 45, 183) Needles, m sol water, v sol alcohol and ether —NaA' 2aq —LiA' aq —BaA' 2aq —ZnA' 1¹/₂ aq S (of ZnA') 28 at 22° —CdA' aq —PbA' 2aq —CuA' minute green needles

Ethyl ether EtA' (213 6° cor) SG ^g 9873, ^h 9718

Methyl derivative $\text{C}(\text{C}_6\text{H}_5)_3(\text{OMe})\text{CO}_2\text{H}$ Yields the salts BaA' 2aq —PbA' 1¹/₂ aq —CuA' aq and the ether EtA' (216°) SG ^g 9607 $\mu_s = 1.4556$ $R_\infty = 89.52$ (Barataeff, J pr [2] 35, 1)

Heptic acid [151°], obtained by the action of bromine and potash on isobutyl acetoacetic ether (Demarçay, A Ch [5] 20, 472), is perhaps an oxyoctinoic acid It yields CaA' 5aq, crystals in needles

α OXY-OCTOIC ACID $\text{C}_8\text{H}_{15}\text{O}_4$ 2e

$\text{CPr}_2(\text{OH})\text{CO}_2\text{H}$ [73°] (K' a S), [81°] (R) Formed by the action of zinc and propyl iodide on oxalic ether (Rafalsky, B 14, 2068, Bn 1, 527) Formed also by heating butyrom with conc KOHAq (Klinger a Schmitz, B 24, 1273) Needles —KA' —BaA' scales

Ethyl ether EtA' (209° cor) Oil

α-Oxy-octenoic acid $\text{CPr}_2(\text{OH})\text{CO}_2\text{H}$ [111°]

Formed from its ether, which is made from oxalic acid, isopropyl iodide, and zinc (Markownikoff, Z [2] 6, 516) Needles (from water), volatile with steam —BaA' 3aq, small efflorescent needles

Ethyl ether EtA' (203° cor) Oil

α-Oxy octoic acid

$\text{CH}_2\text{PrCH}_2\text{CH}_2\text{CH}(\text{OH})\text{CO}_2\text{H}$ [69 5°] Formed from canthanol and HCl, followed by boiling HClAq (Erlenmeyer a Sigel, B 7, 697, 1108, A 177, 102) Plates, v sl sol water Split up by HClAq at 120° into canthanol and formic acid. —AgA'

Ethyl ether EtA' (230°) at 715 mm

Amide [150°] Satiny plates

Nitrile $\text{C}_8\text{H}_{11}\text{CH}(\text{OH})\text{CN}$ SG ^h 905

Formed from canthanol and anhydrous HCl in the cold Liquid, sl sol water

α-Oxy-octoic acid $\text{CMe}_2\text{CH}_2\text{CMe}(\text{OH})\text{CO}_2\text{H}$

[107°] A product of the oxidation of 'isodibutylene' with KMnO_4 (Butleroff, J R 1892, 190, O J 42, 936, Bl [2] 88, 554) Needles or prisms, v sol hot water —AgA' needles

β -Oxy octoic acid $\text{CH}_2\text{CH}(\text{OH})\text{CET}_2\text{CO}_2\text{H}$
 Formed by reducing di ethyl acetoacetic ether with sodium amalgam and water (Schnapp, *A* 201, 62) Hygroscopic syrup, v sol alcohol, m sol water Decomposed on distillation into aldehyde and di ethyl acetic acid $-\text{NaA}', 6\text{aq} - \text{CaC}_2\text{H}_3\text{O}_2 - \text{AgA}'$ flocculent pp

β Oxy octoic acid $\text{CH}_2\text{CH}(\text{OH})\text{CMePr CO H}$
 Formed by reducing methyl propyl acetoacetic ether (Jones, *A* 226, 288) Split up on distillation into aldehyde and methyl propyl acetic acid $-\text{ZnA}_2$ crusts

β Oxy octoic acid $\text{CPrH}(\text{OH})\text{CMe CO H}$ [108°] Formed by reducing isobutyryl isobutyric ether (Wohlbruck, *B* 20, 2334, Hantzsch, *A* 249, 54) Prisms, v sol ether $-\text{BaA}', 3\text{aq}$

γ Oxy octoic acid $\text{CET}_2(\text{OH})\text{CH}_2\text{CH}_2\text{CO}_2\text{H}$
 Formed from succinyl chloride and ZnEt_2 (Wischnin, *A* 143, 262) $-\text{CaA}', 2\text{aq} - \text{BaA}'$

Lactone $\text{C}_8\text{H}_{14}\text{O}_2$ (228°-233°) (Emmert, *B* 15, 1852) Oil, v sol alcohol and ether

γ Oxy-octoic acid
 $\text{CH}_2\text{CH}(\text{OH})\text{CMeH CETH CO}_2\text{H}$ Formed by reducing acetyl hexoic acid (Young, *C J* 43, 177, *A* 216, 43) $-\text{BaA}'_2$ amorphous

Lactone $\text{C}_8\text{H}_{14}\text{O}_2$ (227°)

γ Oxy-octoic acid **Lactone** $\text{C}_8\text{H}_{14}\text{O}_2$ (134° at 36 mm) A product of the action of iodoforn or iodine on NaOCH_2Pr (Gorboff, *J pr* [2] 41, 236, 261) Liquid

Oxy octoic acid [92°] A product of the action of alcoholic potash on isobutyric aldehyde (Fossek, *M* 4, 676) Crystals $-\text{CaA}'_2$ sol water

Di oxy-octoic acid
 $\text{CHMe}(\text{OH})\text{C}(\text{CH Pr})(\text{OH})\text{CO H}$ A product of the action of alcoholic potash on bromo isobutyl acetoacetic ether (Demarçay, *C R* 86, 1135) Liquid, m sol water $-\text{BaA}'$ needles $-\text{CaA}'$

Di oxy octoic acid
 $(\text{CHMe}(\text{OH})\text{CH})_2\text{CH CO H}$ Formed by boiling barium di oxy di propyl malonate with water (Hjelt, *A* 216, 70) **Anhydride** $\text{C}_8\text{H}_{14}\text{O}_2$ Liquid, v e sol water

Tri oxy octoic acid **Lactone**
 $\text{CH}_2\text{CH}(\text{OH})\text{CH C}(\text{OH})\text{CH}_2\text{CHMe}$ Formed by the action of H_2SO_4 on a oxy octoic acid [47°] (Bulitsch, *J pr* [2] 39, 89) Syrup, sol alcohol $-\text{Ba}(\text{C}_2\text{H}_3\text{O}_2)_2$

Tetra oxy octoic acid $\text{C}_8\text{H}_{14}\text{O}_6$ etc
 $(\text{CH}_2(\text{OH})\text{CH}(\text{OH})\text{CH}_2)_2\text{CH CO}_2\text{H}$ Formed by boiling tetra oxy di propyl malonic acid with baryta water (H) Readily changes to the only anhydride $\text{C}_8\text{H}_{14}\text{O}_6$

Penta-oxy-octoic acid
 $(\text{CH}_2(\text{OH})\text{CH}(\text{OH})\text{CH}_2)_3\text{C}(\text{OH})\text{CO}_2\text{H}$
Lactone $\text{C}_8\text{H}_{14}\text{O}_6$ Formed by heating α -oxy-octoic ('diallyl oxalic') acid with nitric acid on a water bath (Bulitsch, *J pr* [2] 39, 65) Yellow liquid, v sol water and alcohol, insol ether

References $-\text{Bromo-}$ and **Di chloro oxy-octoic acid**

DI-OXY OCTYLENE $\text{C}_8\text{H}_{14}(\text{OH})_2$ **Conylene-glycol** **Acetyl derivative** $\text{C}_8\text{H}_{14}(\text{OAc})_2$ (225°) $\text{SG } 12.5$ 9887 Formed by conylene bromide and AgOAc (Wertheim, *A* 180, 298) Oil

OXY-OLEIC ACID $\text{C}_{18}\text{H}_{34}\text{O}_2$ An oily product of the action of Ag_2O on dibromostearic acid (Burg a Overbeck, *A* 140, 69) An isomeric acid [58°]

is formed by the action of H_2SO_4 on oleic acid (Liechti a Suida, *B* 16, 2455)

OXY-OXALACETIC ETHER **Ethyl derivative** $\text{CO}_2\text{Et CH}(\text{OEt})\text{CO CO}_2\text{Et}$ (155° at 17 mm) Formed by the action of sodium ethylate on a mixture of oxalic ether and $\text{CH}_3(\text{OEt})\text{CO}_2\text{Et}$ in ether (Wislicenus a Scheidt, *B* 24, 432) Thick liquid, miscible with alcohol and ether

α OXY-PALMITIC ACID $\text{C}_{16}\text{H}_{32}\text{O}_2$ [83°] Formed by boiling α bromo palmitic acid with alcoholic potash (Hell a Jordanoff, *B* 24, 939) Small crystals (from alcohol)

Acetyl derivative [62.5°] **Scale**

OXYPARACONIC ACID $\text{C}_8\text{H}_{14}\text{O}_2$ An acid formed by boiling calcium chlorotamalatate with water (Morawski, *J pr* [2] 11, 450) Thick liquid $-\text{CaA}'_2$ 2aq rhombohedral

TETRA OXY PENTAMETHENYL HYDRIDE
 $\text{CH}(\text{OH})\text{CH}(\text{OH})\text{CH}(\text{OH})\text{CH}(\text{OH})$ Formed by the action of HOAc on the Ba salt of its carboxylic acid (Hantzsch, *B* 20, 2792) Oil $-\text{BaA}' 3\frac{1}{2}\text{aq}$ amorphous powder, almost insol water

Carboxylic acid $\text{CH}(\text{OH})\text{CH}(\text{OH})\text{CH}(\text{OH})\text{CH}(\text{OH})\text{CO}_2\text{H}$
 Formed from ammonium tri chloro di oxy hexenoic acid and baryta water at 60° $-\text{BaA}'_2$ 4aq bulky pp, insol water

OXY PENTANE v **AMYL ALCOHOL**
Di-oxy pentane $\text{CH}_2(\text{CH CH}_2\text{OH})_2$ Mol w 104 (260°) Formed from pentamethylenediamine by HNO_2 (Gustavson, *J pr* [2] 39, 542)

Di oxy-pentane
 $\text{CHMe}(\text{OH})\text{CH}_2\text{CH}_2\text{CH}_2(\text{OH})$ (220°) at 713 mm $\text{SG } 1.0003$ Formed by reducing aceto-propyl alcohol by sodium amalgam (Fieer a. Perkin, *C J* 51, 836, *B* 19, 2568, Lupp, *B* 22, 2567) Liquid, miscible with water, decomposing at 236° HBrAq at 100° forms an anhydride (78° at 716 mm), and finally $\text{C}_8\text{H}_{14}\text{Br}_2$ (201°)

Di-oxy pentane $\text{CHEt}(\text{OH})\text{CHMe}(\text{OH})$ (187.5°) $\text{SG } 0.9945$, $\frac{1}{2}$ 9800 Formed from the corresponding di bromo pentane by successive treatment with AgOAc and baryta water (Wagner a Saytzeff, *A* 179, 308) Formed also by heating the oxide $\text{O} < \text{CHEt} < \text{CHMe}$ with water at 100° (Eltehoff, *J R* 1882, 355) Yields α oxy butyric acid on oxidation

Di oxy pentane $\text{CHPr}(\text{OH})\text{CH}_2(\text{OH})$ (206°). $\text{SG } 0.999$ Formed from the corresponding di-bromo pentane (Flavitzky, *B* 10, 230, 2240, *A* 179, 351) Yields oxy valeric acid on oxidation P_2O_5 forms valeric aldehyde and methyl isopropyl ketone

Di-oxy pentane $\text{CMe}_2(\text{OH})\text{CHMe}(\text{OH})$ (177°) $\text{SG } 0.987$ Made from the corresponding di bromo pentane (Wurtz, *A Ch* [3] 51, 458) Formed also by heating methyl isopropenyl carbinol with dilute H_2SO_4 (Kondakoff, *J R* 20, 32) Syrup, with bitter taste Miscible with water, alcohol, and ether Yields oxy-isobutyric acid on oxidation Yields methyl-isopropyl ketone when heated at 220°

Acetyl derivative $\text{C}_8\text{H}_{14}(\text{OAc})_2$ (above 200°) Oil, readily saponified

Benzoyl derivative $\text{C}_8\text{H}_{14}(\text{OBz})_2$ [123°]. Laminæ, sol alcohol and ether (Meyer, *C R* 59, 444)

Di oxy-pentane

$\text{CH}_3\text{CH}(\text{OH})\text{CH}_2\text{CH}(\text{OH})\text{CH}_3$ (177°). Formed by reducing methylene-di-methyl-diketone (acetyl acetone) with sodium amalgam (Combes, *A Ch.* [6] 12, 229). Liquid, sol water

Tri-oxy-pentane

$\text{CHMe}(\text{OH})\text{CH}_2\text{CH}(\text{OH})\text{CH}_2(\text{OH})$ (180° at 27 mm) $\text{SG } \frac{3}{4}$ 1185, $\frac{2}{3}$ 1120 Got by oxidising $\text{CH}_3\text{CH}(\text{OH})\text{CH}_2\text{CHMe}(\text{OH})$ with KMnO_4 . The yield is 82 p c (Wagner, *B* 21, 3351) Syrup not volatile with steam

Tri-acetyl derivative $\text{C}_6\text{H}_5(\text{OAc})_3$ (270°) at 740 mm $\text{SG } \frac{3}{4}$ 1120, $\frac{2}{3}$ 1101

Tri oxy-pentane

$\text{CHEt}(\text{OH})\text{CH}(\text{OH})\text{CH}_2(\text{OH})$ (192° at 68 mm) $\text{SG } \frac{3}{4}$ 10851 Made by oxidising $\text{CHEt}(\text{OH})\text{CHCH}_3$ (114°) with KMnO_4 (Wagner, *B* 21, 3349) The yield is 63 p c Thick, sweet syrup, miscible with water, alcohol, and ether

Tri-acetyl derivative (265°) $\text{SG } \frac{3}{4}$ 1122, $\frac{2}{3}$ 1103 Smells like onions

Tri oxy-pentane

$\text{CHMe}(\text{OH})\text{CMe}(\text{OH})\text{CH}_2(\text{OH})$ 'Pentenyl glycerol' (158° at 24 mm) Formed from tiglic alcohol by addition of bromine and subsequent distillation (Lieben & Zeisel, *M* 7, 68) Thick liquid

Tri acetyl derivative (149° at 18 mm)

Tetra-oxy pentane $\text{C}_6\text{H}_{12}\text{O}_4$ [253°] $\text{S } 6$ at 16° Formed by the action of lime on a solution of formic and acetic aldehydes (Tollens & Wigand, *A* 265, 316) Prisms Inactive to light HI and P form $\text{C}_6\text{H}_5\text{I}_2(\text{OH})_2$ [180°] and $\text{C}_6\text{H}_5\text{I}_2(\text{OH})$ [62°]

Tetra-acetyl derivative [84°]

Penta-oxy pentane

$\{\text{CH}_2(\text{OH})\text{CH}(\text{OH})\}_2\text{CH}(\text{OH})$ [102°] Formed by reducing arabinose with sodium amalgam, the liquid being kept neutral with H_2SO_4 (Kiliani, *B* 20, 1233) Prisms or needles, v e sol water, v sl sol cold alcohol

OXY-PENTANE TRICARBOXYLIC ACID

$\text{C}_6\text{H}_4(\text{OH})\text{C}(\text{CO}_2\text{H})_2\text{CH}_2\text{CO}_2\text{H}$ Formed from $\text{C}_6\text{H}_5\text{C}(\text{CO}_2\text{H})_2\text{CH}_2\text{CO}_2\text{H}$ by successive treatment with HBr and KOH (Hjelt, *B* 16, 1258)

Anhydride $\text{C}_6\text{H}_4\text{O}_5$ Dicarboxyprolactonic acid [163°] Triclinic crystals (from water).— $\text{BaC}_6\text{H}_4\text{O}_5$.— $\text{Ag}_2\text{C}_6\text{H}_4\text{O}_5$ pulverulent pp

OXY-PENTANE PHOSPHONIC ACID

$\text{C}_6\text{H}_4(\text{OH})_2\text{PO}(\text{OH})_2$ [184°] Formed from isovaleric aldehyde and PCl_5 followed by water (Fossek, *M* 5, 627, 7, 20) Monoclinic scales Yields isovaleric aldehyde on distillation and on treatment with KMnO_4 , PCl_5 forms $\text{C}_6\text{H}_4\text{Cl}_2\text{POCl}_2$ (184°–140° at 22 mm) Reduced by HI and P to isopentane phosphonic acid $\text{C}_6\text{H}_{11}\text{PO}_3$ [161°]— $\text{BaA}''2\text{aq}$ — $\text{Ba}(\text{HA}'')$ — $\text{Ag}_2\text{A}''$ amorphous pp

OXY-PENTANE SULPHONIC ACID

$\text{C}_6\text{H}_4(\text{OH})_2\text{SO}_3\text{H}$ Amylene isethionic acid Formed from chloro amyl alcohol (amylenic chlorhydrin) and Na_2SO_3 (Falk, *J pr* [2] 2, 272) Decomposed on evaporation of its solution— NaA' — CaA'_2 — $\text{CuA}'_22\text{aq}$ pale blue laminae

An isomeric acid is formed by the action of SO_3 on isomyl alcohol (Schwarz, *B* 3, 691)— BaA'_2 crystalline

OXYPENTIC ACID $\text{C}_6\text{H}_4\text{O}_5$ 3aq or $\text{C}_6\text{H}_4\text{O}_5$ [198°] Formed from ethyl acetoacetic ether, bromine, and alcoholic potash (Demarcay, *C R* 88, 289) Crystals, m sol cold water Ammonia converts its ether into a substance [77°],

and its chloride into another body [204°] Zn and H_2SO_4 yield $\text{C}_6\text{H}_4\text{O}_5$ [95°] Oxyptic acid is perhaps $\Delta\gamma$ hydromuconic acid (Gorboff, *J R* 1887, 605, this vol p 443) *

OXYPENTINOIC ACID $\text{C}_6\text{H}_4\text{O}_5$ v e

$\text{CH}_3\text{COCH}(\text{OH})\text{CH}_2\text{CO}_2\text{H}$ Aceto-acrylic acid. [125°] Prepared by boiling an aqueous solution sodium β bromo-lavulate for a few minutes, neutralising with Na_2CO_3 , again boiling, neutralising again, and so on till the solution finally remains alkaline after boiling (Wolff, *B* 20, 426) Glistening plates V sol alcohol and ether, less sol cold water and chloroform

Salts— CaA'_2 — ZnA'_2 amorphous— AgA' needles

OXY-PHENANTHRAQUINONE v PHENANTHRAQUINONE

DI OXY-PHENAZINE $\text{C}_{12}\text{H}_8\text{N}_2\text{O}_2$ Formed by heating di-amido phenazine with conc HClAq at 200° (Fischer & Hepp, *B* 23, 843, Nietzki & Hasterlik, 24, 1337) Reddish yellow needles (containing 3aq)— $\text{B}'\text{H}_2\text{SO}_42\text{aq}$

Di-acetyl derivative [230°] (F & H), [226°] (N & H) Pale yellow tables

Dioxytriphenazine

$\text{C}_6\text{H}_4\left\langle\begin{array}{c} \text{NH} \text{ C } \text{C } \text{C } \text{N} \\ | \quad | \quad | \\ \text{N} \quad \text{C } \text{C } \text{C } \text{N} \end{array}\right\rangle\text{C}_6\text{H}_4$ Quinone homo-

fluorindin Formed by heating di o nitro diphenyl quinone with alcoholic ammonium sulphide at 100°, the product being oxidised by exposure to air (Leicester, *B* 23, 2794) Dark green crystals, yielding a violet powder Its solution in HOAc exhibits brownish red fluorescence

OXYDIPHENYLS All these compounds yield diphenyl when distilled with zinc dust

p Oxy diphenyl $\text{C}_6\text{H}_4\text{C}_6\text{H}_4\text{OH}$ [165°] (c 307°) Formed by fusing diphenyl sulphonic acid with potash (Latschinoff, *B* 6, 194, Osten, *B* 7, 170), by warming diazobenzene chloride with phenol, and in small quantity in the decomposition of diazobenzene chloride by water (Hirsch, *B* 23, 3705) Plates, volatile with steam Yields a mono nitro derivative [67°] and a di nitro derivative [154°]

Acetyl derivative [89°] Plates, v sol alcohol and ether (Kaiser, *A* 257, 101)

Benzoyl derivative [152°] Tables

An isomeric or identical oxydiphenyl [152°] is obtained from p amido diphenyl by the diazo reaction (Hubner, *A* 209, 348)

oo Di-oxy-diphenyl $\text{C}_6\text{H}_4(\text{OH})\text{C}_6\text{H}_4(\text{OH})$ [99°] Formed by potash fusion from diphenyl disulphonic acid (Lumprecht, *Z* 1871, 261, and by dropping fluorene into fused KOH at 400° (Hodgkinson & Matthews, *C J* 43, 168) Crystals (from benzene ligroin)

pp Di-oxy-diphenyl [272°] (above 360°) Formed from benzidine by the diazo reaction (Griess, *J* 1866, 461, Hirsch, *B* 22, 336), and by potash fusion from diphenyl di-p sulphonic acid (Engelhardt & Latschinoff, *Z* 1871, 261, Doebner, *B* 9, 180) Got also by distilling di-oxy-phenyl benzoic acid with lime (Schmidt & Schultz, *A* 207, 334, *B* 12, 490) Obtained also by passing a rapidly alternating electric current through a solution of phenol (Drechsel, *J. pr* [2] 29, 237) Laminae (from alcohol)

Di-acetyl derivative [160°]

Ethyl ether [176°] (Hirsch)

op-Di oxy-diphenyl [161°] (342°) Formed in small quantity by potash fusion from phenol *o*- and *p*-sulphonic acids (Lincke, *J pr* [2] 8, 43, Herzog, *B* 1st, 2234), and by the action of nitrous acid on *op* di amido diphenyl (Schmidt, *A* 207, 357, *B* 12, 497) Yields a di acetyl derivative [94°]

By fusing phenol with potash two more di oxy diphenyls [123°] and [190°] have been obtained (Barth a Schroeder, *A* 156, 93, *B* 11, 1336) The one melting at 123° yields $C_{12}H_8(O_2Me)_2$ (310°-320°) and $C_{12}H_8O_2(SO_3H)_2$, whence $Na.A''$ 2aq and KA'' aq

Tri oxy-diphenyl $C_6H_4(OH)_2C_6H_4(OH)_2$ Two isomerides of this formula [180°] and [205°] are formed by fusing fluorene sulphonic acid with potash at a little above 400° (Hodgkinson a Matthews, *C J* 43, 167) Neither gives any colour with $FeCl_3$ The compound [180°] forms an acetyl derivative [100°]

Tetra oxy diphenyl $C_6H_4(OH)_2C_6H_4(OH)_2$ *Dipyrocatechin* [84°] Formed from diphenyl (a) disulphonic acid by potash fusion (Barth, *B* 11, 1336) Needles $FeCl_3$ colours its solution green

Tetra oxy diphenyl $C_{12}H_8O_4$ *Diresorcin* [310°] Formed in small quantity by fusing resorcin or phenol with NaOH (Barth, *B* 12, 503, Benedikt, *M* 1, 355, 5, 177, Herzog, *M* 11, 419) Needles (containing 2aq) Gives a blue colour with $FeCl_3$ Phthalic anhydride and H_2SO_4 at 120° form $C_{12}H_{10}O_5$ 4aq, which gives blue solutions with alkalis

Acetyl derivative $C_{12}H_8(OAc)_2$ [159°]

Benzoyl derivative $C_{12}H_8(OBz)_2$ [199°] Formed from phloroglucin, $BzCl$, and NaOH (Skraup, *M* 10, 721, Pukall, *B* 20, 1143)

Ethyl ether $C_6H_4(OEt)_2$ [114°] Plates

Tetra oxy-diphenyl $C_6H_4(OH)_2C_6H_4(OH)_2$ *Dihydroquinone* [237°] Formed by fusing hydroquinone with NaOH (Barth, *M* 5, 600) Plates Tastes sweet $FeCl_3$ colours it red, and then forms diquinhydrone $C_{12}H_8O_4$ and diquinone $C_{12}H_8O_2$ [187°]

Hexa-oxy diphenyl $C_6H_4(OH)_2C_6H_4(OH)_2$ Formed by heating hydrocærulignone with $HClAq$ at 200° (Liebemann, *A* 169, 239, *B* 9, 1887) Plates, m sol water $FeCl_3$ gives a bluish grey pp Iodine yields $C_{12}H_8O_6$, crystallising in blue needles

Tri methyl ether $C_{12}H_8(OMe)_3(OH)_2$ Formed from cærulignone and H_2SO_4 (Fischer, *B* 8, 158)

Tetra methyl ether $C_{12}H_8(OMe)_4(OH)_2$ *Hydrocærulignone* [190°] Formed by reduction of cærulignone (Liebemann, *A* 169, 226) Monoclinic prisms (from alcohol) $FeCl_3$ oxidises it to cærulignone Yields $C_{12}H_8(OMe)_4(OAc)_2$ [217°-225°] — $Na_2C_4H_8O_6$ — K_2A'' 4aq (Ewald, *B* 11, 1623)

Hexa-methyl ether $C_{12}H_8(OMe)_6$ [126°]

Tetra-ethyl ether $C_{12}H_8(OH)_2(OEt)_2$ [176°] Crystals (from alcohol) (Hofmann, *B* 11, 802)

Hexa-acetyl derivative $C_{12}H_8(OAc)_6$ [145°]

Hexa-propionyl derivative Needles

Di-bromo hexa-oxy-diphenyl Tetra-methyl derivative $C_{12}H_8Br_2(OH)_2(OMe)_2$ [262°] Formed from $C_{12}H_8Br_2(OAc)_2(OMe)_2$ [178°], which is got from di acetyl cærulignone,

$HOAc$, and bromine (Hayduck, *B* 9, 929) It yields $C_{12}H_8Br_2(OMe)_6$ [140°]

References — Di bromo and Di chloro-~~hexa~~ OXY DIPHENYL

(8) **Hexa oxy diphenyl** Formed by fusing ellagic acid with potash (Barth a Goldschmidt, *B* 12, 1244) Needles, blackening at 250°, and melting at a higher temperature Its alkaline solution is blue, changing in air to red

(7) **Hexa-oxy-diphenyl** Formed from ellagic acid by fusion with NaOH (*B* a *G*), or by treatment with sodium amalgam (Gobenzl, *M* 1, 672) Needles, blackening at 230° Its alkaline solution is red, changing in air to green

(8) **Hexa oxy diphenyl** Formed by fusing hydroquinone with NaOH (Barth a Schroeder, *M* 5, 597) Plates, blackening and melting near 290° Its alkaline solution is red

Acetyl derivative [172°] Needles

OXY PHENYL ACETAMIDINE $C_6H_5N_2O$ 1e $CHPh(OH)C(NH)(NH_2)$ [110°] Formed from mandelic imido ether and alcoholic NH_3 (Beyer, *J pr* [2] 28, 191) Needles (from ether) — $B'HCl$ [214°] Prisms (from water)

Di acetyl derivative

$CHPh(OAc)C(NH)NHAc$ [210°] (Pinner, *B* 23, 2948)

OXY-PHENYL ACETAMIDOXIM

$C_6H_5CH(OH)C(NH_2)(NOH)$ [159°] Formed from the nitrile of mandelic acid and alcoholic hydroxylamine (Tiemann, *B* 17, 126, Gross, *B* 18, 1074) Crystals (from alcohol) Yields the derivatives $C_6H_5NaNO_2$, $B'HCl$, $C_6H_5EtN_2O_2$ [89°], and $C_6H_5(CH_2Ph)N_2O_2$ [103°]

Reactions — 1 $COCl_2$ forms the compound $CHPh(OH)C(NH_2)NO_2CO$ [121°] — 2 Its hydrochloride reacts with *potassium cyanate*, forming $CHPh(OH)C(NOH)NHCONH_2$ [127°] — 3 *Phenyl cyanate* forms the compound $CHPh(OH)C(NOH)NHCONHPh$ [155°] — 4 $ClCO_2Et$ yields $CHPh(OH)C(NH_2)NOCO_2Et$ [107°]

Acetyl derivative

$CHPh(OH)C(NH_2)NOAc$ [140°] Crystals (from alcohol)

Di acetyl derivative [113°] Plates

Benzoyl derivative $C_6H_5N_2O_2$ [149°].

Acetyl benzoyl derivative [165°]

OXY PHENYL-ACETIC ACID

[2 1] $C_6H_4(OH)CHCO_2H$ [137°] Formed from *o* oxy phenyl-glyoxylic acid by reduction with sodium amalgam to $C_6H_4(OH)CH(OH)CO_2H$, and further reduction of this acid with HI (Baeyer a Fritsch, *B* 17, 975) Needles, v sol water Gives a violet colour with $FeCl_3$ On heating it yields the lactone

$C_6H_4 < \begin{smallmatrix} CH_2 \\ O \end{smallmatrix} > CO$, crystallising in tables [49°] (237°)

m **Oxy phenyl-acetic acid** [129°] Formed by saponification of its nitrile and from *m*-amido-phenyl acetic acid by the diazo reaction (H Salkowski, *B* 17, 507) Needles, v sol water Coloured violet by $FeCl_3$

Nitrile [3 1] $C_6H_4(OH)CH_2CN$ [53°] Formed from $C_6H_4(NH)CH_2CN$ and nitrous acid Trimetric tables, v sol alcohol and hot water

p-**Oxy phenyl acetic acid** [148°] Formed by the action of nitrous acid on *p* amido phenyl-acetic acid (Salkowski, *B* 12, 1438), and by

saponifying its nitrile, which is a product of the action of AgNO_3 on sinalbin (Will a Laubenheimer, *A* 199, 156, Salkowski, *B* 22, 2137) Occurs in human urine (Baumann, *B* 13, 280) Prismatic needles, sol water and alcohol FeCl_3 gives a dark colouration $-\text{NH}_4\text{A}' - \text{CaA}'_2, 4\text{aq} - \text{PbA}'_2, -\text{AgA}'$ minute needles

Methyl ether $\text{C}_6\text{H}_5(\text{OH})\text{CH}_2\text{CO Me}$ (310° i v) $\text{SG } \frac{1}{2}$ 11948, $\frac{3}{4}$ 11786 Oil
Ethyl ether EtA' (314° i v) $\text{SG } \frac{1}{2}$ 11886, $\frac{3}{4}$ 11226

Nitrile [70°] (330.5° i v) Made like stem isomeride (Salkowski, *B* 22, 2137) Trichinic crystals Yields $\text{C}_6\text{H}_5(\text{OMe})\text{CH}_2\text{CN}$ (287° i v) $\text{SG } \frac{1}{2}$ 11001

Methyl derivative $\text{C}_6\text{H}_4(\text{OMe})\text{CH}_2\text{CO}_2\text{H}$ [86°] Formed from $\text{C}_6\text{H}_5(\text{OMe})\text{CH}_2\text{Cl}$ and KCy , the product being saponified (Cannizzaro, *A* 117, 243) Pearly plates $-\text{AgA}'$

Ethyl derivative [88°] Plates
Amide $\text{C}_6\text{H}_5(\text{OH})\text{CH}_2\text{CONH}_2$ [175°]

Methyl derivative of the amide
 $\text{C}_6\text{H}_5(\text{OMe})\text{CH}_2\text{CONH}_2$ [189°] Scales

o-Oxy phenyl acetic acid *v* MANDELIC ACID

oo-Di oxy-phenyl acetic acid

$\text{C}_6\text{H}_4(\text{OH})\text{CH}(\text{OH})\text{CO}_2\text{H}$ Formed from salicylic aldehyde, HCy , and HCl (Plöschl, *B* 14, 1317), and also by reducing oxy phenyl glyoxylic acid (Bayer a Fritsch, *B* 17, 974) Syrup Yields a crystalline anhydride

Methyl derivative of the nitrile

$\text{C}_6\text{H}_5(\text{OMe})\text{CH}(\text{OH})\text{CN}$ [71°] Formed from methyl salicylic aldehyde and HCy (Voswinckel, *B* 15, 2025) Colourless crystals

Di-oxy phenyl-acetic acid

[1 3 5] $\text{C}_6\text{H}_3(\text{OH})_3\text{CH}_2\text{CO}_2\text{H}$ [54°] Formed from $\text{C}_6\text{H}_3(\text{OH})_3(\text{CO}_2\text{Et})_2\text{CH}_2\text{CO}_2\text{Et}$ and alcoholic potash (Pechmann, *B* 19, 1449) Crystals (containing aq) $-\text{PbA}'_2, 2\text{aq}$ needles (from water)

op-Di oxy phenyl-acetic acid *Methyl derivative*

$\text{C}_6\text{H}_4(\text{OMe})\text{CH}_2\text{CH}(\text{OH})\text{CO}_2\text{H}$ [93°] Formed from anisic aldehyde, HCy , and HCl (Tiemann a Kohler, *B* 14, 1976) Needles $-\text{CaA}'_2, -\text{AgA}'$ amorphous pp

Amide $\text{C}_6\text{H}_4(\text{OMe})\text{CH}_2\text{CH}(\text{OH})\text{CONH}_2$ [159°]

Nitrile $\text{C}_6\text{H}_4(\text{OMe})\text{CH}_2\text{CH}(\text{OH})\text{CN}$ [63°]

Di oxy-phenyl acetic acid

$\text{C}_6\text{H}_3(\text{OH})_3\text{CH}_2\text{CO}_2\text{H}$ [4 3 1] *Homoprotocatechuic acid* [127°] Formed by heating its methyl derivative with HClAq at 175° (Tiemann a Nagai, *B* 10, 207) Slender needles (from benzene) FeCl_3 gives a green colour

Di acetyl derivative

$\text{C}_6\text{H}_3(\text{OAc})_3\text{CH}_2\text{CO}_2\text{H}$ [90°] (Nagai, *B* 11, 658)

Methyl derivative

$\text{C}_6\text{H}_3(\text{OMe})_3\text{CH}_2\text{CO}_2\text{H}$ [3 4 1] *Homovanillic acid* [143°] Made from its acetyl derivative [140°], which is got by oxidising acetyl eugenol

Di-methyl derivative

$\text{C}_6\text{H}_3(\text{OMe})_3\text{CH}_2\text{CO}_2\text{H}$ *Homoveratric acid* [99°] (Tiemann a Matsumoto, *B* 11, 143) Needles (containing aq)

Methylene derivative

$\text{C}_6\text{H}_4(\text{OCH}_3)_2\text{CH}_2\text{CO}_2\text{H}$ [128°] Got from saffrol, KMnO_4 , and dilute HOAc (Tiemann, *B* 24, 2882) Yields MeA' (279°), EtA' (291°), and an amide [173°]

A di-oxy phenyl-acetic acid [168°] occurs in urine after poisoning by phosphorus (Baumann, *H* 6, 192) It crystallises in small needles

Tri-oxy phenyl-acetic acid. Methylene derivative [4 3 1] $\text{C}_6\text{H}_3(\text{O}_2\text{CH}_3)_3\text{CH}(\text{OH})\text{CO}_2\text{H}$ [153°] Made from piperonal by treatment with HCy and saponification (Lorenz, *B* 14, 793)

Hexa-oxy-di-phenyl acetic acid Hexa methyl derivative

$(\text{C}_6\text{H}_5(\text{OMe}))_2\text{CH}(\text{OH})\text{CO}_2\text{H}$ [175°] Formed by boiling the hexa methyl derivative of hexa oxybenzil with potash (Marx, *A* 263, 255) Prisms, *v* sol alcohol, *sl* sol water

o-Oxy-di-phenyl-acetic acid *v* BENZILIC ACID

Tri-oxy-di-phenyl-acetic acid *Di methyl derivative v* ANISILIC ACID

(*B* 2) *OXY (A) PHENYL-ACRIDINE*

$\text{C}_6\text{H}_5\text{C}(\text{N})\text{CPh} > \text{C}_6\text{H}_5(\text{OH})$ [c $275^\circ - 280^\circ$] Formed

by heating *p* oxy di-phenylamine with benzoic acid and ZnCl_2 Formed also by heating (*B* 2) amido (*A*) phenyl acridine with HCl at $200^\circ - 220^\circ$ (Hess a Bernthsen, *B* 18, 695) Thin yellow crystals *v* sol alcohol and acetic acid, less sol ether Dissolves in caustic alkalis and in acids The ethereal and alcoholic solutions have a slight bluish-violet fluorescence The salts are all rather sparingly soluble, and have a tendency to separate in a gelatinous form

Acetyl derivative $\text{C}_{10}\text{H}_{13}\text{N}(\text{OAc})$ [174°], glistening four sided prisms

OXY - PHENYL - ACRYLIC ACID v OXY CINNAMIC ACID and COUMARIC ACID

Oxy-di-phenyl-acrylic acid Methyl derivative $\text{C}_6\text{H}_4(\text{OMe})\text{CH CPh CO}_2\text{H}$ [189°] Made from anisic aldehyde, sodium phenyl acetate, and Ac_2O at 160° (Oghaloro, *G* 9, 533) Prisms, yielding $\text{C}_6\text{H}_4(\text{OMe})\text{CH CPh}$ when heated

Methyl derivative of the nitrile $\text{C}_6\text{H}_4(\text{OMe})\text{CH CPh CN}$ [93°] Formed from anisic aldehyde, benzyl cyanide, and alcoholic NaOEt (Frost, *A* 250, 159) White needles

Reference —PHENYL COUMARIN

TRI OXY-PHENYL ALLYL KETONE According to Schiff (*A* 253, 336), flicic acid (vol u p 548) is the isobutryl derivative of this ketone [5 3 1 2] $\text{C}_6\text{H}_2(\text{OH})_3(\text{O COEt})\text{CO CO}_2\text{H}$

OXY TRI PHENYL-ALLYL-PYRROLE

$\text{N}(\text{C}_6\text{H}_5)_3\text{C}(\text{CPh})\text{CH} > \text{CO CPh}_2$ [112°] Formed by heating anhydriacetophenone - benzil PhCBz CHBz with alcoholic allylamine at 100° (Japp a Klingemann, *C* 57, 707) Monoclinic prisms (from alcohol), a b c = 1 665 1 844, β = $88^\circ 53'$

o *OXY PHENYL-ALLYL THIO UREA*

$\text{C}_6\text{H}_4(\text{OH})\text{NH CS NHC}_6\text{H}_5$ [99°] Formed from *o* amido phenol and allyl thiocarbimide in alcoholic solution (Von Chelmicki, *J pr* [2] 42, 442) *v* *sl* sol cold, *v* sol hot, water

o *OXY PHENYL AMIDO ACETIC ACID*

$\text{C}_6\text{H}_4(\text{OH})\text{CH}_2\text{CO}_2\text{H}$ *Oxy - phenyl - glycolic acid* Formed by boiling chloro acetic acid (1 mol) with *o* amido phenol (2 mols) and water (Vater, *J pr* [2] 29, 289) Plates (containing aq), *sl* sol water, *v* sol alcohol FeCl_3 colours its solutions red At 105° it yields an anhydride which crystallises from alcohol in cubes

Methyl derivative

$\text{C}_6\text{H}_4(\text{OMe})\text{NH CH}_2\text{CO}_2\text{H}$ [142°] Formed from *o* anisidine and chloro acetic acid Rectangular plates $-\text{PbH}_4\text{A}'$, $-\text{B'HCl}$ crystals

Ethyl derivative $\text{C}_6\text{H}_4(\text{OEt})\text{NH CH}_2\text{CO}_2\text{H}$ [120°] Formed from chloro acetic acid and

$C_6H_5(OEt)NH_2$ — EtA' long waxy needles— $EtA'HCl$

***p*-Oxy-phenyl amido-acetic acid**

$C_6H_5(OH)NHCH_2CO_2H$ Formed from chloro-acetic acid and *p* amido phenol (V) Plates, sl sol water and alcohol $FeCl_3$ colours its solutions red— NaA'

Methyl derivative

$C_6H_5(OMe)NHCH_2CO_2H$ Formed in like manner from *p* anisidine (V), and also by saponifying its nitrile, which is got by heating anisaldehyde cyanhydrin with alcoholic NH_3 (Tiemann & Kohler, B 14, 1976)—Needles, sol hot water— CuA' — $HA'HCl$

Ethyl derivative

$C_6H_5(OEt)NHCH_2CO_2H$ [163°] Crystals (containing $\frac{1}{2}$ aq) Yields, on heating, $C_6H_5(OEt)NHCH_2CO_2NHC_6H_5(OEt)$ [140°] and other bodies (Bischoff & a Navstogel, B 22, 1788)

***p*-OXY-PHENYL-AMIDO-BENZENE SULPHONIC ACID** $C_6H_5(OH)(NPh)SO_3H$ Formed from $C_6H_5(OH)NPh$ [1 4] and conc H_2SO_4 at 100° (Limpicht, B 22, 2910) Small grey prisms Does not melt below 260°

OXY-PHENYL AMIDO CROTONIC ETHER

Methyl derivative $C_{12}H_{13}NO_2$ [46°] Formed from *p* anisidine and acetoacetic ether (Conrad & Limpach, B 21, 1649)

DI OXY-PHENYLAMIDOETHYL BENZOIC ACID

Methylene derivative of the anhydride $C_{16}H_{13}NO_4$ *re*

$CH_2 < \begin{smallmatrix} O \\ \diagup \quad \diagdown \end{smallmatrix} C_6H_5 < \begin{smallmatrix} CO \\ \diagup \quad \diagdown \end{smallmatrix} NPh$ [157°] Formed from CH_2O , $C_6H_5(C_6H_5Cl)CO_2Me$ and alcoholic aniline (Perkin, jun, C J 57, 1036) Prisms, v sl sol hot alcohol

OXY PHENYL AMIDO NAPHTHOQUINONE

$C_{16}H_9O_5(OH)(NPh)$ ***Oxy naphthoquinone anilide*** [210°] Formed from $C_{10}H_7ClO_2(NPh)NO$ and boiling $KOHAq$ (Plagemann, B 16, 896) Deep blue needles with metallic lustre, sol alkalis

***o*-OXY-PHENYLAMIDO PHENYL-ACETO**

NITRILE ***Methyl derivative*** $C_6H_5(OMe)CH(NPh)CN$ [61°] Formed by the action of aniline on $C_6H_5(OMe)CH(OH)CN$ (Voswinkel, B 15, 2027) Six sided tables, sol alcohol

OXY PHENYL AMIDO QUINONE ANILIDE

$C_6H_5(OH)(NPh)O(NPh)$ Formed by heating its ethers with alcoholic potash (Zincke & Hagen, B 18, 788) Small scales, sl sol alcohol Cold, very dilute potash, converts it into the quinone $C_6H_5(OH)(NPh)O_2$, a blue crystalline pp decomposing above 200°

Methyl ether $C_6H_5(OMe)(NPh) < \begin{smallmatrix} O \\ \diagup \quad \diagdown \end{smallmatrix} NPh$

[191°] Formed by warming the compound $C_6H_5(NPh)O(NPh)$ with $MeOH$ and H_2SO_4 (Z & H), and also from azophenine, $MeOH$, and H_2SO_4 (Fischer & Hepp, B 21, 677, A 262, 253) Brownish red plates, forming blue salts with acids, $C_6H_5(OMe)(NPh)O_2$ [189°]

Ethyl ether $C_6H_5(OEt)(NPh)O(NPh)$

[137°] Formed in like manner, using $EtOH$

Isobutyl ether [138°] Needles

OXY-PHENYL-AMIDO-THYMOQUINONE

$C_6Me(C_6H_5)(OH)(NPh)O_2$ [135°] Formed by boiling oxy thymoquinone with aniline and alcohol (Schulz, B 16, 902). Lustrous, dark-

violet needles (from alcohol), forming a violet solution in NH_3Aq

OXY-PHENYL AMIDO-TOLUQUINONE

$C_6HMeO_2(OH)(NPh)$ Formed from di phenyl di amido toluquinone by boiling with alcoholic H_2SO_4 (Hagen & Zincke, B 16, 1560) Deep blue needles (from alcohol), decomposing at 250°

Anilide $C_6HMe(OH)(NPh) < \begin{smallmatrix} O \\ \diagup \quad \diagdown \end{smallmatrix} NPh$

Formed by treating its ethers with acids or alkalis Brown needles (from $HOAc$) The ethers $C_{11}H_{11}MeN_2O_2$ [131°], $C_{11}H_{11}EtN_2O_2$ [116°] and $C_{11}H_{11}(C_6H_5)N_2O_2$ [117°] are crystalline

***o*-OXY-PHENYL-*o*-AMIDO-TOLYL GLYCOLIC ACID** $NPhCH(OH)C_6H_5OCH_2CO_2H$

Formed from *o* aldehyde phenoxy acetic acid and aniline (Rossing B 17, 2992) Very unstable in alkaline solution— $HA'HCl$ [191°]— $HA'H_2SO_4$ [186°] Yellow needles, v sol hot water

***m* OXY DIPHENYLAMINE**

$C_6H_5NH C_6H_5(OH)$ ***Phenyl amido phenol*** [82°] (c 340°) Formed by heating resorcin (1 mol) with aniline (4 mols) and $CaCl_2$ (2 mols) for eight hours at 280°, the yield being 85 p c of the theoretical (Calm, B 10, 2786, Kohler, B 21, 908) White pearly plates, v sol alcohol Yields a crystalline nitrosamine $NPh(NO)C_6H_5OH$ [115°]— $HA'HCl$ needles— $H_2A_2H_2SO_4$ needles— BaA' 5aq colourless plates, sol water

***p* Oxy diphenylamine** $C_6H_5NH C_6H_5OH$

[70°] (330°) Prepared by heating hydroquinone (1 mol) with aniline (4 mols) and $CaCl_2$ (2 mols) for ten hours at 260°, the yield being 90 p c (Calm, B 16, 2799, 17, 2431) Formed also by heating hydroquinone with aniline for sixteen hours at 300° (the yield being 85 p c), and by the action of aniline on quinone phenylimide (Bandrowski, M 9, 133, 416) Plates, v sol aqueous alkalis and acids, sl sol water Yields a nitrosamine [95°] On oxidation with HgO in benzene it yields $C_{12}H_9NO$ [97°] (B)— $HA'HCl$ slender needles, decomposed by water— $HA'HB$ needles

Isobutyl ether C_6H_5A' [68°]

Formyl derivative $C_{12}H_9NO(CHO)$ [178°] White needles, sol alcohol

Di acetyl derivative $NPhAc C_6H_5OAc$

[120°] Monoclinic prisms

Di-benzoyl derivative [175°] Prisms

Di oxy-diphenylamine $NH(C_6H_5O)_2$

Formed by heating resorcin with ammoniacal $CaCl_2$ at 200° (Seyevitch, C R 109, 946). Minute crystals (from dilute alcohol)

***o* OXY PHENYL-ANGELIC ACID**

$C_{11}H_{11}O_2$ *re* $C_6H_5(OH)CHCMeCH_2CO_2H$ [73°] Formed from salicylic aldehyde, sodium pyrotartrate, and Ac_2O (Fittig & Brown, A 265, 290) Large plates, v sol chloroform— BaA' 4aq— AgA' flocculent pp

***o*-Oxy-phenyl-angelic acid**

$C_6H_5(OH)CHCETCO_2H$ [174°] Formed by boiling its anhydride with potash (Perkin, A 150, 84, C J 21, 472) Flat prisms (from dilute alcohol), sl sol water and chloroform.— AgA' yellowish pp

Anhydride $C_6H_5 < \begin{smallmatrix} CHCET \\ \diagup \quad \diagdown \end{smallmatrix} O < \begin{smallmatrix} CHCET \\ \diagup \quad \diagdown \end{smallmatrix} CO_2H$ [71°] (299°).

Formed by boiling sodium salicylic aldehyde

with butyric anhydride (Perkin, *A* 147, 233, *C J* 21, 53, 472, Fletcher, *C J* 39, 447) Formed also by distilling coumaryl propionic acid (Fittig, *A* 255, 288) Monoclinic crystals, $a b c = 1.192 \ 1.694, \beta = 67^\circ 18'$

(a) *Methyl derivative*

$C_6H_5(OMe)CHCHEtCO_2H$ [88°] Formed from the anhydride $NaOH$, and MeI , the resulting $C_6H_5(OMe)CHCHEtCO_2Me$ (282°) being saponified (Perkin, *C J* 39, 435). Monoclinic crystals

(b) *Methyl derivative*

$C_6H_5(OMe)CHCEtCO_2H$ [105°] Formed from methyl-salicylic aldehyde, butyric anhydride, and sodium butyrate. Needles (from alcohol) Yields $C_6H_5Br_2(OMe)C_6H_5Br_2CO_2H$ [159°] and $C_6H_5(OMe)CHCEtCO_2Me$ (292°) HI forms a compound whence Na_2CO_3 produces *o* butenyl-phenol

p-Oxy-phenyl-angelic acid

$C_6H_4(OH)CHCEtCO_2H$

Methyl derivative [124°] Formed from anisic aldehyde, butyric anhydride, and sodium butyrate (Perkin) Needles (from alcohol)

γ Oxy-phenyl-angelic acid *Anhydride*

$C_6H_5CH<\begin{smallmatrix} CH \\ CO-O \end{smallmatrix} CMe$ [53°] Made by distilling a phenyl levulic acid (Erdmann, *A* 254, 219) Plates (from CS_2 and ligroin)

Di oxy-phenyl-angelic acid *Methylene derivative*

$CH_2<\begin{smallmatrix} O \\ \diagup \diagdown \end{smallmatrix} C_6H_5CH_2CHCHCH_2CO_2H$ *Hydropiperic acid* [78°] Formed by reducing piperic acid with sodium amalgam (Foster, *A* 124, 117, *Burn*, *A* 216, 172, *Weinstein*, *A* 227, 32) Needles (from hot water) Changes on heating with $NaOH$ aq into a less soluble isomeric acid [181°] Both acids combine with bromine — $NH_4A' - KHA' - BaA' - AgA'$ crystalline pp

Di-oxy-phenyl angelic acid *Methylene*

derivative $CH_2<\begin{smallmatrix} O \\ \diagup \diagdown \end{smallmatrix} C_6H_5C_6H_5CO_2H$ [c 160°] Prepared by heating piperonal with butyric anhydride and $NaOAc$ (Lorenz, *B* 14, 786) Long felted needles (from dilute alcohol), sl sol water — AgA' curdy pp

Di-*p*-oxy di-phenyl angelic acid *Di-methyl derivative*

$C_6H_4(OMe)CHCHCH(CO_2H)CH_2C_6H_4(OMe)$ [111°] Made by reducing $C_6H_4(OMe)OH$, $C_6H_5CO_2H$ with sodium amalgam (Fittig a Politis, *A* 255, 302) Hard needles (from CS_2), sl sol water, v sol alcohol Yields a dibromide $C_{18}H_{15}Br_2O_4 - CaA'2aq - AgA'$ flocculent pp

DI-OXY-DI-PHENYL-ANTETRAZINE DI-HYDRIDE $N=C(OH)CCH_2C<\begin{smallmatrix} N \\ CPh \end{smallmatrix} CPh$

Formed from dioxyphterphthalic ether, benzamide hydrochloride, and dilute (10 p c) $NaOH$ aq (Pinner, *B* 22, 2624) Crystalline powder, insol water and alcohol — $C_{22}H_{15}N_4Na_2O_4aq$ white needles

OXY-PHENYL-ANTHRANOL $C_{10}H_7O_2$ *is*

$C_6H_4<\begin{smallmatrix} C(OH) \\ C(C_6H_5OH) \end{smallmatrix} C_6H_5$ Formed by the action of H_2SO_4 on oxy-triphenyl-methane carboxylic acid (Fechmann, *B* 13, 1616) Exhibits green fluorescence in ethereal solution.

Di-oxy-phenyl-anthranol $C_{12}H_9O_3$ *is*

$C_6H_5<\begin{smallmatrix} C(C_6H_5OH) \\ C(OH) \end{smallmatrix} C_6H_5$ *Phenolphthalidin* A sticky mass got by dissolving phenol phthalin in H_2SO_4 , and pouring into water (Baeyer, *A* 202, 91) Its ethereal solution fluoresces green Water at 170° converts it into phenol phthalin Potash-fusion gives di oxy benzophenone

OXY-PHENYL-ARSINE v *ARSINE*

DI-OXY-PHENYL-TRIAZOLE $C_8H_5N_3O_2$ *is*

$C_6H_5N-C(OH)<\begin{smallmatrix} N \\ N=C(OH) \end{smallmatrix} N$ or $C_6H_5NCO<\begin{smallmatrix} N \\ HNCO \end{smallmatrix} N$ *Phenylurazol* [263°]

Formation — 1 By heating phenyl hydrazine hydrochloride (1 pt) with urea (2 pts) for 4 or 5 hours at 160°, the yield is 70–80 p c of the hydrazine hydrochloride used — 2 By heating phenyl semicarbazide (1 pt) with urea (2 pts) for 4 hours at about 160°

Properties — Colourless glistening plates M sol hot water, sl sol cold water, v sol hot alcohol, sl sol ether Dissolves in cold aqueous alkalis and ammonia, and is reppd by acids unaltered It does not reduce alkaline solutions of Ag or Cu It is not affected by boiling with alkalis or acids (Pinner, *B* 20, 2358)

TETRA-OXY-PHENYL-BENZOPYRAZOLE

$NPh<\begin{smallmatrix} CO \\ N=C(OH) \end{smallmatrix} C(OH)C=N >NPh$ [c 150°]

Formed by the action of HCl aq on the phenylhydrazide which is made by warming di oxy quinone dicarboxylic ether with alcohol and phenyl hydrazine (Böniger, *B* 22, 1291) Begins to decompose at 125° — $(NH_4)A'$ ochre yellow powder, sol water

OXY-DI-PHENYL-BENZYLIDENE PYRAZOLE

$NPhCO<\begin{smallmatrix} N \\ N=CPh \end{smallmatrix} C>CHPh$ [147°] Formed from oxy di phenyl pyrazole and benzoic aldehyde (Knorr a Klotz, *B* 20, 2548) Needles

Di oxy-tetra phenyl benzylidene dipyrazole $(NPhCO<\begin{smallmatrix} N \\ N=CPh \end{smallmatrix} C>CH)_2OHPh$ [c 220°] Formed from benzoic aldehyde and excess of oxy-di phenyl pyrazole

EXO OXY PHENYL-BENZYL KETONE v *BENZOLIN*

Tri oxy-phenyl-benzyl ketone *Di-methyl-derivative* v *ANISOIN*

Hexa oxy-phenyl-benzyl-ketone *Hexa-methyl derivative*

$C_6H_4(OMe)COCH_2C_6H_4(OMe)_6$, [162°] Formed from hexamethoxy benzil, $HOAc$, and zinc dust (Marx, *A* 263, 255) Needles, insol cold water

OXY-PHENYL-BENZYL-METHYL PYRIMIDINE

$IDINE$ $CPh<\begin{smallmatrix} N \\ N=C(OH) \end{smallmatrix} C<\begin{smallmatrix} N \\ N=C(OH) \end{smallmatrix} CCH_2Ph$ [243°]

Formed from benzamidine hydrochloride, benzyl-acetoacetic ether, and dilute (10 p c) $NaOH$ aq (Pinner, *B* 22, 1626) Needles, insol water, v sl sol alcohol

Di oxy-phenyl-benzyl-methyl-pyrimidine

Ethyl derivative

$C_6H_4(OEt)C<\begin{smallmatrix} N \\ N=C(OH) \end{smallmatrix} C<\begin{smallmatrix} N \\ N=C(OH) \end{smallmatrix} C(OH)$ [242°] Formed from *p*-ethoxy-benzamidine hydrochloride, benzyl-acetoacetic ether, and $NaOH$ aq (Pinner, *B* 23, 2955) Needles, v sl. sol. hot alcohol

OXY-PHENYL BENZYL-PYRIMIDINE

$\text{CH}_2\text{Ph} \text{C} \begin{smallmatrix} \text{N} \text{CPh} \\ \text{N} \text{C(OH)} \end{smallmatrix} \text{CH}$ [233°] Made from phenylacetamidine hydrochloride, benzyl acetoacetic ether and dilute (10 pc) NaOHAq (Pinner, B 22, 1623) Needles, v sl sol water

Oxy-phenyl di-benzyl pyrimidine

$\text{CH}_2\text{Ph} \text{C} \begin{smallmatrix} \text{N} \text{C(CH}_2\text{Ph)} \\ \text{N} \text{C(OH)} \end{smallmatrix} \text{CPh}$ [180°] Formed by heating 'cyanbenzyl' (derived from benzyl cyanide) with HClAq (Wache, J pr [2] 39, 258)

Di oxy phenyl benzyl pyrimidine

$\text{CHPh(OH)} \text{C} \begin{smallmatrix} \text{N} \text{CPh} \\ \text{N} \text{C(OH)} \end{smallmatrix} \text{CH}$ [218°] Formed from a oxy phenyl acetamidine and benzoyl-acetic ether (Pinner, B 23, 2951) Felted needles, v sl sol water, sol alkalis and acids

OXY PHENYL BIURET

$(\text{NHPh CO})_2\text{N(OH)}$, [178°] Formed by the action of an aqueous solution of hydroxylamine on phenyl cyanate (von der Kall, A 263, 263) Needles, sl sol hot water Does not reduce Fehling's solution

p-OXY PHENYL-BROMO *p*-TOLYL THIO-UREA $\text{C}_6\text{H}_4(\text{OH}) \text{NH CS NH C}_6\text{H}_4\text{Br}$

Acetyl derivative [156°] Formed by the action of acetyl oxy phenyl thiocarbamide on bromo-*p* toluidine (Kalkhoff, B 16, 1832) V sol acetic acid, sl sol alcohol and ether, insol water

OXY PHENYL BUTANE *v* BUTYL PHENOL

Di oxy phenyl butane

$\text{CHPh(OH)} \text{C}_2\text{H}_5 \text{CH}_2\text{OH}$ (c 200°) A thick syrup formed by reduction of $\text{C}_6\text{H}_5 \text{CO C}_2\text{H}_5 \text{CHO}$ (Buicker, A Ch [5] 26, 469)

Di oxy di phenyl butane

$\text{CPhMe(OH)} \text{CPhMe(OH)}$ [120°] Formed by adding sodium amalgam to a solution of acetophenone in dilute alcohol (Emmerling, Engler, B 4, 147, 6, 1005, Buchka, B 10, 1714) Long prisms, insol water, v sol alcohol Split up by long boiling in acetophenone and phenyl methyl carbinol (Zincke, A Thorne, B 13, 641)

Di oxy tetra phenyl butane

$\text{CH}_2\text{Ph CPh(OH) CPh(OH) CH}_2\text{Ph}$ [213°] Formed, together with an isomeride [172°], by reducing benzoin with zinc (Limpriecht, A Schwanert, A 155, 60, Zagoumenny, B 7, 1651, Wischenus, A 248, 5) The compound [213°] is formed, together with an isomeride (?) [61°], by reducing benzoin with Na and alcohol (Limpriecht, A 155, 98) The compounds [172°] and [213°] are both converted by heat into phenyl benzyl ketone and phenyl benzyl carbinol

OXY-DI-PHENYL-BUTANE CARBOXYLIC ACID $\text{C}_6\text{H}_5 \text{CH}(\text{CO}_2\text{H}) \text{CH}_2 \text{CH}(\text{OH}) \text{CH}_2 \text{C}_6\text{H}_5$, *Tetrahydrocornicularic acid* Formed by reducing hydro cornicularic acid with sodium amalgam (Spiegel, A 219, 95) Thick oil, decomposed by boiling with water, yielding the lactone *Lactone* $\text{C}_{11}\text{H}_{14}\text{O}_3$ [71°]

Oxy phenyl butane dicarboxylic acid *v*. Oxy-benzyl-pyrotartaric acid

Di-oxy-di-phenyl-butane dicarboxylic acid $\text{C}_6\text{H}_4(\text{OH}) \text{CH}_2 \text{CH}(\text{CO}_2\text{H}) \text{CH}(\text{CO}_2\text{H}) \text{CH}_2 \text{C}_6\text{H}_4\text{OH}$ Formed by heating salicylic aldehyde with sodium succinate and Ac_2O , saponifying the resulting 'dicoumarin', and reducing the product with sodium amalgam (Fittig & Dyson, A

255, 281) Colourless crystals, v sol alcohol.— $\text{CaC}_{11}\text{H}_{14}\text{O}_6$ 6aq — $\text{Ag}_2\text{A}''$ bulky white pp

Lactone

$\text{C}_6\text{H}_5 \text{C} \begin{smallmatrix} \text{O CO} \\ \text{CH}_2 \end{smallmatrix} \text{CH CH} \begin{smallmatrix} \text{CO O} \\ \text{CH}_2 \end{smallmatrix} \text{C}_6\text{H}_5$, *Dicoumarin tetrahydride* [224°] Formed by heating the acid above 100° Needles (from chloroform alcohol)

Di oxy-di phenyl butane dicarboxylic acid.

Lactone $\text{C}_6\text{H}_5 \text{C} \begin{smallmatrix} \text{CH CH}_2 \text{CH}_2 \text{CH} \\ \text{CO O CO O} \end{smallmatrix} \text{C}_6\text{H}_5$

[210°] Formed by treating $\text{C}_6\text{H}_5(\text{CO C}_6\text{H}_5 \text{CO}_2\text{H})_2$ with sodium amalgam (Gabriel & Michael B 10, 2209) Needles (from alcohol), insol water The corresponding acid is unstable

DI *p*-OXY-DI-PHENYL BUTINENE *D*.

methyl ether $\text{C}_6\text{H}_4(\text{OMe}) \text{CH CH CH CH C}_6\text{H}_4(\text{OMe})$ [225°] A body formed on heating anisic aldehyde with sodium succinate and Ac_2O at 120° (Fittig & Politz, A 255, 307) Crystals, insol water, m. sol HOAc

DI-OXY-DI-PHENYL-BUTINENE DI-CARBOXYLIC LACTONE

$\text{C}_6\text{H}_5 \text{C} \begin{smallmatrix} \text{O CO CO O} \\ \text{CH C-C CH} \end{smallmatrix} \text{C}_6\text{H}_5$, *Dicoumarin* [above 330°] Made by heating salicylic aldehyde with sodium succinate and Ac_2O at 140° (Fittig & Dyson, A 255, 276) Needles (from HOAc) Not attacked by cold alkalis

O OXY PHENYL BUTYLENE DICARBOXY-

LIC ACID *Lactone* $\text{C}_6\text{H}_5 \text{C} \begin{smallmatrix} \text{O CO} \\ \text{CH C CHMe CO}_2\text{H} \end{smallmatrix}$ [171°] Made by heating salicylic aldehyde with sodium pyrotartrate and Ac_2O for 30 hours at 120° (Fittig & Brown, A 255, 285) Plates (from water) Yields $\text{Ba}(\text{C}_{11}\text{H}_{14}\text{O}_6)_2$ 3aq, CaA' 5aq, and AgA'

Di oxy di phenyl butylene dicarboxylic acid

Lactonic acid $\text{C}_6\text{H}_5 \text{C} \begin{smallmatrix} \text{O CO} \\ \text{CH C CH} \end{smallmatrix} \text{C}_6\text{H}_5$ Formed by boiling dicoumarin with NaOHAq and adding sodium amalgam (Fittig & Dyson, A 255, 277) Needles, sl sol hot water — BaA' 2aq — $\text{AgC}_{11}\text{H}_{14}\text{O}_6$ curdy pp

Lactone $\text{C}_6\text{H}_5 \text{C} \begin{smallmatrix} \text{O COCO O} \\ \text{CH C CH CH} \end{smallmatrix} \text{C}_6\text{H}_5$

[256°] Obtained by heating the lactonic acid at 130° Crystals, insol water, Na_2CO_3 aq, and NaOHAq Yields an unstable dibromide

O OXY- β -PHENYL-*iso*-BUTYRIC ACID

$\text{CH}_2\text{Ph CMe(OH) CO}_2\text{H}$ [99°] Made by the action of KCy on the bisulphite compound of benzyl methyl ketone, the product being saponified (Gabriel & Michael, B 12, 814) Long prisms, sol water and alcohol

 β -Oxy- β -phenyl-isobutyric acid

$\text{CHPh(OH) CHMe CO}_2\text{H}$ [125°] Formed by reducing a benzoyl-propionic acid with sodium-amalgam (Perkin, jun., a Calman, C J 49, 161) Needles, v e sol hot water — AgA' crystals

 β -Oxy- γ -phenyl-butyric acid

$\text{CH}_2\text{Ph CH(OH) CH}_2 \text{CO}_2\text{H}$ [98°] Formed by boiling phenyl-isocrotonic acid with NaOHAq (Fittig, B 24, 84) Flat needles.

γ -Oxy- γ -phenyl-butyric acid $\text{CHPh(OH) CH}_2 \text{CH}_2 \text{CO}_2\text{H}$ *Benzhydrylic propionic acid* [75°] Formed by reducing benzoyl-propionic acid with sodium-amalgam (Buroker,

Bl [2] 87, 5, *A Ch* [5] 26, 455), and also by boiling its lactone with baryta. Flat crystals (from CS_2), splitting up at 70° into water and lactone. Chromic acid mixture oxidises it to benzoyl propionic acid— BaA'_2 — CuA'_2 — AgA' crystalline pp

Lactone $\text{CHPh} \begin{smallmatrix} \text{O} \\ \diagup \text{---} \text{CO} \\ \diagdown \end{smallmatrix} \text{CH}_2\text{CH}_2$ [37°] (306°)

Formed from γ bromo γ -phenyl butyric acid by treatment with Na_2CO_3 or boiling with water (Jayne, *A* 216, 103). Formed also by boiling phenyl paraconic acid with diluted (1:1) H_2SO_4 (Erdmann, *A* 228, 178). Six sided trimetric tables (from CS_2), $a b c = 611.1 \ 426$. It has a pleasant odour, and is volatile with steam.

Amide $\text{CHPh}(\text{OH})\text{CH}_2\text{CH}_2\text{CO}\text{NH}_2$ [86°] Formed by heating the lactone with alcoholic NH_3 at 100° (Fittig, *A* 256, 155). Monoclinic prisms, v sol hot water. Forms unstable $\text{C}_6\text{H}_5\text{NO}_2\text{HCl}$, crystallising in needles.

***o*-Oxy phenyl butyric acid** *Methyl derivative* $\text{C}_6\text{H}_4(\text{OMe})\text{CH}_2\text{CH}_2\text{CO}_2\text{H}$ [56°] Obtained by reducing either (α)- or (β) methoxy-phenyl crotonic acid with sodium amalgam (Perkin, *C J* 39, 433). Transparent prisms (from light petroleum). May be distilled— BaA'_2 (dried at 100°).

***a*-Oxy-di phenyl-isobutyric acid** *Di benzyl-glycollic acid*

γ -Oxy-di- γ phenyl butyric acid $\text{CPh}_2(\text{OH})\text{CH}_2\text{CH}_2\text{CO}_2\text{H}$ [145°] Made from its lactone (Auger, *A Ch* [6] 22, 316). Thin leaflets (from alcohol). Changes to lactone on melting, but is stable at 100° — BaA'_2 , micaceous plates, sol water.

Lactone $\text{CPh}_2 \begin{smallmatrix} \text{O} \\ \diagup \text{---} \text{CO} \\ \diagdown \end{smallmatrix} \text{CH}_2\text{CH}_2$ [90°] Formed by

the action of succinyl chloride on benzene in presence of AlCl_3 . Leaflets, insol water, sl sol cold alcohol.

Oxy-tri-phenyl-butyric acid

$\text{CHPh}(\text{OH})\text{CH}(\text{OH})\text{CH}_2\text{CO}_2\text{H}$ Formed by reducing di-phenyl-benzoyl propionic acid in alkaline solution with sodium amalgam (Japp a Klingemann, *C J* 87, 680, *B* 22, 2882). White solid, sol $\text{Na}_2\text{CO}_3\text{Aq}$.

Lactone $\text{C}_6\text{H}_5\text{O}_2$ [153°] Small needles, sol hot alcohol.

β -Di-oxy- γ -phenyl butyric acid

$\text{CHPh}(\text{OH})\text{CH}(\text{OH})\text{CH}_2\text{CO}_2\text{H}$ [118°] Made from its lactone, which is got by oxidising phenyl isocrotonic acid with KMnO_4 (Fittig, *B* 21, 920). Crystals, forming the lactone on melting.

Lactone $\text{CHPh} \begin{smallmatrix} \text{O} \\ \diagup \text{---} \text{CO} \\ \diagdown \end{smallmatrix} \text{CH}(\text{OH})\text{CH}_2$ [87°]

Melts when hydrated (with $\frac{1}{2}$ aq) at 77° .

Di oxy phenyl butyric acid

Methyl derivative

[4:1] $\text{C}_6\text{H}_4(\text{OMe})\text{CH}(\text{OH})\text{CH}_2\text{CH}_2\text{CO}_2\text{H}$ Got from its lactone [53.5°], which is made by the action of sodium amalgam on bromo *p* methoxy-phenyl-butyrolactone (Fittig a Politz, *A* 256, 299). The free acid is solid, and yields the lactone when heated to 80° — BaA'_2 , amorphous mass.

Di-oxy-phenyl-isobutyric acid

Methyl derivative

[4:3:1] $\text{C}_6\text{H}_4(\text{OH})(\text{OMe})\text{CH}_2\text{CHMeCO}_2\text{H}$ *Hydrohomoferric acid* [115°] Obtained by reducing $\text{C}_6\text{H}_4(\text{OH})(\text{OMe})\text{CHMeCO}_2\text{H}$ (Tiemann a.

Kraaz, *B* 15, 2070). Sol water, alcohol, and ether.

Di-methyl derivative

$\text{C}_6\text{H}_4(\text{OMe})\text{CH}_2\text{CHMeCO}_2\text{H}$ *Methyl hydrohomoferric acid* [59°] Got in like manner. Crystals.

Methylene derivative

$\text{C}_6\text{H}_4(\text{O}_2\text{CH}_2)\text{CH}_2\text{CHMeCO}_2\text{H}$ *Hydrohomocaffic acid* [77°] Formed by reduction of the methylene derivative of di oxy phenyl methacrylic acid (Lorenz, *B* 13, 760). Thick prisms, sl sol water, v sol alcohol and ether.

Di-oxy di phenyl butyric acid?

$\text{CH}_2\text{CPh}(\text{OH})\text{CPh}(\text{OH})\text{CO}_2\text{H}$ Formed from its nitrile, which is made from acetophenone, KOH , and aqueous K_2FeCy_4 (Buchka, *B* 20, 389)— BaA'_2 , 3:1 aq.

***o*-OXY-PHENYL-CARBAMIC ACID** *Ethyl*

ether $\text{C}_6\text{H}_4(\text{NH})\text{CO}_2\text{Et}$ *ie* $\text{C}_6\text{H}_4(\text{OH})\text{NHCO}_2\text{Et}$ [85°] Formed from *o* amido phenol and ClCO_2Et (Groenink, *Bl* [2] 25, 177). Triclinic prisms (from ether alcohol), almost insol cold water. Yields, on distillation, alcohol and the anhydride.

Anhydride $\text{C}_6\text{H}_4 \begin{smallmatrix} \text{NH} \\ \diagup \text{---} \text{CO} \\ \diagdown \end{smallmatrix} \text{O}$ or

$\text{C}_6\text{H}_4 \begin{smallmatrix} \text{N} \\ \diagup \text{---} \text{C} \text{OH} \end{smallmatrix} \text{O}$ *Oxycarbonyl Oxymethenyl amido phenol* *Anhydro o amido phenyl carbonic acid* *Carbonyl amido phenol* [138°] (above 360°).

Formation—1 As above—2 By heating oxy phenyl urea (Kalkhoff, *B* 16, 1828)—3 By distilling *o* amido phenyl ethyl carbonate (Bender, *B* 19, 2269, 2950)—4 By heating a mixture of urea and *o* amido phenol (Sandmeyer, *B* 19, 2655)—5 By the action of phosgene on *o*-amido phenol in benzene (Chetnicki, *B* 20, 177, Jacoby, *J pr* [2] 37, 29)—6 By heating *o* amido phenyl di phenyl carbamate (Lellmann a Bonhoeffer, *B* 20, 2126).

Properties—Needles (from water), sol alcohol, ether, and alkalis. Decomposed by heating with HClAq above 150° into CO_2 and *o* amido phenol. Bleaching powder and HCl yield crystalline $\text{C}_6\text{H}_4\text{Cl} \begin{smallmatrix} \text{NCl} \\ \diagup \text{---} \text{CO} \\ \diagdown \end{smallmatrix} \text{O}$, whence

further action of HClAq forms $\text{C}_6\text{H}_4\text{Cl} \begin{smallmatrix} \text{NH} \\ \diagup \text{---} \text{CO} \\ \diagdown \end{smallmatrix} \text{O}$ [196°]. Br forms $\text{C}_6\text{H}_4\text{BrO} \begin{smallmatrix} \text{NH} \\ \diagup \text{---} \text{CO} \\ \diagdown \end{smallmatrix} \text{O}$ [196°]. HNO_3 produces $\text{C}_6\text{H}_4(\text{NO}_2)_2\text{NH}$ [206°]. Injected into rabbits, it becomes $\text{C}_6\text{H}_4(\text{OSO}_3\text{H}) \begin{smallmatrix} \text{NH} \\ \diagup \text{---} \text{CO} \\ \diagdown \end{smallmatrix} \text{O}$

(Nencki, *M* 11, 253)— AgO_2HNO_2 , curdy pp.

Acetyl derivative $\text{C}_6\text{H}_4\text{O}_2\text{Nac}$ [98°]

Crystallises from water.

Phenyl-hydrazide

$\text{C}_6\text{H}_4 \begin{smallmatrix} \text{NH} \\ \diagup \text{---} \text{C} \text{N}_2\text{HPh} \end{smallmatrix} \text{O}$ [208°] Yellow needles.

Ethyl derivative $\text{C}_6\text{H}_4 \begin{smallmatrix} \text{NEt} \\ \diagup \text{---} \text{CO} \\ \diagdown \end{smallmatrix} \text{O}$.

[29°] (300°) Formed from the silver salt and EtI . Crystalline. Converted into ethyl amido-phenol by heating with fuming HClAq at 180° .

Ethyl ether $\text{C}_6\text{H}_4 \begin{smallmatrix} \text{N} \\ \diagup \text{---} \text{COEt} \end{smallmatrix} \text{O}$ (225°).

230° Formed from amido phenol hydrochloride and $\text{NH}_4\text{C}(\text{OEt})_2$ (Sandmeyer). Liquid, converted by HClAq into EtCl and $\text{C}_6\text{H}_4 \begin{smallmatrix} \text{NH} \\ \diagup \text{---} \text{CO} \\ \diagdown \end{smallmatrix} \text{O}$.

p-Oxy phenyl-carbamic acid Ethyl ether
 $C_8H_{11}NO_3$, *ie* $C_6H_4(OH)NHCO_2Et$ [120°]
 Formed from p-amido phenol and $ClCO_2Et$
 (Groenink, *Bl* [2] 25, 179) Monochmic tables,
 v sol alkalis. The ethyl derivative
 $C_8H_9(CEt)NHCO_2Et$, formed from $ClCO_2Et$ and
 $C_6H_5(NH_2)(OEt)$ (Köhler, *J pr* [2] 29, 257),
 crystallises in needles [94°], insol water

Reference — CHLORO OXY-PHENYL CARBAMIC
 ACID

p OXY-DI-PHENYL-CARBINOL $C_{15}H_{13}O_3$, *ie*
 $C_6H_5CH(OH)C_6H_4(OH)$ [161°] Formed by
 reducing oxy benzophenone with sodium amal
 gam (Doebner, *A* 210, 253) Silky needles (from
 water) Coloured red by $FeCl_3$

Di-oxy di-phenyl carbinol
 $CH(OH)(C_6H_5)_2$, [160°-165°] Made by
 reducing di oxy benzophenone (salicyl phenol)
 with sodium amalgam (Michael, *B* 14, 657)
 Amorphous powder, sol hot alcohol

Di oxy tri phenyl carbinol
 $C_6H_5C(OH)(C_6H_5)_2$, [c 100°]. Made by
 warming $PhCCl_3$ with dry phenol and treating
 the product with water (Doebner, *A* 217, 227)
 Formed also by warming phenyl glyoxylic acid
 with phenol and H_2SO_4 at 120° (Homolka, *B*
 18, 988) Brick red powder, insol cold water,
 melts under hot water Its alcoholic solution
 dyes silk feebly yellow In alkalis it dissolves
 with deep violet red colour, and is reppd by
 acids

Di acetyl derivative
 $C_6H_5C(OH)(C_6H_5OAc)_2$, [119°] Colourless
 prisms (from dilute alcohol)

Tri oxy tri phenyl carbinol Anhydride
 $C_{15}H_9O_3$, *ie* $(C_6H_5(OH))_2C < \begin{smallmatrix} C_6H_5 \\ O \end{smallmatrix} >$ Aurin
 Rosolic acid

Formation — 1 By heating phenol with
 oxalic acid and H_2SO_4 (Kolbe a Schmitt, *A* 119,
 169, Dale a Schorlemmer, *A* 196, 79) — 2 By
 heating phenol with formic acid and $ZnCl_2$ at
 120° (Nencki a Schmid, *J pr* [2] 23, 549, 25,
 273) — 3 By the action of $AlCl_3$ on a mixture of
 phenol and tri chloro nitro methane (Elbs, *B*
 16, 1275) — 4 From para rosaniline by the
 diazo reaction (Fischer, *A* 194, 268) — 5 From
 di p oxy benzophenone by treatment with PCl_5 ,
 and heating the product with phenol and H_2SO_4
 (Caro a Graebe, *B* 11, 1350)

Preparation — Phenol (10 pts) is heated with
 dried oxalic acid (7 pts) and H_2SO_4 (5 pts) at
 120°-130° about 24 hours, until gas no longer
 comes off rapidly The product is poured into
 water, the pp dissolved in $NaOH$ aq, saturated
 with SO_2 , and mixed with much water The
 filtrate from ψ rosolic acid (which amounts to
 70 p c of crude product) is saturated at 70° with
 HCl , and on cooling it deposits aurin sulphite
 This is recrystallised from dilute alcohol, which
 deposits methyl aurin on cooling, while the
 mother liquor, saturated with SO_2 , deposits aurin
 sulphite, which is freed from SO_2 by heat (Zul-
 kowsky, *A* 194, 119, 202, 184)

Properties — Dark red trimetric crystals
 (from alcohol $HOAc$) or red needles with green
 lustre (from alcohol) Not melted below 220°
 Its alkaline solution is crimson

Reactions — 1 Reduced by *smc-dust* and
 $HOAc$ to tri-oxy-tri phenyl methane — 2 Aque-
 ous NH_3 at 120° yields para-rosaniline. — 3.

Vol. III

Water at 250° yields phenol and di p-oxy
 benzophenone — 4 On warming with KOH aq
 and KCy and adding HCl , a product is got
 which, when heated with Ac_2O , yields tri-acetyl
 hydrocyanaurin [194°]

Salts — $(NH_4)_2C_{15}H_{13}O_3$, dark red needles
 with steely lustre — $C_{15}H_{13}O_3 \cdot HCl$, $HOEt$ red
 crystals — $(C_6H_5O_2HCl)HOAc$ red needles —
 $(C_{15}H_{13}O_3)_2 \cdot H_2SO_4$, 4aq brick red cubes, sl sol
 cold alcohol — $C_{15}H_{13}O_3 \cdot (NH_4)HSO_4$ —
 $C_{15}H_{13}O_3 \cdot NaHSO_4$ — $C_{15}H_{13}O_3 \cdot KHSO_4$, minute
 colourless tables — $C_{15}H_{13}O_3 \cdot H_2SO_4$ —
 $(C_{15}H_{13}O_3)_2 \cdot H_2SO_4$, bluish violet needles.

Di acetyl-derivative
 $(C_6H_5(OH))_2C(OAc)_2$, C_6H_5OAc [168°] Formed
 from aurin and Ac_2O at 100° Colourless tables
 (from alcohol)

Tetra-oxy-tri-phenyl-carbinol
 $C_{15}H_9O_4$, *ie* $C_6H_5C(OH)(C_6H_5(OH))_2$

Anhydride $C_{15}H_5O_4$, *Resorcin benzetrin*.
 Formed by heating benzo trichloride with re-
 sorcin at 180° (Doebner, *B* 13, 610, *A* 217,
 234) Large crystals (from alcohol and $HOAc$)
 Yellow by transmitted, violet red by reflected
 light, at 130° it loses $2H_2O$, and at 200° it is
 decomposed Dilute alkaline solutions show
 yellowish green fluorescence, but less intense
 than fluorescein, they dye wool yellow Insol
 water, v sol alcohol Zinc dust and HCl re-
 duce it to tetra oxy tri phenyl methane Bromine
 in alcohol and $HOAc$ forms a fiery red pp
 $C_{15}H_9Br_2O_4$, which closely resembles eosin Its
 salts dissolve readily in alcohol and dye wool
 and silk like eosin

Penta-oxy-tri-phenyl-carbinol
 $(C_6H_5(OH))_3C(OH)C_6H_5OH$
Anhydride $C_{15}H_5O_5$, Formed by heating
 resorcin with formic acid and $ZnCl_2$ at 140°
 (Nencki a Schmid, *J pr* [2] 23, 547) Hygro-
 scopic brick red powder, v sol alcohol

p OXY-DI-PHENYL-CARBINOL o-CARB-
OXYLIC ACID

Methyl derivative of the anhydride
 $C_6H_5 < \begin{smallmatrix} CH \\ CO \\ O \end{smallmatrix} C_6H_5OMe$ [117°] Formed by re-
 ducing methoxy benzophenone carboxylic acid
 with zinc and alcoholic HCl (Nourrisson, *B* 19,
 2103) Flat white needles, v sol warm alcohol,
 insol water

Oxy-tri-phenyl-carbinol carboxylic acid

Anhydride $C_{15}H_9(OH)CPh < \begin{smallmatrix} C_6H_5 \\ O \end{smallmatrix} > CO$

Oxy di phenyl phthalide [155°] Prepared by
 heating o benzoyl benzoic acid with phenol and
 $SnCl_4$ at 120°, yield 100 p c of the benzoyl-
 benzoic acid (Fechmann, *B* 13, 1613) Colour-
 less crystals Sol all ordinary solvents except
 water and ligroin By fusion with KOH it gives
 benzoic acid and oxy-benzophenone With
 alkalis it forms a deep-red solution By strong
 H_2SO_4 it is converted into oxy-phenyl anthranol
 $C_6H_5 < \begin{smallmatrix} C(OH) \\ C(C_6H_5OH) \end{smallmatrix} > C_6H_5$ On reduction it
 gives oxy triphenyl methane carboxylic acid

Acetyl derivative $C_{22}H_{15}O_5(OAc)$, [186°].
 Colourless crystals

Di-bromo-derivative $C_{20}H_{11}Br_2O_4(OH)$.
 [196°] Spikes

Acetyl di bromo-derivative
 $C_{20}H_{11}Br_2O_4(OAc)$ [172°] Colourless prisms.

***o*-di-oxy-di-phenyl-carbinol carboxylic anhydride** *Methyl derivative*

$C_6H_5 \begin{array}{c} \text{CH} \begin{array}{c} \text{CO} \end{array} \begin{array}{c} \text{CH} \end{array} \begin{array}{c} \text{O} \end{array} \end{array}$ [117°] Made by reducing '*p* anisolphthaloylic' acid (Nourrisson, *B* [2] 46, 206) Needles, insol water, v e sol alcohol

Di-oxy-tri-phenyl-carbinol carboxylic acid
Anhydride $C_6H_5(OH)_2CPh \begin{array}{c} \text{CO} \end{array}$

Benzene resorcin phthalate [176°] Made by heating benzoyl benzoic acid with resorcin (Pechmann, *B* 14, 1859) Prisms (containing $CHCl_3$) melting at 114° (from chloroform) Its alcoholic solution turns greenish blue on addition of HCl On heating with H_2SO_4 it yields anthraquinone, H_2SO_4 acting on its solution in $HOAc$ forms $C_{20}H_{12}O_2$ [285°], which yields $C_{20}H_{12}O_2$, Ac_2O , [245°]

Di acetyl derivative $C_{20}H_{12}O_2(OAc)_2$, [137°]

Di-bromo-derivative $C_{20}H_{12}Br_2O_4$, [219°]

Tri-oxy-tri-phenyl-carbinol-carboxylic acid
Anhydride $(C_6H_5(OH)_3)CPh \begin{array}{c} \text{CO} \end{array}$

Benzene pyrogallol-phthalate Prepared by heating benzoyl benzoic acid with pyrogallol and crystallising the product from acetic acid (Pechmann, *B* 14, 1864) Four-sided tables (containing $AcOH$), [190°] Sol most solvents except ligroin, sl sol hot water It dissolves in alkalis with a green colour

Tri acetyl derivative $C_{20}H_{12}O_2(OAc)_3$, [231°] Fine needles Sol Ac_2O

DI-OXY DI-PHENYL-CARBOLACTONE *v.* *ISO-EUXANTHIC ACID*

OXY-DI-PHENYL-DI-CARBOXYLIC ACID
Hexahydrate $C_{12}H_{10}O_6$ An unstable acid formed by boiling hydro oxybenzyluric acid with potash (Otto, *A* 184, 330) Yields Et_2A (206°), a heavy oil

Di-*p*-oxy-diphenyl carboxylic acid $C_{12}H_{10}O_6$, *s.e.* [4 1] $C_6H_5(OH)(CO_2H)(CO_2H)$ [1 4 2] [270°] Made by fusing diphenylene ketone disulphonic acid with potash (Schmidt & Schultz, *B* 12, 496) Sl sol water, v sol hot alcohol

Di-*p*-oxy-diphenyl dicarboxylic acid
 $C_6H_5(OH)(CO_2H)C_6H_5(OH)CO_2H$ [151°] S 0052 at 15°. Formed by heating sodium di-oxy di phenyl with CO_2 at 200° under high pressure (Schmitt & Kretschmer, *B* 20, 2703) Minute needles, sl sol water, $FeCl_3$ colours its solution bluish violet

Tetra oxy-diphenyl dicarboxylic acid
 $C_6H_5(OH)_2(CO_2H)C_6H_5(OH)_2CO_2H$ *Di-resorcin dicarboxylic acid* Formed by heating tetra-oxy diphenyl (diresorcin) with $KHCO_3$ and a little water at 130° (Will & Albrecht, *B* 17, 2105) Yellowish powder, decomposing, without melting, above 300°. — K_2A —. — BaA ° 6aq — Ag_2A °. white pp

Tetra-oxy-diphenyl dicarboxylic acid
 $C_6H_5(OH)_2(CO_2H)C_6H_5(OH)_2(CO_2H)$ *Dehydro-diprotocatechus acid* [above 300°] Formed by fusing 'dehydrodivanillin' with potash (Tiemann, *B* 18, 3495). Amorphous, v. sl. sol water, sl sol alcohol

Penta-oxy-diphenyl carboxylic acid
 $C_{11}H_{10}O_6$ Formed by the action of $POCl_3$ on *o*-tri-oxy-benzoic acid (Schiff, *G* 17, 552, *A*.

245, 87) Yellowish astringent powder — BaA °. greyish white powder

Penta-acetyl derivative $C_{11}H_4Ac_5O_6$. White powder, rapidly darkening in air

Ethyl ether EtA° [102°]

Penta-oxy di phenyl carboxylic acid

$C_{11}H_{10}O_6$ Formed by heating phloroglucin carboxylic acid with $POCl_3$ (Schiff) Reddish brown hygroscopic mass, resembling tannin and the preceding isomeride

TETRA-OXY-DIPHENYL DICARBOXYLIC

ALDEHYDE *Di-methyl derivative*

$C_6H_5(OMe)(OH)(CHO)C_6H_5(OMe)(OH)CHO$

Divanillin [304°] Formed by boiling vanillin $C_6H_5(OMe)(OH)(CHO)$ [3 4 1] with $FeCl_3$ (Tiemann, *B* 18, 3498) Slender white needles, sl sol alcohol, sol alkalis

Tetra methyl derivative [138°]

***p* OXY-PHENYL-CINNAMIC ACID** *Me-*

thyl derivative $C_6H_5(OMe)CHCPhCO_2H$ [189°] Formed from sodium phenyl acetate, anisic aldehyde, and Ac_2O (Oghaloro, *G* 9, 533, 10, 481) Prisms (from alcohol), sl sol water

***a* OXY- γ PHENYL- α CROTONIC ACID**

$C_6H_5CHCHCH(OH)CO_2H$ [115°] Formed by boiling cinnamic aldehyde with aqueous H_2Cy and HCl (Matsumoto, *B* 8, 1144, Peine, *B* 17, 2114) Formed also by heating the ethyl ether of styryl ψ hydantoin with concentrated baryta water (Pinner & Spulker, *B* 22, 690) Styryl ψ

hydantoin $CHPhCHCH \begin{array}{c} \text{CO NH} \\ \text{O} \end{array} \begin{array}{c} \text{O} \end{array} \begin{array}{c} \text{CNH} \end{array}$ [198°] is obtained by the action of warm alcoholic potash on styryl hydantoin [172°], an isomeric body previously described by Pinner (*B* 20, 2353) as oxystyryl pyrazole Styryl hydantoin is got by the action of boiling dilute $HClAq$ on (*a*) ur

amido phenyl crotonic nitrile $CHPhCHCH(NHCO_2NH_2)CN$ [160°], which is made by heating oxy phenyl isocrotonic nitrile with urea

Properties — Needles, sl sol cold water, v sol alcohol and ether

Salts — PbA °, 2aq needles — AgA ° minute needles

Methyl ether MeA°. (290°) Liquid

Ethyl ether EtA° (295°) (Peine)

Nitrile $CHPhCHCH(OH)CN$ [81°]

Formed from cinnamic aldehyde, KCy , and HCl Crystalline grains (from benzene ligroin) Converted by hydroxylamine into the crystalline amidoxim $CHPhCHCH(OH)C(NH_2)NOH$ (Bornemann, *B* 19, 1513)

***o*-Oxy-phenyl-crotonic acid** (*a*) *Methyl*

derivative $C_6H_5(OMe)CHCO_2H$ [118°]

Made from its methyl ether by boiling with alcoholic potash (Perkin, *C* J 33, 213, 39, 431) Monoclinic crystals (from alcohol), $abc = 863 \ 1 \ 1 \ 252$, $\beta = 64^\circ 54'$ (Fletcher) Not converted into its (*B*) isomeride by light With PCl_5 it gives $MeCl$, HCl , and the anhydride Conc H_2SO_4 also forms propionic coumarin Sodium amalgam reduces it to methoxy phenyl butyric acid With Br and with HI it behaves like the (*B*) acid

Methyl ether of the (a)-methyl derivative $C_6H_5(OMe)CHCO_2H$ (275°) S $\frac{1}{2}$ 1 1112, $\frac{1}{2}$ 1 1061 Formed from sodium propionic coumarin and MeI Liquid

(*B*) *Methyl derivative*

$C_6H_5(OMe)CHCO_2H$ [107°] Formed

from methyl salicylic aldehyde, sodic propionate and propionic acid

Properties — Monoclinic crystals, $a b c = 1.281 \ 1.762 \ \beta = 84^\circ 18'$ (Fletcher) With PCl, it gives the chloride of the acid Conc H_2SO_4 appears to polymerise it Behaves like the (a) isomeride with sodium amalgam With dry bromine vapour it forms the compound $C_6H_4Br_2(OMe)CHBrCMeBrCO_2H$ [c 200°] With HI (S G 1 94) it unites, forming a product whence Na_2CO_3 liberates the methyl derivative of o-allyl phenol — AgA' white pp

Methyl ether of the (B)-methyl derivative MeA' (c 286°) SG $\frac{15}{16}$ 1.1279, $\frac{30}{32}$ 1.1136

(B) **Ethyl derivative** $C_6H_4(OEt)CHCMeCO_2H$ [133°] Formed from ethyl-salicylic aldehyde, sodic propionate, and propionic anhydride Large tables (from alcohol) Its Ba salt forms satiny needles

Anhydride $C_6H_4 \begin{smallmatrix} CHCMe \\ \diagup \quad \diagdown \\ O \quad CO \end{smallmatrix}$ **Propionic**

coumarin Methyl-coumarin [90°] (292 5°) Formed by heating sodium salicylic aldehyde with propionic anhydride (Perkin, C J 28, 10) Trimetric crystals, $a b c = 2.1950 \ 1.4001$ Smells like coumarin, m sol alcohol, nearly insol cold KOHAq Fuming H_2SO_4 forms $C_{10}H_8O_4(SO_3H)$ whence BaA' , 10aq

Oxy-phenyl-crotonic acid **Anhydride** $C_6H_4 \begin{smallmatrix} CHCMe \\ \diagup \quad \diagdown \\ O \quad CO \end{smallmatrix}$ β **Methyl coumarin** [126°]

Formed by the action of H_2SO_4 on a mixture of acetoacetic ether and phenol (Pechmann a. Duisberg, B 16, 2127)

p-Oxy-phenyl-crotonic acid **Methyl derivative** $C_6H_4(OMe)CHCMeCO_2H$ [154°] Formed from anisic aldehyde, sodium propionate, and propionic anhydride Rectangular tables (from alcohol) — AgA'

p-Oxy-phenyl-isocrotonic acid **Methyl derivative** $C_6H_4(OMe)CHCHCH_2CO_2H$ [106 5°] Formed by heating anisic aldehyde with sodium succinate and Ac_2O at 120° (Fittig a. Politis, A 255, 294) Plates, m sol hot water — BaA' , 3aq — CaA' , 2aq — AgA' sol hot water

γ-Oxy-tri phenyl crotonic acid **Lactone** $C_6H_4O_3 \begin{smallmatrix} CHCPh \\ \diagup \quad \diagdown \\ CPh_2CO \end{smallmatrix} > O$ (?) [118°] Mol w

329 (by Raoult's method), 312 (calc) Formed by heating aB di benzoyl styrene at 310° (Japp a. Klingemann, C J 57, 679, 702, 59, 148) Lustrous needles or monoclinic prisms Not attacked by phenyl hydrazine or Ac_2O at 150° Bromine forms $C_{22}H_{15}BrO_2$ [109°] Chromic acid mixture oxidises it to benzophenone May be reduced to oxy tri phenyl butyric acid Alcoholic potash at 100° yields $CH_2BzCPh_2CO_2H$, which at 310° changes to the original lactone Alcoholic methylamine forms $CH_2BzCPh_2CONHMe$ [156°] Ethylamine acts in like manner

γ-Oxy-tetra phenyl-crotonic acid. Lactone $C_6H_4CPhCPh \begin{smallmatrix} CHCPh \\ \diagup \quad \diagdown \\ CPh_2CO \end{smallmatrix} > O$ **Tabular oxylepiden** [136°] S (alcohol) 7 at 78° Formed by heating 'acoular oxylepiden' $CPhBzCPhBz$ at 340° (Zinn, J R 5, 16, Bn. 3, 113, Japp a. Klingemann, C J 57, 665, Klingemann a. Laycock, C J 59, 140) Tables (from alcohol), insol. water. Converted by alcoholic potash into $C_{22}H_{15}KO_3$. Methyl-

amine in alcohol at 100° forms $CHPhBzCPh_2CONHMe$ [267°] crystallising in triclinic plates

Di oxy-phenyl-crotonic acid $C_6H_4(OH)_2CMeCHCO_2H$ (B) **methyl-umbelliferone acid**

Methyl derivative [4 2 1] $C_6H_4(OH)(OMe)CMeCHCO_2H$ [140°] Formed by boiling the methyl derivative of (B)-methyl umbelliferone with conc KOHAq for 5 hours (Pechmann a. Duisberg, B 16, 2125). Four sided tables, insol water, v sol. alcohol. Reconverted into its lactone by boiling with acids or heating with NH_3Aq

Di-methyl derivative $C_6H_4(OMe)_2CMeCHCO_2H$ [145°] Formed by saponification of its methyl ether (310°-320°) which is obtained by further methylation of the methyl derivative (Pechmann a. Cohen, B 17, 2132) Small needles, v sol alcohol — AgA' . white powder

Anhydride $C_6H_4(OH) \begin{smallmatrix} CHCMe \\ \diagup \quad \diagdown \\ O \quad CO \end{smallmatrix}$ (B)-

Methyl umbelliferone [186°] Formed by the action of H_2SO_4 on a mixture of acetoacetic ether and resorcin (Pechmann a. Duisberg, B 16, 2119, A 261, 169, cf Schmid, J pr [2] 25, 82) Formed also by warming acetoacetic ether with resorcin and alcoholic potash, and heating the resulting methyl-umbelliferone carboxylic acid [191°] above its melting point (Michael, J pr [2] 37, 469) (B) methyl umbelliferone is

also obtained by heating citric acid with resorcin and H_2SO_4 (Wittenberg, J pr [2] 24, 123) Plates (by sublimation), sol alcohol and hot water, sl sol ether Its dilute alkaline solution has blue fluorescence Potash-fusion forms di oxy acetophenone Yields a nitro compound, converted by reduction into amido-(B) methyl umbelliferone [247°], whence $B'H_2SO_4$, 2aq and $C_6H_5(NO)NO$, Sodium-amalgam reduces (B)-methyl umbelliferone to $C_{10}H_{10}O_2$ [259°], whence $C_{10}H_8AcO_2$ [222°] (Michael, Am. 5, 436)

Acetyl derivative of the anhydride $C_{12}H_{10}O_4$ [150°] Long white needles, v sol. alcohol

Benzoyl derivative of the anhydride $C_{10}H_8BzO_2$ [160°] Needles (from alcohol)

Methyl derivative

$C_6H_4(OMe) \begin{smallmatrix} CHCMe \\ \diagup \quad \diagdown \\ O \quad CO \end{smallmatrix}$ [159°] Needles (from alcohol), insol water On reduction with sodium amalgam it yields $C_6H_4(OMe) \begin{smallmatrix} CHMe \\ \diagup \quad \diagdown \\ O \quad CO \end{smallmatrix}$ [244°] Bromine in chloroform reacts forming $C_6H_4(OMe) \begin{smallmatrix} CHMeBr \\ \diagup \quad \diagdown \\ O \quad CO \end{smallmatrix}$ [235°]

(4 3 1) **Di-oxy-phenyl-crotonic acid** $C_6H_4(OH)_2CHC(CH_3)CO_2H$ [4 3 1] **Homocafferic acid**

p-Propionyl-m methyl derivative $C_6H_4(OCOCH_3)(OMe)CHC(CH_3)CO_2H$ **Propiohomoferrulic acid** [129°] Formed by heating vanillin with sodium propionate and propionic anhydride (Tieman a. Kraaz, B 15, 2060). White needles, sol alcohol, ether, and benzene, insol water and ligroin.

m-Methyl derivative $C_6H_4(OH)(OMe)CHC(CH_3)CO_2H$ **Homoferrulic**

acid [168°]. Large trimetric tables Sol alcohol and ether, al. sol chloroform, benzene, and hot water, nearly insol cold water and ligroin. On heating the acid it splits off CO_2 forming isoeugenol— A^1Ba yellow needles.

Di-methyl derivative
 $\text{C}_6\text{H}_4(\text{OMe})_2\text{CH}(\text{CO}_2\text{H})\text{CO}_2\text{H}$ *Methyl-homoferric acid* [141°] Needles Sol alcohol, ether, and hot water— A^1Ag white crystalline pp

Di-methyl-derivative-methyl ether
 $\text{C}_6\text{H}_4(\text{OMe})_2\text{CH}(\text{CO}_2\text{Me})\text{CO}_2\text{Me}$ [66°] Colourless plates

Methylene derivative

$\text{CH}_2\text{C}(\text{O})\text{C}_6\text{H}_4\text{CH}(\text{CO}_2\text{H})\text{CO}_2\text{H}$ [194°] Formed by boiling piperonal with propionic anhydride and sodium propionate (Lorenz, *B* 13, 759) Prisms (from dilute alcohol)— AgA^1 pp

Tri-oxy-phenyl-crotonic acid Anhydride

$\text{C}_6\text{H}_3(\text{OH})_3\text{C}(\text{OMe})\text{CH}(\text{CO})\text{CO}$ [235°] Prepared by the action of H_2SO_4 on a mixture of acetoacetic ether and pyrogallol (Wittenberg, *J pr* [2] 26, 68, Pechmann & Duisberg, *B* 16, 2127) Needles, sol. alcohol and hot water. Coloured green by FeCl_3 . Yields a di-acetyl derivative [176°]. An isomeride [284°] is formed by using phloroglucin instead of pyrogallol, it yields a di acetyl derivative [140°] (Pechmann & Cohen, *B* 17, 2189)

Tetra-oxy-phenyl-crotonic acid. Methylene-di-methyl derivative

$\text{CH}_2\text{C}(\text{O})\text{C}_6\text{H}_2(\text{OMe})_2\text{CH}(\text{CO}_2\text{H})\text{CO}_2\text{H}$ [209°].

Formed by heating apionic aldehyde with propionic aldehyde and sodium propionate (Ciamician & Silber, *B* 22, 2488) Yellow needles, almost insol water— CaA^1 , 5aq— AgA^1 white gelatinous pp

OXY-PHENYL-CUMINYLAMINE v CUMINYLAMIDO-PHENOL

OXY-PHENYL-CUMYLAMINE v CUMINYLAMIDENE-AMIDO-PHENOL

o-OXY-PHENYL-CYANAMIDE Ethyl derivative $\text{C}_6\text{H}_4(\text{OEt})\text{NHCO}$ [94°] Made by passing gaseous ClCN into an ethereal solution of *o*-amido phenetole (Berlinerblau, *J pr* [2] 30, 100). Crystals Insol water, v sol alcohol and ether. Does not appear to polymerise With fuming HCl at 120° it gives *o*-amido phenol

Salts— $\text{C}_6\text{H}_4\text{N}_2\text{ONa}$ Microscopic needles, got by mixing alcoholic solutions of NaOEt and ethoxy phenyl cyanamide Its aqueous solution does not absorb CO_2 — $\text{C}_6\text{H}_4\text{N}_2\text{OAg}$ Curdy pp

p-Oxy-phenyl-cyanamide Ethyl derivative $\text{C}_6\text{H}_4(\text{OEt})\text{NHCO}$. [78°] Made in the same way as the *o*-compound Colourless crystals Insol water, v sol. alcohol and ether— $\text{C}_6\text{H}_4\text{N}_2\text{OAg}$.

OXY-PHENYL-CYANATE. Ethyl derivative $\text{C}_6\text{H}_4(\text{OEt})\text{NCO}$ [219°] Made by distilling $\text{C}_6\text{H}_4(\text{OEt})\text{NHCO}_2\text{Et}$ (Köhler, *J pr* [2] 29, 259) White needles (from HOAc), sol alcohol and chloroform

DI-OXY-PHENYL-DI-CUMYL-METHANE $\text{CHPh}(\text{C}_6\text{H}_4\text{MePrOH})_2$. [146°] Formed from benzoic aldehyde, thymol, and H_2SO_4 (Russanoff, *B* 22, 1949). Thin tables (containing EtOH), v. sol. chloroform

Di-acetyl derivative [196°]. Crystals

OXY-DIPHENYLENE-ACETIC ACID

$\text{C}_6\text{H}_4\text{C}(\text{OH})\text{CO}_2\text{H}$. [162°] Formed by boiling phenanthraquinone with NaOHAq (Baeyer, *B* 10, 125, Friedlander, *B* 10, 584) Plates (containing 4aq), v al sol cold water, v sol alcohol Conc H_2SO_4 forms a blue solution on warming Chromic acid mixture yields diphenylene ketone NaOHAq at 160° splits it up into fluorene alcohol and CO_2 . HIAq and P at 140° reduce it to diphenylene acetic acid Bromine forms $\text{C}_{14}\text{H}_8\text{Br}_2\text{O}_2$ [225°], whence $\text{C}_{14}\text{H}_8\text{Br}_2\text{EtO}$ [151°]— CaA^1 , 2aq crystals *Ethyl ether EtA* [92°] Prisms.

DI-OXY-PHENYLENE-DIAMINE

$\text{C}_6\text{H}_4(\text{OH})(\text{NH}_2)_2$ Got by reducing the dioxim of di oxy-quinone with SnCl_4 and HCl (Nietzki & Schmidt, *B* 22, 1656) Yields on oxidation crystalline $\text{C}_6\text{H}_4(\text{OH})(\text{NH}_2)_2\text{B}^1\text{H}_2\text{SO}_4$

Tetra-acetyl derivative [225°] Needles.

Isomeride v DI-AMIDO-HYDROQUINONE

DI-OXY-DIPHENYLENE-DIHYDRAZINE

$\text{C}_6\text{H}_4(\text{OH})(\text{N}_2\text{H}_4)_2$ $\text{C}_6\text{H}_4(\text{OH})(\text{N}_2\text{H}_5)_2$ [140°] Formed by reducing the diazo-compound from di-oxy di amido-diphenyl with SnCl_4 (Kunze, *B* 21, 3333) Thin plates, forming with acetone a compound $\text{C}_{18}\text{H}_{12}\text{N}_4\text{O}_2$ [200°]

OXY-DIPHENYLENE KETONE $\text{C}_{12}\text{H}_8\text{O}_2$

$\text{CO}\text{C}(\text{OH})\text{C}_6\text{H}_4\text{OH}$ [96°] (G), [91°] (R) Formed by warming dry sodium salicylate with excess of POCl_3 and distilling (R Richter, *J pr* [2] 23, 294) It is also one of the products of the action of phenol on the sulphate of *o*-diazobenzoic acid (Griess, *B* 21, 981) Needles, insol water, v sol hot alcohol Its vapour passed over red-hot lime yields diphenylene ketone and di phenylene ketone oxide [82°] Distilled over red hot zinc dust it is reduced to diphenyl Gives a nitro-derivative [224°], a di nitro derivative [c 235°], and a bromo derivative [193°]

OXY-DIPHENYLENE-KETONE OXIDE

$\text{C}_{12}\text{H}_8\text{O}_2$ v $\text{C}_6\text{H}_4\text{C}(\text{O})\text{C}_6\text{H}_4(\text{OH})$ *Oxy-xanthone* [147°] Formed by heating resorcin with salicylic acid and ZnCl_2 (Michael, *Am* 5, 91). Formed also by heating salicylic acid with (β -resorcylic acid and Ac_2O (Graebe, *A* 254, 290) Yellow needles (from alcohol) Split up by fusion with potash into resorcin and salicylic acid. Gives diphenylene methane oxide [99°] when distilled with zinc dust Yields tri oxy-benzophenone [183°] when fused with NaOH — $\text{NaC}_6\text{H}_4\text{O}_2$ — $\text{Na}_2\text{C}_6\text{H}_4\text{O}_4$ (dried at 100°) Lemon-yellow needles

Acetyl derivative $\text{C}_{12}\text{H}_8\text{AcO}$. [168°].

Di-oxy-diphenylene ketone oxide v EUXANTHONE, vol II p 529 (β)-*Iso euxanthone* $\text{C}_6\text{H}_4(\text{OH})\text{C}(\text{O})\text{C}_6\text{H}_4(\text{OH})$, got from di nitro-

diphenylene ketone oxide [260°], yields a crystalline di-acetyl derivative [175°] (Graebe, *A* 254, 301) An isomeride $\text{C}_6\text{H}_4\text{C}(\text{O})\text{C}_6\text{H}_4(\text{OH})_2$ [247°] is obtained from phloroglucin and salicylic acid (von Kostanecki & Nessler, *B* 24, 1896).

A second isomeride with the formula $\text{C}_6\text{H}_4\text{C}(\text{O})\text{C}_6\text{H}_4(\text{OH})_2$ [1 6] [240°], got by heating tetra-oxy-benzophenone with water at 200° , yields a diacetyl derivative [161°] and dyes cotton

mordanted with alumina, yellow, and with iron, greyish black (Graebe, *B* 24, 969) Another isomeride $[1\frac{3}{4}]C_6H_4(OH) < \begin{smallmatrix} O \\ CO \end{smallmatrix} > C_6H_4(OH) [1\frac{5}{6}]$ [246° cor] made by heating resorcylic acid with Ac_2O yields a diacetyl derivative [c 127°] (G).

Tetra oxy-diphenylene ketone oxide *v* Anhydride of Hexa oxy-benzophenone

DI-OXY PHENYLENE-DI METHYL DIKETONE $C_6H_4(OH)_2(COCH_3)_2$ [180°] Obtained from di oxy acetophenone (10 g), $ZnCl_2$ (20 g), $HOAc$ (50 g), and $POCl_3$ (10 g), at 140° (Crepieux, *Bl* [3] 6, 152) Crystals (from hot water), yields with phenyl hydrazine the compound $C_6H_4(OH)_2(CMeN,HPH)_2$ [c 231°]

Tri oxy-phenylene di-methyl diketone $C_6H_3(OH)_3(COCH_3)_2$ [189°] Formed in like manner from tri oxy acetophenone (C) Yields a di phenyl di hydrazide [246°] and also an acetyl derivative [209°] which yields a di phenyl di hydrazide [265°]

DI OXY PHENYLENE DI PHENYL DIKETONE $C_6H_4(OH)_2(CO C_6H_5)_2$ *Di-benzoresorcin* [149°] Formed, together with dioxybenzophenone, by heating resorcin with $ZnCl_2$ and $ZnCl_2$ (Doebner a Stachmann, *B* 11, 2270, *A* 210, 259) Leaflets, insol water

Di acetyl derivative [150°] Needles

Di benzoyl derivative [151°] Needles

Di oxy phenylene di phenyl diketone

$C_6H_3(OH)_3(CO C_6H_5)_2$ *Dibenzohydroquinone* [207°] Formed from hydroquinone, $BzCl$, and $AlCl_3$ (D) Golden needles (from alcohol)

Di-benzoyl derivative [146°] Plates

TETRA OXY DIPHENYLENE DI PHENYL-DI THIO-DI UREA *Tetra methyl derivative*

$NHPh.CS.NH.C_6H_4(OMe)_2.C_6H_4(OMe)_2.NH.CS.NHPh$ [184°] Formed by warming the compound $C_6H_3(OMe)_3(NH)C_6H_4(OMe)_2(NH_2)$ with phenylthiocarbamide (Bossler, *B* 17, 2128) Flakes, sol hot alcohol

TETRA - OXY - DIPHENYLENEQUINONE *Tetra methyl ether v* CEREULIGNON

DI OXY DIPHENYLENE QUINOXALINE *Di ethyl ether*

$[6\frac{3}{4}]C_6H_4(OEt)_2 < \begin{smallmatrix} N & C & C_6H_4 & [1\frac{2}{3}] \\ N & C & C_6H_4 & [1\frac{2}{3}] \end{smallmatrix} > [260°]$

Formed by the action of phenanthraquinone on the hydrochloride of the diethyl ether of (a)-diamido hydroquinone (Nietzki a Reehberg, *B* 23, 1212) Yellowish needles, sl sol water

OXY DI-PHENYL ETHANE

C_6H_5O *ie* $C_6H_5CH_2CH_2C_6H_4(OH)$ or $CH_3CH(C_6H_5)C_6H_4(OH)$ [58°] Formed by allowing a mixture of styrene, phenol, H_2SO_4 , and $HOAc$ to stand (Koenigs, *B* 23, 3144) Crystals, sol dilute $NaOHaq$

Benzoyl derivative [83°]

Methyl derivative $CH_3PhCH_2C_6H_4OMe$ [61°] Formed from α -phenyl methoxy cinnamic acid, alcohol, and Na (Freund a Remse, *B* 23, 2865) Plates, sol alcohol

Sulphonic acid $C_{12}H_{12}(OH)(SO_3H)$ Formed by potash-fusion from *s*-di phenyl-ethane disulphonic acid (Kade, *B* 7, 239) Plates, sol hot water

Di-p oxy-s-di phenyl-ethane

$C_6H_4(OH)CH_2CH_2C_6H_4(OH)$ [189°] Formed from di-phenyl-ethane disulphonic acid by potash-fusion (Kade). Plates or needles, almost

insol cold water Does not give a dye stuff on oxidation (Heumann a Wiernik, *B* 20, 914)

Di-oxy-di-phenyl-ethane $CH_3CH(C_6H_4OH)_2$ *Ethylidene diphenol* [122°] (F), [125°] (Claus, *B* 19, 3004) Formed by the action of $SnCl_4$ or HCl on a mixture of phenol and paraldehyde (Fabiny, *B* 11, 283) Needles (containing $\frac{1}{2} C_6H_5$) Reduces ammoniacal $AgNO_3$ forming a mirror

Di benzoyl derivative [152°] Prisms

Di methyl derivative [140°] Formed from acetyl chloride, anisole, and $AlCl_3$ (Gattermann, *B* 22, 1129) Micaceous scales

Di ethyl derivative [142°] Scales

Isomerides *v* **DI OXY ETHYL-BENZENE**

Tri oxy-tri phenyl-ethane

$C_6H_4(OH)CH_2CH(C_6H_4OH)_2$ *Ethynyl-triphenol* Formed from $CH_3ClCHCl(OEt)$ and phenol (Wischneus, *A* 243, 153) Amorphous resin, v. sl sol ether Yields iso rosolic acid on oxidation with $FeCl_3$ Forms a tri acetyl derivative

Tetra oxy di-phenyl ethane

$C_6H_4(OH)CH(OH)CH(OH)C_6H_4(OH)$ *Anhydride* $C_{12}H_{10}O_2$ Two isomerides [68°] and [117°] are formed by the action of zinc dust and $HOAc$ on *o* oxy benzoic aldehyde (Tiemann, *B* 24, 3172, Harries, *B* 24, 3175)

Tetra oxy-tetra-phenyl ethane $C_{24}H_{18}(OH)_4$ [248°] Formed by fusing tetra phenyl-ethane tetra sulphonic acid with potash (Engler, *B* 11 930) Plates (from dilute alcohol)

Tetra oxy-tetra-phenyl ethane

$(HO C_6H_4)_4CHCH(C_6H_4OH)_2$ Formed from di oxy benzophenone (Baeyer, *A* 202, 133) Resin, yielding a crystalline tetra acetyl derivative

Hexa oxy tri phenyl ethane

$(HO)_2C_6H_3CH_2CH_2C(C_6H_4OH)_2$ Three amorphous isomerides are formed from di-chloro-di-ethyl oxide by the action of pyrocatechin, resorcin, and hydroquinone respectively (Wischneus, *A* 243, 181) They are v sol alcohol

OXY DI-PHENYL ETHANE CARBOXYLIC ACID $C_6H_4(OH)CH_2CHPhCO_2H$ [120°] Formed from phenyl-coumarin, dilute alcohol, and sodium amalgam (Sardo, *G* 13, 273) Small prisms— AgA' sl sol hot water

Oxy-di-phenyl ethane dicarboxylic acid $C_6H_4(CO_2H)CH_2CH(OH)C_6H_4CO_2H$ *Hydroxy-diphthalic acid* [c. 170°] Formed from its anhydride, which is a product of the action of zinc-dust and $HOAc$ on phthalic anhydride (Wischneus, *B* 17, 2181) Prisms— Ag_2A'' ; decomposes at 225° *in vacuo* (Hasselbach, *A* 243, 249)

Ethyl ether Et_2A'' Crystalline

Anhydride $C_{12}H_{10}O_4$ *Hydrodiphthalic acid* [198 5°] Prisms. Converted by KOy at 215° into $C_6H_4(CO_2H)CHCHClC_6H_4CO_2H$ — AgA' pp

Di-oxy-di-phenyl-ethane di-o-carboxylic acid $C_6H_4(CO_2H)CH(OH)CH(OH)C_6H_4CO_2H$ Formed by dissolving hydrodiphthalyl in $KOHaq$ (Hasselbach, *A* 243, 266) Hydrodiphthalyl is a product of the action of zinc-dust and $HOAc$ on diphthalyl The free acid is unstable, at once yielding the anhydride $C_{12}H_{10}O_4$, which on heating to 190° yields its second anhydride hydrodiphthalyl $C_{12}H_{10}O_4$ [250°]— AgA'' Pp

Di-oxy-di-phenyl-ethane di-p-carboxylic acid $C_6H_4(CO_2H)CH(OH)CH(OH)C_6H_4CO_2H$ Formed by reducing benzoic dicarboxylic acid

with sodium-amalgam (Oppenheimer, *B* 19, 1817). M sol water Infusible

o-OXY-PHENYL-ETHYL-AMIDO-ACETIC ACID *Ethyl derivative*

$C_6H_4(OEt)NEtCH_2CO_2H$ Formed from chloroacetic acid (1 mol), $C_6H_4(OEt)NH_2$ (2 mols), and alcohol (Vater, *J pr* [2] 29, 296) Oil — $C_6H_4(NO_2)HCl$ Very hygroscopic — *Ethylchloride* $C_6H_4(NO_2)HCl$ Formed from $C_6H_4(OEt)NEt$, and chloroacetic acid Oil — $(C_6H_4(NO_2)EtCl)_2PtCl_4$ yellow crystals

p-OXY-PHENYL ETHYLAMINE

$C_6H_4(OH)CH_2CH_2NH_2$ Formed by heating tyrosine at 270° (Schmitt & Nasse, *A* 133, 214) Solid. Yields *p* oxy benzoic acid on fusion with potash (Barth, *A* 152, 101) — $B'HCl$ Needles

Oxy di-phenyl ethylamine

$CHPh(OH)CHPh(NH_2)$ [161°] Formed by reducing the oxim of hydrobenzoin or the mono oxim of benzoin with sodium amalgam and alcohol (Goldschmidt & Polonowska, *B* 20, 492, 21, 488) Needles (from alcohol) — $B'HCl$ [210°] — $B_2H_4PtCl_4 \cdot 2aq$ — $B'HOAc$ [156°] Prisms

Di-acetyl derivative [159°]

p-Oxy-di-phenyl-ethyl-amine *Ethylether* $EtN(C_6H_4)(C_6H_4OEt)$ [319°] Formed from *p*-oxy-diphenylamine, EtI , and alcoholic potash Oil, with an odour between that of geraniums and violets (Philip & Calm, *B* 17, 2434)

β -OXY- β -PHENYL-ETHYL ISOAMYL MALONIC ACID $CHPh(OH)CH_2C(CO_2H)_2C_6H_5$ Formed by reducing phenacyl isoaeryl-malonic acid with sodium amalgam (Paal & T Hoffmann, *B* 23, 1503) Thick oil, almost insol water, yielding the lactone of γ oxy γ phenyl-heptonic acid on distillation

OXY PHENYL ETHYL CARBAMIC ANHY.

DRIDE $C_6H_4 \begin{smallmatrix} \diagup \\ \text{NH} \\ \diagdown \end{smallmatrix} \begin{smallmatrix} \text{NEt} \\ \diagup \\ \text{O} \end{smallmatrix} \diagdown \text{CO}$ [29°] Formed from $C_6H_4 \begin{smallmatrix} \diagup \\ \text{NH} \\ \diagdown \end{smallmatrix} \begin{smallmatrix} \text{O} \\ \diagup \\ \text{O} \end{smallmatrix} \diagdown \text{CO}$, alcoholic potash, and EtI (Bender, *B* 19, 2952) Not affected by HCl

o-OXY-PHENYL-ETHYL CARBAZIDE

Methyl derivative $C_6H_4N_2O_2$ *re* $C_6H_4(OMe)N_2HCO_2NH_2$ [110°] Formed from $C_6H_4(OMe)N_2H$, and phenyl cyanate (Reisenegger, *A* 221, 323) Needles (from Aq)

o-OXY-DI-PHENYL ETHYLENE $C_{12}H_{10}$ *re* $C_6H_4CHCHC_6H_4(OH)$ [136°] Formed in small quantity by heating salicylic aldehyde with phenylacetic acid and $NaOAc$ at 200° (Michael, *Am* 1, 315) Needles (from alcohol)

p-Oxy-di-phenyl-ethylene *Methyl derivative* $C_{12}H_{11}(OMe)$ [186°] Formed by heating the methyl derivative of *p*-coumaric acid (Ogliastro, *G* 9, 586) Plates, sl alcohol

Di-o-oxy-di-phenyl ethylene

$C_6H_4(OH)CHCHC_6H_4(OH)$ [95°] Formed from salicylic aldehyde, zinc-dust, and $HOAc$ (Harnies, *B* 24, 8178, Tiemann, *B* 24, 3176) Needles Its alkaline solutions show blue fluorescence Yields a di-benzoyl derivative [108°] which forms a dibromide [59°]

Di-p-oxy-di-phenyl-ethylene

$C_6H_4(OH)CHCHC_6H_4(OH)$ *Di-oxy-stilbene*. [280°]. Formed by boiling an alcoholic solution of $COCl_2CH(C_6H_4OH)_2$ with zinc-dust (Ester Meer, *B* 7, 1200, Elbs & Hoermann, *J pr* [2] 39, 498) Small crystals (from $HOAc$) Yields a di acetyl derivative [218°].

Di-oxy-di-phenyl ethylene *Di benzoyl derivative* $C_6H_4C(OBz)C(OBz)C_6H_5$ *Isobenzil* [159°] Formed by the action of sodium on a mixture of benzoic aldehyde and $BzCl$, or on $BzCl$ alone, in ether (Klinger & Schmitz, *B* 24, 1276) Decomposed by HCl into benzil and benzoic acid Alcoholic potash forms benzoil

Di p oxy di phenyl ethylene *Di methyl derivative* $CH_3C(C_6H_4OMe)_2$ [140°] Formed from anisole, $AcCl$, and $AlCl_3$ (Gattermann, *B* 22, 1132) Plates, v sl sol cold alcohol Yields $CO(C_6H_4OMe)_2$ on oxidation by CrO_3

Di-ethyl derivative $CH_3C(C_6H_4OEt)_2$ [142°] Formed from C_6H_4OEt , $AcCl$, and $AlCl_3$

Tetra oxy tetra-phenyl ethylene $C_{24}H_{20}O_4$ Formed by potash fusion from tetra phenyl-ethylene tetra sulphonic acid (Behr, *B* 5, 278) Plates (from $HOAc$) Not melted at 300° $FeCl_3$ in $HOAc$ forms green metallic crystals of $C_{24}H_{20}O_4 \cdot 3aq$, sl sol alcohol

DI OXY-DI-PHENYL ETHYLENE DIAMINE

Di ethyl derivative $C_6H_4(NH C_6H_4OEt)_2$ [98°] Formed by the action of ethylene bromide and Na_2CO_3 on the hydrochloride of $C_6H_4(OEt)NH_2$ (Bischoff, *B* 23, 1979) Plates (from ether alcohol)

OXY-DI PHENYL-ETHYLENE CARBOXYLIC ACID

$C_6H_4CH_2CH(OH)C_6H_4CO_2H$ [96°] Formed from deoxybenzoic o carboxylic acid by sodium amalgam (Gabriel & Michael, *B* 11, 1020, 18, 3480) Plates (from dilute alcohol) At 100° it forms an anhydride $C_{12}H_{10}O_2$ [60°]

The isomeric acid $C_6H_4CH(OH)CH_2C_6H_4CO_2H$, [127°], formed by reducing the isomeric deoxybenzoic o carboxylic acid, also yields an anhydride $C_{12}H_{10}O_2$ [90°]

p OXY-PHENYLETHYLENE-QUINOLINE $C_6NH_4CHCHC_6H_4OH$ [253°] Formed from (*Py* 1) quinoline (lepidine) by heating with *p* oxy benzoic aldehyde and $KHSO_4$ at 160° (Heyman & Koenigs, *B* 21, 1424) Crystals, sl sol dilute $NaOHaq$

p OXY PHENYL ETHYL KETONE

$C_6H_4(OH)CO C_6H_5$ *Propionyl phenol* [148°] *S* 034 at 15°, 33 at 100° Formed by heating phenol with propionic acid and $ZnCl_2$ (Goldzweig, *J pr* [2] 43, 86), and also by the action of propionyl chloride on phenol (Perkin, *C I* 55, 546) Needles or prisms, v e sol alcohol Potash fusion yields phenol and *p* oxy benzoic acid Yields a di bromo derivative [100°] and a nitro derivative [180°]

Methyl derivative $C_6H_4(OMe)CO C_6H_5$ [27°] (274°) Formed from anisole (10g), propionyl chloride (10g), and $AlCl_3$ (12g) in CS_2 (Gattermann, *B* 23, 1203) Yields an oxim [67°] crystallising from alcohol

Ethyl derivative [30°] Prisms Yields an oxim $C_{12}H_{11}(OEt)C(NOHEt)Et$ [97°]

Di-oxy-phenyl ethyl ketone

[134] $C_6H_4(OH)CO C_6H_5$ [95°] Formed from resorcin (1 pt), propionic acid (1 pt), and $ZnCl_2$ (Goldzweig) Needles Gives a red colour with $FeCl_3$. Yields a phenyl-hydrazide [115°]

Di-ethyl derivative $C_6H_4(OEt)_2CO C_6H_5$ [76°] Formed from propionyl chloride, $C_6H_4(OEt)_2$, and $AlCl_3$ (Gattermann). Yields an oxim [183°] crystallising in needles

Di-oxy-phenyl ethyl ketone

[521] $C_6H_4(OH)CO C_6H_5$ [92°] Formed from hydroquinone (1 pt), propionic acid (1 pt), and

ZnCl_2 (2 pts) at 190° (Goldzweig) Needles, giving a yellowish red colour with FeCl_3 Yields a phenyl hydrazide [100°]

OXY DI-PHENYL-ETHYL NITRITE

$\text{C}_6\text{H}_5\text{NO}_2$ 16 $\text{C}_6\text{H}_5(\text{OH})\text{CH}_2\text{ONO}$ [106°] Formed from di phenyl ethane (1 pt), HOAc (10 pts), and HNO_3 (1 pt of S G 15) in the cold (Anschutz a Romig, A 233, 327) Needles, oxidised by CrO_3 to benzophenone and di-phenyl-nyl nitrite [87°]

DI OXY DI PHENYL-DI-ETHYL PYRAZINE

INE DIHYDRIDE $\text{NPh} \begin{array}{c} \text{CHET CO} \\ \text{CO CHET} \end{array} \text{NPh}$ [260°] Formed, together with an isomeride [163°], by the action of Ac_2O on a phenylamido butyric acid (Nastvogel, B 22, 1795, 23, 2014) Needles (from alcohol)

OXY-PHENYL ETHYL PYRIMIDINE

$\text{Cet} \begin{array}{c} \text{N-CPh} \\ \text{C(OH)} \end{array} \text{CH}$ [238°] Formed from propionamidine hydrochloride, benzoyl aceto acetic ether, and dilute (10 p c) NaOHAq (Pin ner, B 22, 1621) Needles, v sl sol water

OXY TRI-PHENYL ETHYL PYRROLE

$\text{CPh} \begin{array}{c} \text{CH CPh} \\ \text{CO NEt} \end{array}$ [123°] and [129°] Formed from $\text{C}_6\text{H}_5\text{CBzCHBz}$ and alcoholic ethylamine (Japp a Klingemann, C J 57, 705) Formed also by heating the ethylamide of di phenyl-benzoyl propionic acid at 340° Trichino prisms [123°], $a b c = 912 \ 1 \ 952$, $\alpha = 78^\circ 48'$, $\beta = 89^\circ 10'$, $\gamma = 68^\circ 2'$ From dilute solutions it also crystallises in monoclinic prisms [129°], $a b c = 1 \ 690 \ 1 \ 1 \ 958$, $\beta = 86^\circ 54'$ (Tutton, C J 57, 731) Yields a biomo derivative $\text{C}_{14}\text{H}_{18}\text{BrNO}$ [142°] crystallising in monoclinic prisms

OXY PHENYL (Py 1) ETHYL QUINOLINE

$\text{C}_6\text{H}_5 \begin{array}{c} \text{C(CH}_3\text{CH}_2\text{C}_6\text{H}_5\text{OH)} \\ \text{N CH CH} \end{array}$ Formed by condensing *o* oxy benzoic aldehyde with (Py 1) methyl quinoline and reducing the product with HI in HOAc and P (Koenigs, B 21, 1428, 2167) The *o* compound melts at 181° , the *m* compound at 209° , and the *p* compound at 177° They yield the oxybenzoic acids on fusion with potash

β OXY PHENYL ETHYL SUCCINIC ACID

$\text{CO}_2\text{H CHPh CH(CO}_2\text{H)}\text{CH(OH)Me}$ Formed from phenyl acetosuccinic ether, alcohol and sodium amalgam (Weltner, B 18, 791) The free acid splits up at once, yielding the anhydride $\text{C}_{12}\text{H}_{12}\text{O}_4$ [$167 \ 6^\circ$] whence $\text{Ca}(\text{C}_{12}\text{H}_{11}\text{O}_4)_2 \cdot \text{AgC}_2\text{H}_3\text{O}_2$, flocculent pp

DI OXY DI PHENYL-ETHYL + THIO

UREA *Di methyl derivative* $\text{C}_6\text{H}_5(\text{OMe})\text{N}(\text{C}(\text{SEt})\text{NH C}_6\text{H}_5\text{OMe})$ [83°] Got from di amyl thio-urea and EtI (Foerster, B 21, 1863) Prisms $\text{—C}_{12}\text{H}_{18}\text{N}_2\text{SO}_2\text{HI}$ [163°] Rhombohedra $\text{—B'H}_2\text{P}(\text{Cl})_4$

o OXY PHENYL GLYCIDIC ACID

$\text{C}_6\text{H}_5(\text{OH}) \begin{array}{c} \text{O} \\ \text{CH CH} \end{array} \text{CO}_2\text{H}$ *Salicylglycidic acid* Formed by the action of conc aqueous NaOH NBz

upon benzoyl imido-cumarin $\text{C}_6\text{H} \begin{array}{c} \text{CH CH} \\ \text{CO CO} \end{array}$ (Flöchl a Wolfrum, B 18, 1185) Flat needles or prisms V sol alcohol and ether, sl sol cold water. By boiling with dilute H_2SO_4 , it is con-

verted into oxido-cumarin $\text{C}_6\text{H} \begin{array}{c} \text{O} \\ \text{CH CH} \\ \text{O CO} \end{array}$ [153°].

—CaA' , 6sq prisms

o OXY-PHENYL-GLYOXYLIC ACID

$\text{C}_6\text{H}_5(\text{OH})\text{CO CO}_2\text{H}$ [44°] Formed by adding acid to a solution of NaNO_2 and isatin in dilute NaOH , and heating the solution of the diazo compound thus obtained to 60° (Baeyer a Fritsch, B 17, 973) Yields *o* di oxy phenyl acetic acid on reduction with sodium amalgam

Di-oxy-phenyl-glyoxylic acid Methyl derivative [$4 \ 3 \ 1$] $\text{C}_6\text{H}_5(\text{OH})(\text{OMe})\text{CO CO}_2\text{H}$ [134°] A product of the oxidation of acetyl eugenol by KMnO_4 (Tiemann, B 24, 2878)

Prisms, v sol water and alcohol *Dimethyl derivative* [$4 \ 3 \ 1$] $\text{C}_6\text{H}_5(\text{OMe})_2\text{CO CO}_2\text{H}$ [139°] Formed, together with veratric acid by oxidation of the methyl derivative of eugenol or isoeugenol by KMnO_4 (Tiemann a Matsmoto, B 11, 141, Ciamician a Silber, B 23, 1165) Trimetric tables, melting at 100° when hydrated.

Methylene derivative

$\text{CH}_2 \begin{array}{c} \text{O} \\ \text{O} \end{array} \text{C}_6\text{H}_5\text{CO CO}_2\text{H}$ [149°] Formed by oxidising iso safrol with alkaline KMnO_4 (C a S) Light yellow needles, containing benzene of crystallisation (from benzene) $\text{—C}_6\text{H}_5\text{AgO}_2$

o OXY PHENYL HEXENOIC ACID

$\text{C}_6\text{H}_5(\text{OH})\text{C}_6\text{H}_5\text{CO}_2\text{H}$ *Anhydride* $\text{C}_{12}\text{H}_{16}\text{O}_2$ [54°] (301°) Formed by boiling sodium salicylic aldehyde with valeric anhydride (Perkin, A 147, 235) Prisms (from alcohol), insol cold Aq

An isomeric lactone, prepared by heating phenyl acetyl butyric acid, is oily (Erdmann, A 254, 182)

OXY PHENYL HEXOIC ACID

$\text{CHMe}(\text{OH})\text{CH}(\text{CH}_2\text{Ph})\text{CH}_2\text{CO}_2\text{H}$ [76°] Small prisms (containing aq) —CaA' , 6sq minute prisms *Anhydride* $\text{C}_{12}\text{H}_{16}\text{O}_2$ *Benzyl valero lactone* [86°] Formed by reducing benzyl-acetylpropionic acid with sodium amalgam (Erdmann, A 254, 182) Crystals (from CS_2)

o-OXY-PHENYL-HYDRAZINE

Methyl derivative $\text{C}_6\text{H}_5(\text{OMe})\text{NH NH}_2$ [43°] (240°) By reducing $\text{C}_6\text{H}_5(\text{OMe})\text{N}_2\text{SO}_2\text{Na}$ with Zn and glacial acetic acid there is formed the salt $\text{C}_6\text{H}_5(\text{OMe})\text{N}_2\text{H}_2\text{SO}_2\text{Na}$ aq, which is then warmed with HClAq (Reisenegger, A 221, 314) Needles (from ligroin) Yields B'HCl , $\text{B'H}_2\text{C}_2\text{O}_4$, and $\text{B'C}_6\text{H}_5\text{N}_2\text{O}$

Acetyl derivative $\text{C}_6\text{H}_5\text{N}_2\text{O}_2$ [135°]

Di-o-oxy-di-phenyl hydrazine Di ethyl derivative $\{\text{C}_6\text{H}_5(\text{OEt})\}_2\text{N}_2\text{H}_2$ [89°] Formed

by reducing the azo-compound $\text{N}_2(\text{C}_6\text{H}_5\text{OEt})_2$ by alcoholic ammonium sulphide (Schmitt a Möhlau, J pr [2] 18, 202) Colourless needles, in sol water, sol alcohol and ether The *m* iso meride forms colourless needles [85°] (Buchstab J pr [2] 29, 300)

Tetra-oxy-diphenyl-hydrazine Tetra

methyl derivative $\text{C}_6\text{H}_5(\text{OMe})_2\text{NH NH C}_6\text{H}_5(\text{OMe})_2$ *Hydrazodi-methylhydroquinone* Formed by reducing the di methyl derivative of nitro-hydro quinone in alkaline solution (Baessler, B 17, 2126) Converted by acids into the tetra methyl derivative of tetra oxy-diamido-diphenyl

p-OXY-PHENYL IMIDO DIACETIC ACID

Di ethyl derivative of the oxy anilide

$C_6H_5(OEt)N(CH_2CO_2H)CH_2CONHC_6H_5OEt$ [167°] Got from $C_6H_5(OEt)NHCH_2CO_2H$ by heating at 260° (Bischoff a Nastvogel, *B* 22, 1790) Crystalline, insol hot water

DI-*o*-OXY-DI-PHENYL-IMIDO-ACETO-NITRILE *Di-methyl derivative* $C_{11}H_{11}N_3O_2$, α . $NH(CH_2C_6H_5OMe)_2$ [123°] Formed by heating $C_6H_5(OMe)CH(OH)CN$ with alcoholic NH_3 at 70° (Voswinkel, *B* 15, 2025) Tables, insol water, sol alcohol

***o*-OXY-PHENYL- α -IMIDO-PROPIONIC ANHYDRIDE** *Di-benzoyl derivative* $\{C_6H_5(OH)C_6H_5(NBz)CO\}_2O$ Obtained by heating hippuric acid with salicylic aldehyde dissolved in acetic anhydride with addition of sodium acetate (Plöchl a Wolfrum, *B* 18, 1183) [160°] Crystalline solid By boiling its acetic acid solution with a few drops of aqueous HCl it is converted into the benzoyl derivative of imidocoumarin

DI-*o*-OXY-DI-PHENYL IMIDO THIAZOLE *Tetrahydride of the di-methyl derivative* $CH_2\langle\begin{smallmatrix} N(C_6H_5OMe) \\ CH_2S \end{smallmatrix}\rangle CN C_6H_5OMe$ [128°]

Formed by boiling di-anisyl thio urea with $C_6H_5Br_2$ (Foerster, *B* 21, 1864) Prisms On heating with CS_2 at 200° it yields the compound $CH_2\langle\begin{smallmatrix} N(C_6H_5O) \\ CH_2S \end{smallmatrix}\rangle CS$ [136°] — $B'H_2P_4Cl_4$ unstable when moist

***p*-OXY-PHENYL INDIAZOLE**

$C_6H_5\langle\begin{smallmatrix} N \\ OH \end{smallmatrix}\rangle N C_6H_5OH$ [195°] Formed by boiling its ethyl derivative with HIAq and a little P (Paal, *B* 24, 965) Groups of short prisms, ν sol HClAq, sl sol ether — $B'HI$ [200°] Large yellow plates

Ethyl derivative [118°] Formed by heating a conc alcoholic solution of the ethyl derivative of *o* nitro benzyl *p* amido phenol with tin and HCl Pearly plates (from alcohol), prisms (from benzene ligroin), or needles (from dilute HOAc) The tin double salt crystallises in yellowish plates [138°]

OXY-PHENYL-INDOLE $C_{11}H_{11}NO$ α .

$C_6H_5\langle\begin{smallmatrix} O(OH) \\ NH \end{smallmatrix}\rangle C_6Ph(?)$ [α 163°] Formed by heating the phenyl hydrazide of α oxy aceto phenone with $ZnCl_2$ at 160° (Laubmann, *A* 243, 246) Amorphous pp, ν sl sol ligroin

DI-OXY-DI-PHENYL KETONE ν DI-OXY-BENZOPHENONE

Di-oxy-di phenyl-diketone *Di-methyl derivative ν Anisal* Anisal forms a monooxime [180°] and two di oxims [195°] and [217°] The (α)-dioxim [217°] forms a diacetyl derivative [189°], while the (β) dioxim [195°] forms an isomeric di acetyl derivative [180°] (Suerlin, *B* 22, 877)

***o*-OXY-PHENYL MERCAPTAN**

$C_6H_5(SH)(OH)$ *Thiopyrocatechin* [6°] (217° cor) S G § 1287, 1189 Formed by reducing di oxy-di-phenyl disulphide with sodium-amalgam (Haitinger, *M* 4, 170) Formed also by the action of potassium xanthate on diazophenyl chloride, the oxyphenyl ethyl xanthate being subsequently saponified by sodium sulphide (Leuckart, *J pr* [2] 41, 192) Oil, volatile with steam, sl sol water Attacks the skin

p-Oxy-phenyl mercaptan $C_6H_5(SH)(OH)$ [80°]. (168°) Formed in like manner from

p-diazophenol (L) Crystalline mass, forming a bluish green solution in sulphuric acid — $Pb(S C_6H_5OH)_2$ bulky yellow pp Oxidised in alcoholic ammoniacal solution by air to di *p*-oxy-di phenyl disulphide [151°]

***Acetyl derivative* $C_6H_5(SH)(OAc)$ (280°)**

Di acetyl derivative [86°] Plates *Ethyl ethers* $C_6H_5(SH)(OEt)$, [41°], (277°) and $C_6H_5(SeEt)(OH)$, [41°], (287°) Crystalline **OXY-PHENYL METHACRYLIC ACID** is described as OXY-PHENYL CROTONIC ACID (*q* v)

OXY DI-PHENYL-METHANE ν BENZYL-PHENOL

o Oxy tri-phenyl-methane $CHPh_2C_6H_5OH$ [118°] Formed by passing air through a solution of diazoamido tri-phenol methane sulphate, and boiling the product in a current of CO_2 (O Fischer, *A* 241, 862) Sol alcohol and ether

Di p oxy di phenyl-methane $CH_2(C_6H_5OH)_2$ [158°] Formed by fusing di phenyl methane disulphonic acid with potash (Beck, *A* 194, 318) Plates or needles (from hot water), not volatile with steam Its sodium salts $C_{12}H_{11}NaO_2$ and $C_{15}H_{13}NaO_2$ form green solutions — BaA''

Di-acetyl derivative [70°] Prisms *Di-benzoyl derivative* [156°] Needles *Di methyl derivative* $CH_2(C_6H_5OMe)_2$ [52°] (M), [49°] (B) (335°) Formed from anisole, methylal, HOAc, and H_2SO_4 (Ter Meer, *B* 7, 1200) Small plates (from alcohol)

Di-ethyl derivative $CH_2(C_6H_5OEt)_2$ [89°] Scales, ν sol alcohol

Isomeride ν OXY DI PHENYL CARBINOL

Di p-oxy tri-phenyl methane $CHPh(C_6H_5OH)_2$ *Leucobenzaurn* [161°] Formed from di oxy tri phenyl carbinol, zinc dust and HCl (Doebner, *B* 12, 1462, *A* 217, 230) Formed also from di amido tri phenyl methane by the diazo reaction (O Fischer, *A* 206, 153), and from benzoic aldehyde, phenol, and H_2SO_4 (Russanoff, *B* 22, 1943) Yellowish needles (from dilute alcohol), sl sol hot water Absorbs atmospheric oxygen when heated above 160°, changing to di oxy tri phenyl carbinol Potash fusion gives di oxy benzophenone Yields a di nitro compound [134°]

Di-acetyl derivative [111°] Plates

Di-benzoyl derivative [130°]

Tri oxy tri phenyl methane $CH(C_6H_5OH)_3$, *Leucaurn* Formed by reducing tri oxy tri-phenyl carbinol (aurin) with zinc dust and NaOHaq (Dale a Schorlemmer, *A* 166, 286) Colourless prisms (from HOAc), sl sol water

Tri acetyl derivative [139°] (Zulkowsky, *A* 202, 197) Small needles

Tri-benzoyl derivative Crystals

Tetra-oxy di-phenyl methane $C_{12}H_9O_4$ Formed by fusing orcin with NaOH (Barth, *M* 8, 646) Needles, turning brown at 260°

Tetra-oxy tri phenyl methane $C_{15}H_{11}O_4$ α $CHPh(C_6H_5(OH))_2$ [171°] Formed by reducing the anhydride of the corresponding carbinol ('resorcinbenzein') with zinc dust and HCl (Doebner, *A* 217, 236) Colourless needles (from dilute alcohol) Reoxidised by alkaline K_2FeO_4 to the anhydride of the carbinol

Octo-oxy tri-phenyl-methane *Methyl derivative* $C_6H_5(OMe)(OH)CH(C_6H_5(OH))_2$ Formed from vanillin (1 pt), pyrogallol (1 87 pts), alcohol (20 pts), and conc HClAq (50 pts.) (Ettl,

M 3, 638). Colourless crystals, insol water, turned violet by HCl. An isomeride is formed from vanillin and phloroglucin.

p-OXY DI PHENYL-METHANE o-CARB- OXYLIC ACID *Methyl derivative* $C_6H_4(OMe)CH_2C_6H_4CO_2H$ [111°] Formed by reduction of $C_6H_4(OMe)COCH_2C_6H_4CO_2H$ (Nourisson, B 19, 2105) White needles, insol Aq.

Oxy-di phenyl-methane carboxylic acid $C_6H_4CH_2C_6H_4(OH)CO_2H$ [140°] Formed from sodium benzyl phenol and CO_2 (Paterno a Fiolet, J 1873, 440) Small needles (from water) —AgA' small needles (from water)

Oxy di phenyl-methane di carboxylic acid $OH(OH)(C_6H_4CO_2H)_2$ Formed, as K salt, by heating 'diphthalic acid' with KOHAq at 130° (Jullard, C C 1887, 1148) The free acid at once forms the lactone [205°] which yields a methyl ether [155°], an ethyl ether [99.5°], and an amide [160°]

Oxy-di phenyl methane tri carboxylic acid $C(OH)(C_6H_4CO_2H)_3$ Formed, as K salt, by heating 'diphthalic acid' $C_2O_3(C_6H_4CO_2H)_2$ with KOHAq at 110° (Jullard) The free acid changes to lactone

Ethyl ether [108°]

Oxy tri phenyl methane carboxylic acid $C_6H_4O_3$ *see* CHPh $(C_6H_4OH)_3$ $C_6H_4CO_2H$ [210°] Formed by reducing an alkaline solution of oxy tri phenyl carbinol carboxylic anhydride with powdered zinc (Pechmann, B 13, 1616) Needles

Di oxy-tri phenyl methane carboxylic acid $C_6H_4(OH)_2CHPh(C_6H_4CO_2H)$ [184°] Formed by reduction of di oxy tri phenyl carbinol carboxylic anhydride with zinc dust and NH_4Aq (Pechmann, B 14, 1859) Converted into anthranol by H_2SO_4

Di oxy tri phenyl methane carboxylic acid $C_6H_4(CO_2H)CH(C_6H_4OH)_2$ [225°] Formed by boiling phenol phthalein with zinc dust and NaOHAq (Baeyer, A 202, 80, 212, 350)

Di acetyl derivative [146°] Needles

Anhydride $C_6H_4O_3$ [214°-217°] Formed by reducing phenol phthalein anhydride with zinc dust and alcoholic NaOH. Small needles (from alcohol) Warm H_2SO_4 forms a green solution turning red

OXY-PHENYL PENTAMETHENYL HYDRIDE *see* PHENYL METHYL PURPURINE

OXY DI PHENYL DI METHYL DI AMIDO-BUTYRIC ACID *Lactone*

$O < \begin{smallmatrix} C(NMePh)_2 \\ CO \end{smallmatrix} CH_2$ *Phenyl methyl succinimide*

[157°] Got by heating succinic acid with phenyl-methyl-amine (Putti, G 16, 160) Needles or prisms, sl sol ether

p OXY-DI PHENYL-METHYL AMINE

Methyl derivative $MeN(C_6H_4)(C_6H_4OMe)$ [313°] Formed by methylation of p oxy di-phenyl amine (Philip a. Calm, B 17, 2433) Crl, with an odour like geranium or violets

OXY DI-PHENYL TETRAMETHYLENE-QUINONE (so-called) $C_{12}H_8O_2$ The following derivatives of this body are formed by the action of bases on the quinone $C_{12}H_8O_2$ got by oxidising di-phenyl butinene (Zincke a Breuer, B 18, 631, A 226, 88)

Imide $C_{12}H_8(OH) < \begin{smallmatrix} O \\ NH \end{smallmatrix}$ [174°]. Red plates, yielding a diacetyl derivative [201°] Aqueous SO_3 forms $C_{12}H_8O_3$ [187°]

Methylimide $C_{12}H_8(OH) < \begin{smallmatrix} O \\ NMe \end{smallmatrix}$ [170°].

Ethylimide [130°] Brown needles

Phenylimide $C_{12}H_8(OH)O(NPh)$ [158°]

o-Tolylimide [108°] Red needles

p-Tolylimide [155°] Violet needles

(a)-Naphthylimide [148°]

Isomeride $C_{12}H_8(OH)O_2$ [144°] Formed by warming the quinone $C_{12}H_8O_2$ with dilute NaOH (Z a B) Needles or prisms (from alcohol) Yields phthalic acid on oxidation. Aqueous SO_3 yields an oxyquinhydrone [155°]. HI reduces it to an oxyhydroquinone [73°] — $Ca(C_{12}H_8O_3)_2$ — BAA' — AgA' brownish-red pp

Acetyl derivative $C_{12}H_8O_4$ [111°]

Benzoyl derivative Monoclinic crystals

DI OXY-DI-PHENYL TRI-METHYLENE

THIO UREA *Di-methyl derivative* $CH_3CH_2NC_6H_4OMe$ [114°] Formed from $CH_3S-CNC_6H_4OMe$ [114°] Formed from di anisyl thio urea and trimethylene bromide (Foerster, B 21, 1872) Prisms (from alcohol)

DI - OXY - DI - PHENYL - DI - METHYLENE DIPYRAZOLE $C_8H_8N_4O_2$ *see*

$NPh < \begin{smallmatrix} N-CMe \\ CO CH \end{smallmatrix} CH_2CH_2CH_2CO > NPh$ Formed by splitting off alcohol (2 mols) from the phenylhydrazide of di acetyl-adipic ether (Perkin a Obrembsky, B 19, 2049, C J 57, 206) Small crystals, solid at 250° Sol acids and alkalis

OXY-PHENYL-METHYL-ETHYL-PYRAZ-

OLE $C_{12}H_{14}N_2O$ *see* $NPh < \begin{smallmatrix} CO \\ N \end{smallmatrix} CH_2 > C_2H_5$ [108°]

Formed by heating ethyl acetoacetic ether with phenyl hydrazine at 140° (Knorr a Blank, B 17, 2051) Crystals (containing aq) Oxidised by nitrous acid to $C_{12}H_{12}N_2O_3$ [160°]

OXY-PHENYL-METHYL-ETHYL-PYRIM-

IDINE $CPh < \begin{smallmatrix} N-CMe \\ N-C(OH) \end{smallmatrix} > C_2H_5$ [167°] Formed

from benzamidine hydrochloride, ethylacetoacetic ether, and dilute (10 p c) NaOHAq (Pinner, B 22, 1625) Prisms

Di-oxy-phenyl-methyl-ethyl-pyrimidine

Ethyl derivative

$C_6H_4(OEt) < \begin{smallmatrix} N-CMe \\ N-C(OH) \end{smallmatrix} > OEt$ [194°] Formed

from ethyl-acetoacetic ether and p-ethoxybenzamidine (Pinner, B 23, 2955) Needles (from alcohol), m sol hot alcohol

OXY-PHENYL-DIMETHYL-TETRAHYDRO-

PYRIDINE *see* BENZYLIDENE-DIACETONE-ALGAMINE

OXY PHENYLMETHYL KETONE *see* OXY-

ACETOPHENONE and BENZYL CARBINOL

DI OXY DI-PHENYL-DI-METHYL-PYRAZ-

INE DIHYDRIDE $NPh < \begin{smallmatrix} CHMe \\ CO \end{smallmatrix} > NPh$

[183°] Formed from phenyl- α -amido propionic acid and Ac_2O in the cold (Nastvogel, B 22, 1794, 23, 2012) Needles, insol water

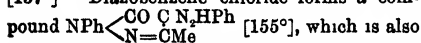
An isomeride [146°] accompanies the preceding body. On heating with KOHAq both compounds yield $C_{12}H_{12}N_4O_2$ [80°]

OXY-PHENYL-METHYL-PYRAZOLE

$NPh < \begin{smallmatrix} CO \\ N=O \end{smallmatrix} CH_2$ [127°] (287° at 265 mm.).

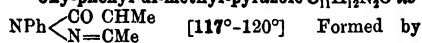
Preparation.—1 By heating acetoacetic ether with phenyl hydrazine (Knorr, B 17, 550, 2082; A 293, 187) —2 By the action of NH_4Aq and H_2S on (β)-phenyl-azo-crotonic ether (Bender, B 20, 2748)

Properties—Crystals, sol hot alcohol Yields bromo-, dibromo-, and di-chloro- derivatives Condenses with cinnamic aldehyde, acetone, and acetoacetic ether Zinc dust reduces it to phenyl methyl-pyrazole Excess of phenyl-hydrazine forms, on boiling, $C_{12}H_{11}N_2O_2$, whence $C_{12}H_{11}N_2O_2(SO_3H)_2$ Nitrous acid forms $C_{12}H_{11}N_2O_2$ [187°] Diazobenzene chloride forms a compound



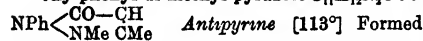
got by heating azobenzene acetoacetic amide with phenyl hydrazine at 130° (Leuckart & Holzapfel, *B* 22, 1406, Buchka, *B* 22, 2541) S_2Cl_2 forms a product which crystallises from alcohol as $C_{12}H_{11}N_2SO_2HClHOEt$ (Sprague, *C J* 59, 334) Alloxan forms a compound $C_{12}H_{11}N_2O_2$, whence dilute potash gives $C_{12}H_{11}N_2O_2$, converted by hot potash into $C_{11}H_9N_2O$, which on heating with alcohol gives rise to oxy phenyl methyl methyl ene-pyrazole [178°] (Pellizzari, *A* 255, 230)

Oxy-phenyl-di-methyl-pyrazole $C_{11}H_9N_2O_2$



heating methyl acetoacetic ether with phenyl-hydrazine at 140° (Knorr, *B* 17, 2050, *A* 238, 162) Formed also by heating oxy phenyl methyl pyrazol acetic acid Crystalline powder

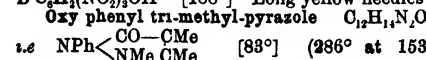
Oxy-phenyl-di-methyl-pyrazole $C_{11}H_9N_2O_2$



by methylation of oxy-phenyl methyl pyrazole (Knorr, *B* 17, 2032) and by heating acetoacetic ether with *s* phenyl methyl-hydrazine (Knorr, *A* 238, 203) Monoclinic crystals (from water), *v* sol water, alcohol, and ether Febrifuge $FeCl_3$ colours its solution deep red Nitrous acid gives a bluish green colour Sodium added to its alcoholic solution liberates methylamine K_2FeCy_4 and HCl give a yellow colour Chromic acid gives an orange pp (Gay & Fortuné, *Ph* 18, 1066) Gives the usual reactions for alkaloids ICl solution forms crystalline $C_{11}H_9IN_2O$ [160°] and $C_{11}H_9IN_2OICl$ [142°] (Dittmar, *B* 18, 1617) Bromine forms $C_{11}H_9Br_2N_2O$ [c 150°], whence water yields $C_{11}H_9BrN_2O$ [117°] HNO_2 forms $C_{11}H_9N_2O_2$, which explodes at 200° Alloxan yields $NMe \text{ CO} \begin{array}{c} \text{CO NH} \\ \text{CO NH} \end{array} \text{ CO}$, decomposing at 261°, whence boiling conc $HClAq$ forms $C_{12}H_{11}N_2O_2$ [238°] (Pellizzari, *G* 18, 340) Benzoin aldehyde in presence of HCl forms benzylidene-di antipyryne $C_{22}H_{21}N_2O_2$ [201°] Combines with chloral

Salts — $B'H_2PtCl_2 \cdot 2aq$ — $B'H_2FeCy_4$ — $B'O_2H_2(NO_2)_2 \cdot OH$ [188°] Long yellow needles

Oxy phenyl tri-methyl-pyrazole $C_{12}H_{11}N_2O$



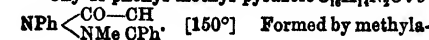
Formed by heating antipyryne with MeI and $MeOH$ (Knorr) Crystals, *v* sol water and alcohol Yields a picrate [94°]

Oxy-phenyl-tri-methyl-pyrazole $C_{12}H_{11}N_2O$



Formed from oxy phenyl methyl pyrazole $NaOMe$, and MeI (Knorr, *A* 238, 165), and also by heating di-methyl-acetoacetic acid with phenyl hydrazine Insol water, *v* sol alcohol

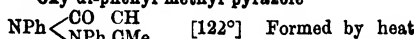
Oxy-di-phenyl-methyl-pyrazole $C_{11}H_9N_2O_2$



tion of oxy di phenyl pyrazole (Knorr & Klotz, *B* 20, 2549) Needles (from ether), *sl* sol boiling water Bromine in chloroform forms a di bromide, whence water liberates $C_{12}H_{11}Br_2N_2O$ [110°-120°]

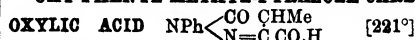
Salts — $B'HCl$ needles — $B'H_2FeCy_4$ — $B'O_2H_2(NO_2)_2 \cdot OH$ [170°] Yellow prisms

Oxy di-phenyl methyl pyrazole



ing hydrazobenzene with acetoacetic ether at 120°, the yield being 60 p.c. of the theoretical (Muller, *B* 19, 1771, Perger, *B* 19, 2140, *M* 7, 191) Crystals (containing $\frac{1}{2}aq$), *v* sol alcohol

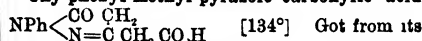
OXY PHENYL METHYL-PYRAZOLE CARB



Made from its ether, which is got by heating oxal propionic ether with phenyl hydrazine at 120° (Arnold, *A* 246, 331) Plates (from alcohol)

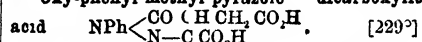
Ethyl ether EtA' [149°]

Oxy phenyl methyl pyrazole carboxylic acid



ether [85°], which is formed by heating acetone dicarboxylic ether with phenyl hydrazine (Pechmann, *A* 261, 171) Prisms, *sol* alcohol

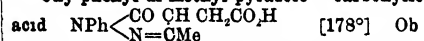
Oxy-phenyl methyl pyrazole dicarboxylic



Formed by saponifying its ether Needles (containing aq), *sl* sol cold water Gives a dark violet colour with $FeCl_3$

Ethyl ether EtA'' [130°] Formed by heating oxalsuccinic ether with phenyl hydrazine at 170° (Wischneus, *B* 22, 888) Needles

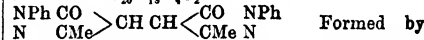
Oxy phenyl di methyl pyrazole carboxylic



tained from its ether [138°], which is got by heating acetyl succinic ether with phenyl hydrazine at 150° (Knorr) Needles (from water)

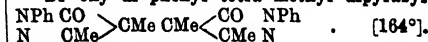
DI OXY-DI PHENYL DI-METHYL DI-

PYRAZOL $C_{22}H_{21}N_2O_2$



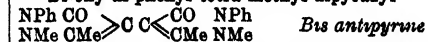
the action of phenyl hydrazine (3 mols) on acetoacetic ether (2 mols) Formed also by the action of phenyl hydrazine on di acetyl succinic ether, on isocarbopyrrolitic ether and on thioacetoacetic ether (Knorr, *B* 17, 2044, 2058, 22, 160, Buchka & Sprague, *B* 22, 2554) Formed also from *s* phenyl azo crotonic ether, alcoholic NH_3 and H_2S (Bender, *B* 20, 2749) Yellowish powder Decomposes on heating without melting Insol neutral solvents, *sol* alkalis Gives the pyrazole blue reaction Yields a bromo derivative [217°]

Di-oxy-di-phenyl-tetra methyl dipyrazyl



Formed by oxidation of oxy phenyl di methyl pyrazole with nitrous acid (Knorr & Blank, *B* 17, 2050) Long prisms (from $HOAc$), insol water and alkalis, *sol* conc H_2SO_4

Di-oxy di-phenyl-tetra methyl dipyrazyl



[245°] Formed by methylation of the preceding body (Knorr, *A* 238, 210) Crystals (from $MeOH$),

nearly insol water —B'H₂Cl₂ 2aq —B'H₂PtCl₆ • orange red prisms Picrate [c 161°]

Oxy di phenyl di-methyl-dipyrasyl sulphide $\text{NPh} \begin{smallmatrix} \text{CO} \\ \text{N} \end{smallmatrix} \begin{smallmatrix} \text{CH} \\ \text{CMe} \end{smallmatrix} \text{CH}_2 \text{S}$ Formed by adding phenyl hydrazine (2 mols) dissolved in HOAc to etho aceto acetic ether (1 mol) in the cold (Buchka a Sprague, B 23, 849) Formed also from oxy phenyl methyl pyrazole and SCl₂ in chloroform Small needles, decomposing at 183° without melting, sl sol alcohol, sol alkalis

OXY PHENYL METHYL PYRIDAZINE

$\text{NPh} \begin{smallmatrix} \text{CO CH} \\ \text{N} \text{CMe} \end{smallmatrix} \text{CH}$ [82°] Formed, together with its chloro- derivative [137°], from the anhydride of the phenyl hydrazide of levulinic acid and PCl₅ at 160° (Ach, A 253, 47) Translucent crystals, v sol alcohol Na added to its alcoholic solutions forms a compound C₁₂H₁₂N₄? [200°], coloured violet by H₂SO₄ and CrO₃ —B'HCl

Oxy phenyl-methyl-pyridasine

$\text{NPh} \begin{smallmatrix} \text{CO CO} \\ \text{N} \text{CMe} \end{smallmatrix} \text{CH}_2$ [196°] Formed from the chloro derivative, mentioned above, by treatment with alcoholic potash, the resulting $\text{NPh} \begin{smallmatrix} \text{CO C(OEt)} \\ \text{N} \text{CMe} \end{smallmatrix} \text{CH}$ [146°] being heated with HClAq at 130° White needles, sl sol hot water

OXY PHENYL-DI-METHYL PYRIDINE

$\text{C}_{13}\text{H}_{13}\text{NO} \pm e$ $\text{NPh} \begin{smallmatrix} \text{CMe CH} \\ \text{CMe CH} \end{smallmatrix} \text{CO}$ *Phenyl lutidone* [197°] (above 360°) Formed by the action of aniline on methyl dehydracetate (Perkin, B 18, 682, C J 51, 498), and also by heating its carboxylic acids (Conrad a Guthzeit, B 20, 161, 22, 85) Needles or prisms (containing aq) v sol hot water and alcohol —B'H₂PtCl₆ yellow needles (from water) —B'O₂H₂(NO)₂OH [95°] Transparent plates

Oxy phenyl di methyl pyridine

$\text{NMe} \begin{smallmatrix} \text{CO CH} \\ \text{CMe CH} \end{smallmatrix} \text{CPh}$ *Methyl phenyl p pco lostyrl* [112°] Formed by heating 'methyl-carbo phenyl lutidylum dehydride' with conc HClAq at 180° (Hantzsch, B 17, 2915) Prisms B'HCl 2aq —B'H₂PtCl₆ 3aq crystalline powder *m-Oxy (Py 3) phenyl (Py 1 5) di methyl-pyridine* C₁₁H₁₁NO ± e C₆H₅(OH) C₂NH₂(CH₃)₂ *Oxy phenyl lutidine* [191°] From the corresponding amido compound by the diazo-reaction White crystalline solid (Lepetit, B 20, 2399) —B'HCl 2aq —B'H₂PtCl₆ [c 200°]

OXY PHENYL DI METHYL PYRIDINE CARBOXYLIC ACID C₁₁H₁₁NO₃ ± e

$\text{NPh} \begin{smallmatrix} \text{CMe CH} \\ \text{CMe C(CO}_2\text{H)} \end{smallmatrix} \text{CO}$ [257°] Formed by heating the dicarboxylic acid at 227° (Conrad a Guthzeit, B 20, 161, 947, 22, 84) Formed also from β phenyl amido crotonic ether and NaOEt at 90° (Knorr, B 20, 1399) Satiny needles Melts at 267° when quickly heated —BaA', 4aq needles, v sol water

Methyl ether MeA' [152°] Formed from methyl dehydracetate and aniline (Perkin, jun, B 18, 682, C J 51, 498) Needles

Oxy-phenyl-di methyl-pyridine dicarboxylic acid $\text{NPh} \begin{smallmatrix} \text{CMe C(CO}_2\text{H)} \\ \text{CMe C(CO}_2\text{H)} \end{smallmatrix} \text{CO}$ Got by electrolysis of its ether Prisms, v sol hot alcohol —Ba(HA'')₂ aq

Ethyl ether EtA' [171°] Formed by

boiling an acetic acid solution of aniline and the product of the action of COCl₂ on cupric acetoacetic ether (Conrad a Guthzeit, B 19, 25, 20, 161) Crystalline solid, v sol alcohol —B'H₂PtCl₆ [120°] Orange yellow pp

m Oxy-(Py 3) phenyl-(Py 1 5)-di-methyl-pyridine-(Py 2 4) di carboxylic acid C₆H₅(OH) C₂N(CH₃)₂(CO₂H)₂ *m Oxy phenyl-lutidine di carboxylic acid*

Di-ethyl-ether A'Et, [174°] From the corresponding amido compound by the diazo reaction (Lepetit, B 20, 2398) White needles

OXY PHENYL METHYL PYRIMIDINE

$\text{CPh} \begin{smallmatrix} \text{N CMe} \\ \text{N CO} \end{smallmatrix} \text{CH}_2$ [216°] Formed by mixing a solution of benzamidine hydrochloride (1 mol) with acetoacetic ether (1 mol) and NaOH (1 mol) Formed also by the action of benzimidio ether CPh(NH)OEt on acetoacetic ether at 100° and of benzamidine on acetyl malonic ether (Pinner, B 18, 759, 2851, 22, 1624, 2617, 23, 3820) Needles, v sol alcohol, sl sol water —B'H₂PtCl₆ 2aq —B'C₆H₅(NO)₂OH [189°] —B'H₂Cr₂O₅ 5aq [177°] Orange red prisms

Acetyl derivative C₁₁H₁₁N₂O₃ [41°]

Ethyl derivative C₁₁H₁₁(OEt)₂N₂ [31°]

(300°) Formed from chloro phenyl methyl-pyrimidine and NaOEt Thick prisms —B'HCl 2aq [86°] —B'HCl [149°] —B'H₂PtCl₆ [197°] —B'HI 4aq [144°] Yellow prisms

Oxy phenyl methyl pyrimidine

$\text{CMe} \begin{smallmatrix} \text{N CPh} \\ \text{N CO} \end{smallmatrix} \text{CH}$ [238°] Formed from acetamidine hydrochloride, benzoyl acetic ether, and dilute (10 p c) NaOHAq (Pinner, B 22, 1618) Needles, m sol hot alcohol

Oxy phenyl-di-methyl-pyrimidine

$\text{CPh} \begin{smallmatrix} \text{N CMe} \\ \text{N CO} \end{smallmatrix} \text{CHMe}$ [203°] Formed from benzamidine hydrochloride, methyl acetoacetic ether, and NaOHAq (Pinner, B 22, 1624) Needles

Oxy di-phenyl-methyl-pyrimidine

$\text{CPh} \begin{smallmatrix} \text{N CPh} \\ \text{N CO} \end{smallmatrix} \text{CHMe}$ [250°] S 195 at 20° Formed from benzamidine and a benzoyl propionic ether (E v Meyer, J pr [2] 39, 197, 40, 303, Schwarze, J pr [2] 42, 12) Minute needles Oxidised by KMnO₄ to oxy di phenyl pyrimidine carboxylic acid [236°] Yields a methyl derivative [132°]

Oxy-phenyl-methyl-pyrimidine *Ethyl derivative* C₆H₅(OEt) C $\begin{smallmatrix} \text{N CMe} \\ \text{N CO} \end{smallmatrix} \text{CH}_2$ [146°]

Formed by mixing o ethoxy benzamidine hydrochloride with NaOH and acetoacetic ether (Pinner, B 23, 2953) Short columns, v sol alcohol

An isomeride [204°] is formed from p ethoxy benzamidine and acetoacetic ether or acetyl-malonic ether

Oxy-phenyl-di methyl-pyrimidine *Ethyl derivative* C₆H₅(OEt) C $\begin{smallmatrix} \text{N CMe} \\ \text{N CO} \end{smallmatrix} \text{CHMe}$ [216°]. Formed from p ethoxy benzamidine and methyl acetoacetic ether (P) Small prisms

Reference — Bromo - oxy - phenyl - methyl - PYRIMIDINE

OXY-PHENYL-DI-METHYL PYRIMIDINE CARBOXYLIC ACID

$\text{CPh} \begin{smallmatrix} \text{N CMe} \\ \text{N CO} \end{smallmatrix} \text{CH CH}_2 \text{CO}_2\text{H}$ [259°] Got by saponifying its ether Needles, sl sol water.

Ethyl ether EtA' [178°] Formed from acetyl succinic ether and benzamidine (Pinner B 22, 2618) Crystals, sl sol water

OXY-PHENYL-METHYL-PYRIMIDYLME THYL METHYL KETONE

$\text{CPh} \begin{smallmatrix} \text{N CMe} \\ \text{CO} \end{smallmatrix} > \text{C CH}_2 \text{CO CH}_2$, [225°] Formed from di-acetyl-succinic ether and benzamidine (Pinner, B 22, 2622) Needles, sl sol alcohol

OXY-PHENYL-METHYL PYRIMIDYL PRO- PIONIC ACID

$\text{CPh} \begin{smallmatrix} \text{N CMe} \\ \text{CO} \end{smallmatrix} > \text{CH CH}_2 \text{CH}_2 \text{CO}_2\text{H}$ [215°] Formed by saponification of its ether [145°], which is got by the action of benzamidine on acetyl glutaric ether (Pinner, B 22, 2620) Powder, sl sol water

OXY-TRI PHENYL-METHYL PYRROLE

$\text{C}_6\text{H}_5\text{NO}$ *se* $\text{NMe} < \begin{smallmatrix} \text{CO CPh}_2 \\ \text{CPh CH} \end{smallmatrix}$ *Tri phenyl-methyl pyrrolone* [139°] and [143°] Formed from $\text{C}_6\text{H}_5\text{CBz CHBz}$ and alcoholic methylamine (Japp a Klingemann, B 22, 2884, C J 57, 698) Colourless rhombohedra [139°], $a c = 1.248$ Occurs also in triclinic forms [143°], $a b c = 906.1 \ 870$, $\alpha = 79^\circ 52'$, $\beta = 86^\circ 3'$, $\gamma = 70^\circ 28'$ (Tutton, C J 57, 724) Oxidised by chromic acid to $\text{C}_6\text{H}_5\text{NO}$, [167°] Bromine in chloroform yields $\text{C}_6\text{H}_5\text{BrNO}$ [150°]

Dihydrate $\text{NMe} < \begin{smallmatrix} \text{CO} - \text{CPh}_2 \\ \text{CHPh CH}_2 \end{smallmatrix}$ *Tri-*

phenylmethylpyrrolidone [153.5°] Mol w (by Raoult's method) 335 (calc 327) Formed by reduction of the preceding body by sodium and boiling amyl alcohol Monoclinic needles, $a b c = 1.655 \ 1.1043$, $\beta = 88^\circ 29'$

Oxy-tetra-phenyl-methyl-pyrrole $\text{C}_6\text{H}_5\text{NO}$ *se* $\text{NMe} < \begin{smallmatrix} \text{CO CPh}_2 \\ \text{CPh CPh} \end{smallmatrix}$ [161°] Formed by heating $\text{C}_6\text{H}_5\text{CBz CHBz C}_6\text{H}_5$ with alcoholic methylamine at 200°, and also from benzoyl-tri-phenyl propiomethylamine and KOH (Klingemann a Laycock, C J 59, 146, B 24, 518) Small yellow monoclinic plates (from boiling alcohol), v sol CS₂.

OXY-PHENYL METHYL PYRROLE CARB- OXYLIC ACID *Ethyl ether*

$\text{NH} < \begin{smallmatrix} \text{CO CPh} \\ \text{CMe CCO}_2\text{Et} \end{smallmatrix}$ [128°] Formed by the action of alcoholic ammonia on phenyl acetyl-succinic ether (Emery, A 260, 152) Needles

Oxy-phenyl-methyl pyrrole dihydride carb- oxylic acid $\text{NPh} < \begin{smallmatrix} \text{CO CH}_2 \\ \text{CMe(CO}_2\text{H)} \end{smallmatrix} > \text{CH}_2$, [188°]

Obtained from its nitrile, which is made from levulic ether, HCy, and alcoholic aniline (Kühling, B 22, 2364) Hard prisms, sl sol water *Amide* [127°] Needles

Nitrile Oil Converted by NH_3 and H_2S into $\text{NPh} < \begin{smallmatrix} \text{CO} - \text{CH}_2 \\ \text{CMe(CS NH)} \end{smallmatrix} > \text{CH}_2$, [193°] Hy- droxylamine forms the amidoxim [120°]

OXY - PHENYL - METHYL - QUINAZOLINE

C_6H_5 $\begin{smallmatrix} \text{CO NMe} \\ \text{N} - \text{CPh} \end{smallmatrix}$ [181°] Formed by the action of KOH and MeI at 120° on o-benzoyl amido-benzamide or on oxy phenyl quinazoline (Körner, J pr [2] 86, 162) Shining lamellae, sol alcohol

Isomeride C_6H_5 $\begin{smallmatrix} \text{CO NPh} \\ \text{N CMe} \end{smallmatrix}$ [147°]. Formed

by heating the acetyl derivative of o amido benzoic anilide (K) Four-sided prisms

OXY - PHENYL - METHYL - QUINOLINE $\text{CMe CH C C(OH) CH} \begin{smallmatrix} \text{CH} \\ \text{CH} - \text{CH C N} - \text{CPh} \end{smallmatrix}$ [291°] Formed by heating its carboxylic acid at 250° (Just, B 19, 1544) Plates (from alcohol)

Isomeride v FLAVENOL

OXY - PHENYL - METHYL - QUINOLINE CARBOXYLIC ACID $\text{CMe CH C C(OH) C CO}_2\text{H} \begin{smallmatrix} \text{CH} \\ \text{CH CH C N} - \text{CPh} \end{smallmatrix}$

Formed by saponifying its ether [236°], which is got by heating to 160° the product of the action of $\text{C}_6\text{H}_5\text{MeN CClPh}$ on sodium malonic ether (Just, B 19, 1542) V sl sol water and alcohol

OXY - PHENYL - METHYL - (Py) - QUINO-

$\text{N} - \text{NPh} \begin{smallmatrix} \text{C} \\ \text{C} \\ \text{N} - \text{C(OH)} \end{smallmatrix} \begin{smallmatrix} \text{C} \\ \text{C} \\ \text{CMe} \end{smallmatrix}$ **PYRAZOL** C_6H_5 *Inner - anhy-*

dride of o amido di phenyl methyl pyrazol-carboxylic acid [261°] Formed by reduction of o nitro di phenyl methyl pyrazol carboxylic acid with SnCl_2 (Knorr a Jödicke, B 18, 2262) Fine needles V sol alcohol, chloroform, and acetic acid, insol water, alkalis, and dilute acids It can be distilled undecomposed It is a very stable body By sodium and alcohol it is reduced to a hydro compound which dissolves in strong H_2SO_4 with a deep green colour

OXY-PHENYL-METHYL-QUINOXALINE

$\text{C}_6\text{H}_5\text{Me}$ $\begin{smallmatrix} \text{N C(OH)} \\ \text{N CPh} \end{smallmatrix}$ [198°] Formed from (1,3,4)-tolylene diamine and phenyl glyoxylic acid (Hunsberg, A 237, 352) Pale yellow needles

γ OXY-γ-PHENYL-S DI-METHYL-SUCCINIC ACID $\text{CHPh(OH) CH(CO}_2\text{H) CHMe CO}_2\text{H}$ *Phenyl homo stamnic acid* The free acid is unstable, but the salts CaA'' 3aq, BaA'' 2aq, and $\text{Ag}_2\text{A}''$ are got by boiling its lactone with bases (Fittig a Penfield, B 216, 119, 20, 3179)

Lactone [177°] *Phenyl homoparaconic acid* Formed from benzoic aldehyde, sodium pyro tartrate, and Ac_2O by heating for 10 hours at 130° Plates (from water) Yields AgA' Forms phenyl-butylene (177°) on distillation Conc HBr at 0° forms $\text{CHPhBr CH(CO}_2\text{H) CHMe CO}_2\text{H}$ [149°] which on warming with water becomes $\text{CHPh CH CHMe CO}_2\text{H}$ The lactone [177°] is accompanied by an isomeride [124.5°] The lactone [177°] yields on distillation methyl-naphthol [89°], while the isomeride yields an isomeric methyl naphthol [92°] Both methyl-naphthols yield (8) methyl naphthalene on distillation with zinc dust

o - Oxy - phenyl - di - methyl succinic acid $\text{C}_6\text{H}_5\text{(OH) CH}_2\text{CH(CO}_2\text{H) CHMe(CO}_2\text{H)}$ [145° 150°] Formed by reducing coumaryl propionic acid in alkaline solution with sodium amalgam (Fittig a Brown, A 255, 288) Crystals, v sol water - BaA'' - CaA'' - $\text{Ag}_2\text{A}''$ curdy pp

Oxy-phenyl-methyl succinic acid is described as **OXY BENZYL-SUCCINIC ACID**

OXY-DI PHENYL-METHYL ψ-THIO UREA.

Methyl derivative $\text{C}_6\text{H}_5\text{(OMe) NH C(SMe) NPh}$ [80°] Formed by the action of MeI on phenyl amyl thio urea (Foerster, B 21, 1870) Long white needles

Di oxy-di phenyl-methyl- ψ thio-urea Di-methyl derivative

$C_6H_5(OMe)NH C(SMe)NC_6H_5OMe$ [87°] Made by heating di-anisyl thio urea with MeI at 100° (Foerster, B 21, 1860) Prisms, v e sol hot alcohol Yields on distillation MeSH and a substance [56°] which on boiling with dilute HCl gives di anisyl urea [184°]— $C_{12}H_{15}N_2SO_2HCl$ white plates— $B^*H_2PtCl_6$ prisms

OXY-PHENYL-NAPHTHYL-ACETIC ACID

$CPh(C_6H_5)(OH)CO_2H$ The salt NaA' 2aq [145°] is formed from sodium phenyl (α) naphthyl ketone, water, and CO_2 (Beckmann, B 22, 915)

α DI OXY PHENYL NAPHTHYL KETONE
 $C_6H_5(OH)CO C_{10}H_7(OH)$ [c 106°] Formed by heating (α) phenonaphthoxanthone with alcoholic potash at 200° (Phomina, A 257, 93) Yields a methyl derivative [66°], an acetyl derivative [137°] and an acetoxim [196°]

The $\alpha\beta$ isomeride [169°] yields K_2A'' and a methyl derivative [68°], an ethyl derivative [c 141°], an acetyl derivative [108°], an acetoxim [188°], and a phenyl hydrazide [198°]

DI OXY DI PHENYL OCTANE

$CPhPr(OH)CPhPr(OH)$ [64°] Formed by the action of sodium amalgam on a solution of phenyl propyl ketone in dilute alcohol (Schmidt & Fieberg, B 6, 499) Needles (from acetone)

Di oxy di phenyl-octane $C_6H_5CMe(C_6H_5OH)_2$ [83 5°] Formed from methyl hexyl ketone, phenol, and HCl (Dianin, J R 1888, 534) Needles Yields a di benzoyl derivative [114°]

OXY PHENYL OXANTHRANOL $C_{12}H_9O_3$

$C_6H_5 \begin{smallmatrix} CO \\ \diagup \quad \diagdown \\ C(OH)(C_6H_5OH) \end{smallmatrix} C_6H_5$ [194°]

Formed by oxidising oxy phenol anthranol with $KMnO_4$ (Pechmann B 13, 1617) Yellowish crystals Yields anthraquinone on oxidation

Acetyl derivative $C_{12}H_9O_4$ Needles

Di oxy-phenyl oxanthranol v PHENOL

PHTHALIDIN**OXY PHENYL-ISO OXAZOLE**

$CPh \begin{smallmatrix} CH \\ \diagup \quad \diagdown \\ N \end{smallmatrix} CO$ [147°] Formed from benzoyl acetic ether and hydroxylamine in acid, neutral, or ammoniacal solution (Claisen & Zedel, B 24, 141, Hantzsch, B 24, 502) Needles, sl sol cold alcohol $HClAq$ at 120° yields the oxim of acetophenone Ammonia forms $CPh \begin{smallmatrix} CH \\ \diagup \quad \diagdown \\ N \end{smallmatrix} CO NH_2$ [168°] Yields a

nitroso derivative $CPh \begin{smallmatrix} C(OH) \\ \diagup \quad \diagdown \\ N \end{smallmatrix} CO$ [143°]

DI OXY DI PHENYL OXIDE $C_{12}H_9O_3$ v e $O(C_6H_5OH)_2$? Formed from phenol and CrO_2Cl_2 (Eliard, B [2] 28, 276) Amorphous powder Yields quinone on oxidation, v RESORCIN

DI OXY DI PHENYL OXINDOLE $C_{20}H_{13}NO_2$

$se NH \begin{smallmatrix} C_6H_5 \\ \diagup \quad \diagdown \\ CO \end{smallmatrix} C(C_6H_5OH)_2$ Phenolsatin [220°] Formed by adding H_2SO_4 to a mixture of isatin and phenol (Baeyer & Lazarus, B 18, 2641) White needles, insol water, sol alkalis Acetyl derivative $C_{20}H_{13}O_3NaO$ [185°] Colourless needles (from HOAc)

Di methyl derivative $C_{20}H_{15}Me_2NO_2$ [65°] Made from anisole, isatin, and H_2SO_4

p OXY-PHENYL PENTADECYL KETONE

Methyl derivative $C_{15}H_{31}CO C_6H_5OMe$ [70 5°] (280° at 15 mm.) Formed from anisole,

palmityl chloride, and $AlCl_3$ (Krafft, B 21, 2269)

Ethyl derivative [69°] (289 at 15 mm) Formed in like manner from phenetole Plates

Di oxy-phenyl pentadecyl ketone Di-methyl derivative [63 5°] (290° at 15 mm) Formed in like manner from the dimethyl derivative of resorcin (K) Plates

DI OXY PHENYL PENTANE

$CHPh(OH)CH(OH)Pr$ [82°] (287°) Formed by the action of alcoholic potash on a mixture of benzoic and isobutyric aldehydes (Swoboda, M 11, 390) White crystals, v sol ether

Di-acetyl derivative $C_{11}H_{14}O_4$ [55°] (297°)

Di oxy phenyl pentane

$CHPh(OH)CH_2CH_2CH_2CH_2OH$ [54°] Formed by reducing benzoyl butyl alcohol with sodium amalgam The glycol is extracted with ether (Kipping & Perkin, jun, C J 57, 312). Needles (from benzene), v sol ether

Di-oxy-di-phenyl-pentane $CEt_2(C_6H_5OH)_2$ [200°] Formed from di ethyl ketone, phenol, and HCl (Dianin, J R 1888, 534) On fusion with NaOH it yields $C_6H_5C_6H_5OH$ [76 5°] (253°)

OXY PHENYL PENTENOIC ACID v OXY-PHENYL ANGELIC ACID**DI OXY DI PHENYL-PENTINOIC ACID**

Di methyl derivative $C_6H_5(OMe)CH CH CO(CO_2H)CH C_6H_5OMe$ [160°] Formed from p methoxy benzoic aldehyde, sodium succinate, and Ac_2O (Fittig & Politis, A 255, 299) Long yellow needles (from HOAc)— BaA' , 2aq silky plates— CaA' , 3aq— AgA light yellow pp

OXY-PHENYLPHENYLENE-ANTHRA-QUINONE DIHYDRIDE $C_{20}H_{12}O_2$ se

$CO \begin{smallmatrix} C_6H(OH) \\ \diagup \quad \diagdown \\ C_6H_2 \end{smallmatrix} CO \begin{smallmatrix} C_6H_5 \\ \diagup \quad \diagdown \\ C_6H_2 \end{smallmatrix} Ph$ [266°] Formed by

fusing truxone with potash (Liebermann & Bergmann, B 23, 321) Yellow needles, sl sol alcohol Yields an acetyl derivative $C_{20}H_{12}AcO_2$ [180°], v sol HOAc

OXY-DI PHENYL PHOSPHINE Phenyl derivative $P(C_6H_5)_2OPh$ (265°-270° at 62 mm) SG $\frac{2}{3}$ 114 VD 1002 (obs) Got by heating PPh_2Cl with phenol (Michaelis & La Coste, B 18, 2109) Oil Readily absorbs oxygen from the air, becoming $(C_6H_5)_2PO OPh$ Sulphur yields $(C_6H_5)_2PS OPh$ [124°] Selenium forms $Ph_2PSe(OPh)$ [115°] Yields a crystalline methyl iodide [c 136°] and a benzylo chloride $Ph_2P(OPh)CH Cl$ [232°-236°]

TETRA-OXY-DI PHENYL-PHTHALIDE**Anhydride v FLUORESCIN****OXY PHENYL-PHTHALIMIDE v. AMIDOPHENOL****DI-OXY-DI-PHENYL-PROPANE**

$OMe_2(C_6H_5OH)_2$ [154° cor] Got from acetone, phenol, and $BzCl$ (Dianin, J R 1888, 534) Flat needles Yields, on soda-fusion, p isopropyl-phenol [61°]

OXY PHENYL-PROPIOLIC ACID v Cog-**MARLIC ACID** **α -OXY β -PHENYL-PROPIONIC ACID**

$C_6H_5CH_2CH(OH)CO_2H$ [98°] Formed from phenyl acetic aldehyde, H_2Cy , and HCl (Erlenmeyer, B 18, 308, A 219, 179) Got also by reducing phenyl glyoxylic acid with sodium-amalgam (Plöchl B 16, 2823) Prisms (from

water) Yields formic acid and phenyl-acetic acid on heating —BaA', aq globular aggregates
Nitrile [58°] Needles (from benzene)

***β* Oxy *β* phenyl propionic acid**

$C_6H_5CH(OH)CH_2CO_2H$ [93°] Formed by the action of sodium amalgam on the acid $C_6H_5CH(OH)CHClCO_2H$ (Glaser, *A* 147, 86) and by boiling *β*-bromo *β*-phenyl propionic acid with water (Fittig a Binder, *A* 195, 138) Got also by reducing benzoyl acetic ether with sodium amalgam (Perkin, *C J* 47, 254) Prisms, v sol cold water Yields cinnamic acid on heating with baryta water (Kast, *A* 206, 26) or with dilute H_2SO_4 at 100° (Erlenmeyer) —KA' —BaA', 1:1 aq —ZnA', 1:1 aq —AgA' needles

***Acetyl derivative* [100 5°] Crystals**

Yields cinnamic acid when heated with Ac_2O at 120° (Slocum, *A* 227, 59)

***β* Oxy-*α*-phenyl-propionic acid**

$CH_3(OH)CHPhCO_2H$ *Tropic acid* Mol w 166 [118°] S 2 at 145°

Formation —1 By decomposing atropine or hyoscyamine with fuming $HClAq$ or with baryta water (Lossen, *A* 138, 230, Kraut, *A* 148, 238, Ladenburg, *B* 13, 607) —2 From atropic acid by union with $HOCl$ followed by reduction with zinc dust, iron filings, and conc $KOHaq$ (Ladenburg a Rüdheimer, *B* 13, 379) —3 From atropic acid by heating it with fuming hydrogen chloride at 100° and digesting the resulting acid ($CHClCHPhCO_2H$) with aqueous Na_2CO_3 at 120° (Ladenburg, *A* 217, 114) —4 From acetophenone cyanhydrin by treatment with HCl , the resulting chloro phenyl propionic acid being heated with sodium carbonate (Spiegel, *B* 14, 237, 1852, Merling, *A* 209, 5)

Properties —Needles or tables, v sol water Yields phenyl acetic acid on fusion with potash Chromic mixture oxidises it to benzoic acid

Salts —CaA', 4aq —AgA' crystals

***Reference* —CHLORO TROPIC ACID**

***α* Oxy *α* phenyl propionic acid**

$C_6H_5CMe(OH)CO_2H$ *Atrolactic acid* [94°]

Formation —1 From atropic acid by successive treatment with HBr and aqueous Na_2CO_3 (Fittig a Wurster, *A* 195, 145, 206, 24) —2 By oxidation of *α*-phenyl propionic acid with alkaline $KMnO_4$ (Ladenburg a Rüdheimer, *B* 13, 373, *A* 217, 107) —3 By reduction of the dibromo derivative obtained by the action of H_2SO_4 on a mixture of dibromo pyruvic acid and benzene (Böttinger, *B* 14, 1238) —4 By dissolving the cyanhydrin of acetophenone in saturated $HClAq$ (Spiegel, *B* 14, 1353, Tiemann, *B* 14, 1980)

Properties —Trimetric needles or tables (containing $\frac{1}{2}aq$), $abc = 72.1$ 57 Becomes anhydrous below 85° Gives rise to atropic acid CH_3CPhCO_2H on distillation with HCl

Salts —CaA', 8aq —BaA', 2aq —ZnA', 2aq. small crystals, v sl sol cold water

Ethyl derivative $CMePh(OEt)CO_2H$ '*Ethyl tropic acid*' [c 62°] Formed from acetophenone by treatment with PCl_5 , acting upon the resulting $PhCCl_2Me$ with KCy and dilute alcohol, and saponifying the product with baryta Small prisms, sol hot water

Nitrile Formed from acetophenone, KCy and HCl (S) Oil

***p*-Oxy-*α*-phenyl-propionic acid**

$C_6H_4(OH)CHMeCO_2H$ *Phloretic acid*. [129°].

Formed from phloretin (7g) and KOH (55 cc. of S G 1.2) by boiling for 3 hours (Hlasiwetz, *J* 1855, 700, Schiff, *A* 172, 357) Formed also from amido phenyl propionic acid by diazo reaction (Trimus, *A* 227, 268) Yellowish needles V sol hot water, sol alcohol and ether, insol CS_2 Yields *p* oxy benzoic acid on fusion with potash On heating with phloroglucin at 180° it yields the crystalline phloroglucide $C_{23}H_{16}O_{11}$ Heating with $POCl_3$ forms crystalline triphloretide $C_{12}H_8O_6$ Phloretic acid gives a green colour with $FeCl_3$, possibly due to phloroglucin
 Salts —BaA', 2aq —CuA', (at 120°)

Ethyl ether EtA' (above 265°)

Isoamyl ether $C_5H_{11}A'$ (above 290°)

Methyl derivative

$C_6H_4(OMe)CHMeCO_2H$ [108°] S 11 at 25°. Got by methylation (Körner a Corbetta, *B* 7, 1732) Yields BaA', 2aq and MeA' (38°) (278°)

Ethyl derivative [106 5°] Scales

Amide $C_6H_4NO_2$ [110°-115°] Prisms

Isophloretic acid, made by boiling isophloretin with $KOHaq$ (Roehleider, *Z* 1868, 711) is probably identical with phloretic acid

***o* Oxy *β* phenyl propionic acid**

$C_6H_4(OH)CH_2CH_2CO_2H$ *Mellitic acid Hy drocoumaric acid* [83°] S 5 at 18°, 109 at 40° Occurs, partly combined with coumarin, in yellow mellot (*Mellotus officinalis*) (Zwenger a Bodenbender, *A* 126, 257, *Suppl* 5, 100)

Formed by reducing coumarin with sodium-amalgam (Zwenger, *A* *Suppl* 8, 32, Dyson, *C J* 51, 70, Hochstetter, *A* 226, 355, Tiemann, *B* 10, 286) Trimetric crystals Its ammoniacal solution turns blue in air Yields salicylic acid when fused with potash Conc $HBrAq$ converts it into the anhydride

Salts —KA', 2aq —BaA', 3aq —CaA', —CaA', 2aq —MgA', 4aq —CuA', 2aq —PbA', —ZnA', aq —AgA' bulky pp

Ethyl ether EtA' [34°] Prisms

Methyl derivative

$C_6H_4(OMe)CH_2CH_2CO_2H$ [92°] Crystals (from alcohol) (Perkin, *C J* 39, 416)

Ethyl derivative

$C_6H_4(OEt)CH_2CH_2CO_2H$ [80°] Needles (from dilute alcohol) Yields the salts $Ba(C_6H_4O_2)_2$ and CaA_2 , 2aq (Fittig a Ebert, *A* 216, 153)

Anhydride $C_6H_4O_2$ *Hydrocoumarin* [25°] (272°) Got by distilling the acid Tables, sl

sol hot water Its oxim $C_6H_4<\frac{C=O}{O}>C(NOH)$ is only (Tiemann, *B* 19, 1664)

Amide $C_6H_4NO_2$ [70°] Needles

***m* Oxy-*β* phenyl propionic acid**

$C_6H_4(OH)CH_2CH_2CO_2H$ [111°] Formed by reduction of *m*-coumaric acid (Tiemann a Ludwig, *B* 15, 2050) Long needles, insol ligroin

Ethyl derivative [c 51°] Needles

***p* Oxy-*β* phenyl propionic acid**

$[4\frac{1}{2}]C_6H_4(OH)CH_2CH_2CO_2H$ *Hydropara coumaric acid* [129°] Occurs in very slight quantity in human urine (Baumann, *H* 4, 307) Formed by reducing paracoumaric acid with sodium amalgam (Hlasiwetz a Malin, *A* 142, 358) Formed also from *p* nitro-cinnamic ether by reduction followed by the diazo reaction (Stöhr, *A* 225, 57, cf. Buchanan a Glaser, *Z* [2] 5, 198) Occurs in putrid meat (Salkowski, *B* 13, 190). Formed also by putrefactive fermentation of tyrosine (Baumann, *B* 12, 1450;

Acetyl derivative $C_{11}H_{11}AcON_2Oaq$ [100°] Prisms

Nitrosamine $C_{10}H_{11}(NO)N_2O$ [109°].

DI-*p*-OXY-DI-PHENYL PROPYLENE *Di-methyl derivative* $CH_3CH(O_2CH_3)_2$ [101°] Formed from anisole, $AlCl_3$, and propionyl chloride (Gattermann, *B* 22, 1130) Plates Yields $CO(O_2CH_3)_2$ on oxidation The homologous $CH_3CH(O_2CH_2OEt)_2$ crystallises in needles [77°]

OXY-PHENYL ISOPROPYL KETONE

Ethyl derivative $C_6H_5(OEt)COCHMe$, [41°] Made from isobutyl chloride, C_6H_5OEt , and $AlCl_3$ (Gattermann, *B* 23, 1206) Tables (from ether) Yields an oxim [111°] crystallising in needles

Oxy phenyl propyl ketone $C_6H_5COCH_2CH_2OH$ [91°] (c 280°) Formed from phenol and butyl chloride (Perkin, *C J* 55, 546) Pearly plates

DI-OXY-PHENYL-ISOPROPYL-PYRIM

IDINE $CH<\begin{smallmatrix} OPh-N \\ O(OH)N \end{smallmatrix}>CMe(OH)$ [198°]

Formed from oxy isobutyramidine hydrochloride, benzoyl acetic ether, and $NaOH_{aq}$ (Pinner, *B* 22, 2626) Prisms (from alcohol or benzene)

OXY-TRI-PHENYL-PROPYL-PYRROLE

$C_{25}H_{23}NO$ *is* $NPr<\begin{smallmatrix} CO CPh_2 \\ CPh CH \end{smallmatrix}>$ [105°] Formed from $C_6H_5CBzCHBz$ and alcoholic propylamine at 100° (Japp a Klingemann, *C J* 57, 706) Monoclinic prisms [95°-98°] or triclinic prisms [105°], $a:b:c = 694:1:765$

DI-*o*-OXY-DI-PHENYL PROPYL- ψ -THIO

UREA *Di methyl derivative* $C(SPr)(NH C_6H_5OMe)N C_6H_5OMe$ [58°] Made from di-anisyl thio urea and propyl chloride (Foerster, *B* 21, 1864) Plates, insol water

OXY-DI-PHENYL PYRAZINE *Tetra*

hydride $NPh<\begin{smallmatrix} CH_2CH_2 \\ CO CO \end{smallmatrix}>NPh$ [148°] Formed from di phenyl ethylene diamine, chloro acetic acid, and $NaOAc$ at 170° (Bischoff a Nastvogel, *B* 22, 1788, 23, 2026) Crystalline insol Aq

Di-oxy di phenyl pyrazine *Dihydrate*

$NPh<\begin{smallmatrix} CH_2CH_2 \\ CO CO \end{smallmatrix}>NPh$ *Di phenyl as diacetyl perazine* [258°] Got by oxidising the above with CrO_3 and $HOAc$, and also by heating di phenyl ethylene diamine with oxalic acid at 200° (B a N) Plates, v sl sol alcohol and ether Nitrous acid forms $C_{16}H_{16}N_4O_2$ [290°]

Di-oxy di phenyl-pyrazine *Dihydrate*

$NPh<\begin{smallmatrix} CO CH_2 \\ CH_2 CO \end{smallmatrix}>NPh$ *Diphenyl as diacetyl perazine* *Di oxy-di phenyl piazine dihydrate* [268°] Made from aniline and chloro acetic acid (Hausdörfer, *B* 22, 1797, cf Meyer, *B* 10, 1967, Abenius, *J pr* [2] 40, 431) Needles Alcoholic potash forms phenyl amido acetic acid [127°] on boiling, and in the cold produces $NHPhCH_2CO NPhCH_2CO_2H$ [130°]

Di-oxy-phenyl-pyrazine *Dihydrate*

$NPh<\begin{smallmatrix} CH_2CO \\ CH_2CO \end{smallmatrix}>NH$ [158°] Made from aniline and chloro-acetic amide (B) Prisms

Di-oxy-di-phenyl-pyrazine *Dihydrate*

$NPh<\begin{smallmatrix} CH_2CO \\ CH_2CO \end{smallmatrix}>NPh$ *Diphenyl-as diacetylperazine* [152°] Formed by heating the mono-amide of phenyl-imido diacetic acid with Ac_2O

at 160° (Bischoff, *B* 23, 1990) Needles Nitrous acid yields $C_{16}H_{16}N_4O_2$ [c 95°] crystallising in golden plates

Di-*p*-oxy-di-phenyl-pyrazine *Di-methyl derivative of the hexahydrate*

$C_6H_5(OMe)N<\begin{smallmatrix} CH_2CH_2 \\ CH_2CH_2 \end{smallmatrix}>N C_6H_5OMe$ [238°].

Formed from *p* anisidine, ethylene-bromide, and $NaOAc$ (Bischoff, *B* 22, 1782) Tables

Di ethyl derivative of the hexahydrate [223°] Gives a nitroso derivative [c 80°]

Tri oxy di phenyl-pyrazine *Di ethyl derivative of the tetrahydrate*

$C_6H_5(OEt)N<\begin{smallmatrix} CH_2CH_2 \\ CH_2CO \end{smallmatrix}>N C_6H_5OEt$ [162°].

Formed from $C_6H_5(NH C_6H_5OEt)_2$ by heating with chloro acetic acid and fused $NaOAc$ at 150° (Bischoff, *B* 23, 2030) Crystals, sl sol ether

Tetra oxy di phenyl-pyrazine *Di ethyl derivative of the dihydrate*

$C_6H_5(OEt)N<\begin{smallmatrix} CH_2CO \\ CO CH_2 \end{smallmatrix}>N C_6H_5OEt$ [265°]

Formed by heating *p* ethoxy phenyl amido-acetic acid at 260° (B a N) Needles

OXY PHENYL PYRAZOLE $C_6H_5(OH)N_2$ [153°] Got from bromo oxy phenyl pyrazole by reduction with sodium amalgam (Fischer a Knoevenagel, *A* 239, 201) Needles Sol conc HCl_{aq} and alkalis The Na salt forms needles

Oxy-phenyl pyrazole tetrahydrate $C_6H_5N_2O$ *is* $NPh<\begin{smallmatrix} NH CH_2 \\ CH_2CH(OH) \end{smallmatrix}>?$ [104°] Formed by the

action of phenyl hydrazine on epichlorhydrin in ethereal solution at 15° (Gerhard, *B* 24, 352) Pearly prisms (from ether), v sol warm water Yields aniline and phenyl pyrazole on boiling with phenyl hydrazine hydrochloride in benzene solution Forms an acetyl derivative—Salt $B'H_2PtCl_2$ 2aq [156°] Yellowish red prisms

Oxy-di-phenyl-pyrazole $NPh<\begin{smallmatrix} NH CO \\ CPh CH \end{smallmatrix}>NPh$ *Di phenylpyrazolone* [251°] Formed by heating cinnamoyl phenyl hydrazine (Knorr, *B* 20, 1108) Needles (from alcohol), exhibiting green fluorescence

Oxy di phenyl pyrazole $NPh<\begin{smallmatrix} N CPh \\ CO CH_2 \end{smallmatrix}>$

[137°] Formed from benzoyl acetic ether and phenyl hydrazine (Knorr a Klotz, *B* 20, 2546) Yields, with nitrous acid, the red compound

$NPh<\begin{smallmatrix} N CPh \\ CO CNOH \end{smallmatrix}>$ [200°] Diazobenzene chloride forms $C_6HPhN_2O(N_2Ph)$ [171°] Phenyl hydrazine converts it into di oxy tetra phenyl dipyrazyl melting above 300° — $B'HCl$ — $B'H_2SO_4$ Powder Decomposes at 237°

Oxy di phenyl-pyrazole $NPh<\begin{smallmatrix} N CH \\ CO CHPh \end{smallmatrix}>$

[196°] Formed from a phenyl glyoxylic acid and phenyl hydrazine (Wishnecus, *B* 20, 2932) Plates (from $EtOH$), sl alkalis, sl sol ether

Di-oxy-phenyl metapyrazole $C_6H_5N_2O$, probably $C_6H_5OH-N=\begin{smallmatrix} O(OH)N \\ C(OH) \end{smallmatrix}$ [182°] Ob-

tained by boiling phenyl uramido aceto nitrile $C_6H_5CH(OH)NHCO NH_2$ (from benzaldehyde-cyanhydrin and urea) with dilute HCl Colourless dimetric plates Dissolves readily in alkalis (Pinner a Lifschutz, *B* 20, 2355)

OXY PHENYL PYRAZOLE CARBOXYLIC

ACID $C_{10}H_9N_3O_3$, $\text{v sol NPh} \left\langle \begin{smallmatrix} \text{N} & \text{C}(\text{CO}_2\text{H}) \\ & \text{CO OH} \end{smallmatrix} \right\rangle$ [263°]

Formed by the action of HCl or NaOH on the phenyl hydrazide of oxalacetic ether (Buchner, *B* 23, 2931) Colourless needles, v sl sol cold water FeCl₃ colours its solution dark blue
• *Ethyl ether* [182°] Formed by heating the above phenyl hydrazide by itself (Wishnecus, *A* 246, 306)

OXYPHENYL PYRIDINE $C_{11}H_9NO$ Formed by heating aniline chelidonate (Lerch, *M* 5, 407) Crystallises from water in needles (containing aq)

Oxy-di-phenyl-pyridine $C_{17}H_{11}NO$ $\text{v sol NH} \left\langle \begin{smallmatrix} \text{CPh CH} \\ \text{CPh CH} \end{smallmatrix} \right\rangle \text{CO}$ [267°] Formed by heat

ing dehydro benzoyl acetic acid with alcoholic NH₃ at 160° (Feist, *B* 23, 3736) Plates

DI OXY-PHENYL PYRIDINE CARBOXYLIC ACID $C_{12}H_9(OH)_2NPh \text{ CO}_2\text{H}$ Made by boiling a solution of comenic acid with aniline (Mennel, *J pr* [2] 32, 177) Tetrahedra (containing aq), m sol water Coloured violet by FeCl₃

OXY PHENYL PYRIMIDINE $C_{10}H_7N_2O$ $\text{v sol CPh} \left\langle \begin{smallmatrix} \text{N} & \text{CH} \\ \text{N} & \text{C}(\text{OH}) \end{smallmatrix} \right\rangle \text{CH}$ [199°] Made by heating its carboxylic acid (Pinner, *B* 22, 2616) Small prisms, sl sol hot Aq, v sol alkalis and acids

Oxy di phenyl pyrimidine $C_{16}H_{11}N_2O$ $\text{v sol CPh} \left\langle \begin{smallmatrix} \text{N} & \text{CPh} \\ \text{N} & \text{C}(\text{OH}) \end{smallmatrix} \right\rangle \text{CH}$ [284°] Formed from benzamididehydrochloride, benzoyl acetic ether, alcohol, and dilute (10 p c) NaOH (Pinner, *B* 22, 1626) Formed also by heating amido di phenyl pyrimidine with HClAq at 180° (Schwarze, *J pr* [2] 42, 15) Needles, v sl sol hot alcohol

Oxy tri phenyl pyrimidine $C_{22}H_{15}N_2O$ [340°] Formed by heating the corresponding amido compound with HClAq at 170° (Wache, *J pr* [2] 39, 255) Needles

Di oxy di phenyl pyrimidine *Ethyl derivative* $C_8H_7(OEt)_2 \text{C} \left\langle \begin{smallmatrix} \text{N} & \text{CPh} \\ & \text{N} & \text{C}(\text{OH}) \end{smallmatrix} \right\rangle \text{CH}$ [274°] Formed from *p* ethoxy benzamidine and benzoyl-acetic ether (Pinner, *B* 23, 2956)

OXY PHENYL PYRIMIDINE CARBOXYLIC ACID $C_{10}H_7 \left\langle \begin{smallmatrix} \text{N} & \text{C}(\text{CO}_2\text{H}) \\ & \text{N} & \text{C}(\text{OH}) \end{smallmatrix} \right\rangle \text{CH}$ [247°] Formed from benzamidine hydrochloride, oxalacetic ether, and dilute (10 p c) NaOH (Pinner, *B* 22, 1628, 2616) Granules, sl sol water—BaA"—CaA'. Prisms—Amide prisms, sl sol water—Benzamidide $C_{10}H_7N_2O_3$ [263°]

Oxy di phenyl-pyrimidine carboxylic acid $C_{16}H_{11} \left\langle \begin{smallmatrix} \text{N} & \text{CPh} \\ & \text{N} & \text{C}(\text{OH}) \end{smallmatrix} \right\rangle \text{CO}_2\text{H}$ [236°] Made by oxidising oxy di phenyl methyl pyrimidine with KMnO₄ (E von Meyer, *J pr* [2] 40, 303) Yellowish prisms (from alcohol) Yields oxy-di phenyl pyrimidine [284°] on heating

Di oxy phenyl pyrimidine carboxylic acid *Ethyl derivative* $C_8H_7(OEt)_2 \text{C} \left\langle \begin{smallmatrix} \text{N} & \text{C}(\text{CO}_2\text{H}) \\ & \text{N} & \text{C}(\text{OH}) \end{smallmatrix} \right\rangle \text{CH}$ [246°] Made from *p*-ethoxy benzamidine and oxalacetic ether (Pinner, *B* 23, 2956) Needles, sl sol hot alcohol—Salt $C_8H_7(OEt)_2 \text{CN}_2\text{H}_4\text{A}'$ [280°]

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OXY-TRI PHENYL-PYRROLE $C_{21}H_{15}NO$ s.e.

$\text{NH} \left\langle \begin{smallmatrix} \text{CO CPh}_2 \\ \text{CPh CH} \end{smallmatrix} \right\rangle$ Mol w 311 [221°] Formed by boiling C_6H_5 , CBz CHBz for a long time with alcoholic ammonia (Japp a Klingemann, *B* 22, 2884, *C J* 57, 682) Triclinic crystals, $\alpha b c = 779 \ 1 \ 512$, $\alpha = 78^\circ 56'$, $\beta = 73^\circ 23'$, $\gamma = 87^\circ 32'$ (Tutton, *C J* 57, 720) Sl sol hot alcohol

Dihydrate $\text{NH} \left\langle \begin{smallmatrix} \text{CO} & \text{CPh}_2 \\ & \text{CHPh CH}_2 \end{smallmatrix} \right\rangle$ [201°]

Formed by reduction with sodium and amyl alcohol Ac_2O at 180° yields an acetyl derivative $\text{NaO} \left\langle \begin{smallmatrix} \text{CO} & \text{CPh}_2 \\ & \text{CHPh CH}_2 \end{smallmatrix} \right\rangle$ [105°].

o Oxy-tri-phenyl-pyrrole

$C_6H_5(OH) \text{N} \left\langle \begin{smallmatrix} \text{CPh CH} \\ \text{CPh CH} \end{smallmatrix} \right\rangle$ [176°] Made by heating its carboxylic acid with lime (Paal a Braikoff, *B* 22, 3094) Yellowish needles, v sol alkalis

Oxy-tetra phenyl-pyrrole

$C_{22}H_{15}NO$ $\text{v sol NH} \left\langle \begin{smallmatrix} \text{CO CPh}_2 \\ \text{CPh CH} \end{smallmatrix} \right\rangle$ [207°]. Formed by heating CPhBz CPhBz with alcoholic NH₃ at 200° (Klingemann a Laycock, *C J* 59, 144, *B* 24, 513) Pale yellow plates

Dihydrate $\text{NH} \left\langle \begin{smallmatrix} \text{CO} & \text{CPh}_2 \\ & \text{CHPh CHPh} \end{smallmatrix} \right\rangle$ [237°]

Oxy tetra phenyl pyrrole $C_6H_5N_2O$ [182°] Formed from CPhBz CHBz and alcoholic NH₃ at 200° Yellow prisms Changed into the iso meride [207°] by heating to 310°

o-OXY TRI PHENYL-PYRROLE CARBOXY-

LIC ACID $C_6H_5(OH) \text{N} \left\langle \begin{smallmatrix} \text{CPh C CO}_2\text{H} \\ \text{CPh CH} \end{smallmatrix} \right\rangle$ [245°] Made from its ethyl ether [159°] which is got by the action of o amido phenol on phenacyl benzoyl acetic ether (Paal a Braikoff, *B* 22, 3093) Nodular groups of needles, v sol ether

OXY-PHENYL-QUINAZOLINE

Dihydrate $C_6H_5 \left\langle \begin{smallmatrix} \text{CH}_2 \text{NPh} \\ \text{NH CO} \end{smallmatrix} \right\rangle$ [143°] Formed from phenyl- ω oxy tolyl urea and HCl (Söderbaum a Widman, *B* 22, 1670) Needles

Oxy phenyl quinazoline

$C_6H_5 \left\langle \begin{smallmatrix} \text{C}(\text{OH}) \text{N} \\ \text{N}=\text{CPh} \end{smallmatrix} \right\rangle$ [234°] S (alcohol) 2. Formed by heating benzoyl o amido benzamid (Korner, *J pr* [2] 36, 157) Needles, insol water—B'H₂PtCl₆ brick red needles

Oxy phenyl quinazoline $C_6H_5 \left\langle \begin{smallmatrix} \text{CO NPh} \\ \text{N}=\text{CH} \end{smallmatrix} \right\rangle$

[189°] Formed by oxidation of phenyl quinazoline dihydride with KMnO₄ (Paal a Busch, *B* 22, 2691) Plates, v sol benzene With hydrazine N₂H₄ it forms $C_6H_5 \left\langle \begin{smallmatrix} \text{C}(\text{N}_2\text{H}_2) \text{NPh} \\ \text{N}=\text{CH} \end{smallmatrix} \right\rangle$ [204°]—B'HCl [214°]—Platinochloride [above 300°] Yellow needles

Di-oxy phenyl quinazoline *Dihydrate* $\text{CH}_2\text{CH}_2\text{C} \left\langle \begin{smallmatrix} \text{N} & \text{CPh} \\ & \text{CO CH}_2 \text{C}(\text{OH}) \text{N} \end{smallmatrix} \right\rangle$ [272°] Formed from

di oxy terephthalic ether, dihydride, and benzamidine (Pinner, *B* 22, 2623) Slender needles, sol alkalis, sl sol alcohol

OXY PHENYL-QUINAZOLINE CARBOXY-

LIC ACID $C_6H_5 \left\langle \begin{smallmatrix} \text{N} & \text{CH} \\ \text{CO N} & \text{CO N} \end{smallmatrix} \right\rangle \text{CO}_2\text{H}$ [320°]. Made by oxidising oxy *p* tolyl-quinazoline (Paal a Busch, *B* 22, 2693) White crystals

(a) **OXY (Py 1)-PHENYL-QUINOLINE**
 $C_{11}H_{11}NO \text{ is } C_6H_5 \cdot \begin{smallmatrix} C(C_6H_5OH)CH \\ \backslash \quad \quad \quad / \\ N \quad \quad \quad CH \end{smallmatrix}$ Phenol-quinoline [248°] Formed by diazotising (a)-amido (Py 1)-phenyl-quinoline [150°], and boiling the dilute H_2SO_4 solution (Koenigs a Nef, B 20, 629) Colourless flat needles or plates Volatilises undecomposed V sol alcohol and chloroform, sl sol benzene, nearly insol ether By CrO_3 and H_2SO_4 it is oxidised to cinchoninic acid The hydrochloride forms glistening plates, the sulphate yellow tables, both are v sol hot water, sl sol cold The platino chloride forms hair-fine yellow needles, v sol conc HCl The K and Na salts are colourless glistening plates, which dissolve in water with a yellow colour

(8) **Oxy (Py 1) phenyl quinoline** $C_{11}H_{11}NO$
 $\text{is } C_6H_5 \cdot \begin{smallmatrix} C(C_6H_5OH)CH \\ \backslash \quad \quad \quad / \\ N \quad \quad \quad CH \end{smallmatrix}$ Phenol quinoline [235°] Formed by diazotising the corresponding amido compound [198°] and boiling the dilute H_2SO_4 solution (Koenigs a Nef, B 20, 630) Colourless granular crystal Sl sol alcohol, nearly insol ether Volatilises undecomposed Dissolves with a yellow colour in alkalis and acids By CrO_3 and H_2SO_4 it is oxidised to cinchoninic acid KOH fusion yields di-oxy-phenyl-quinoline [305°] The sulphate crystallises in glistening yellow needles, the hydrochloride in yellow tables, both are m sol cold water

o Oxy (Py 3) phenyl quinoline
 $C_6H_5 \cdot \begin{smallmatrix} CH \quad CH \\ \backslash \quad \quad \quad / \\ N \quad \quad \quad C \quad C_6H_5 \quad OH \end{smallmatrix}$ [115°] Got by heating its (Py 1)-carboxylic acid [238°] (Doebner, A 249, 101) Light yellow needles (from alcohol), sol alkalis and acids $-B'_2H_2PtCl_4$ $-B'_2C_6H_5N_2O$, [184°] Yellow needles

m Oxy-(Py 3) phenyl-quinoline
 $C_6H_5 \cdot \begin{smallmatrix} CH \quad CH \\ \backslash \quad \quad \quad / \\ N \quad \quad \quad C \quad C_6H_5 \quad OH \end{smallmatrix}$ [156°] Formed by the action of nitrous acid upon m amido phenyl-quinoline (Miller a Kinkel, B 18, 1908) Long needles, v sol alcohol and ether

p Oxy (Py 3) phenyl quinoline
 $C_6H_5 \cdot \begin{smallmatrix} CH \quad CH \\ \backslash \quad \quad \quad / \\ N \quad \quad \quad C \quad C_6H_5 \quad OH \end{smallmatrix}$ [238°] Formed from the amido-compounds (Weidel, M 8, 127) Monoclinic needles, insol water, v sol hot benzene Yields quinoline carboxylic acid [157°] on oxidation Yields a crystalline tetrahydride on reduction $-B'HCl$ 2aq $-B'_2H_2PtCl_4$ Acetyl derivative [123°] Plates

(Py 1, 4) **Oxy-phenyl quinoline**
 $C_6H_5 \cdot \begin{smallmatrix} C(OH) \quad CH \\ \backslash \quad \quad \quad / \\ N \quad \quad \quad CPh \end{smallmatrix}$ [254°] Formed, by heating to 250°, from the phenyl β amido cinnamic ether $NPhH$ CPh CH CO_2Et , got by the action of aniline on benzoyl acetic ether (Conrad a Lumpkin, B 21, 521, Knorr, A 245, 377) Got also by heating its carboxylic acid (Just, B 19, 1462) Plates, v sl sol water, insol ether $-B'HCl$ [230°]. Needles (from alcohol) $-B'HCl$ 2aq (J)

(B 2) **Oxy-(Py 3) phenyl-quinoline** *Methyl derivative*
 $C_6H_5 \cdot \begin{smallmatrix} C(OMe) \quad CH \quad C \quad CH \quad CH \\ \backslash \quad \quad \quad / \\ CH \quad CH \quad \quad \quad C \quad N \quad CPh \end{smallmatrix}$ [133°] Got by heating its carboxylic acid [237°] (Doebner, A 249, 106) Plates (from dilute alcohol) $-B'HCl$ needles, sol. hot water $-B'_2H_2PtCl_4$ $-Picrate$ [205°].

(B 4)-**Oxy (Py 3)-phenyl quinoline** *Methyl derivative*
 $CH \quad CH \quad \quad \quad C \quad OH \quad CH$
 $CH \quad C(OMe) \quad C \quad N \quad CPh$
 Formed by heating its carboxylic acid [216°] (D) Oil $-B'_2H_2PtCl_4$ 2aq orange needles

Di oxy phenyl-quinoline $C_{11}H_{11}NO_2$ etc
 $C_6H_5 \cdot \begin{smallmatrix} C(C_6H_5OH) \quad CH \\ \backslash \quad \quad \quad / \\ N \quad \quad \quad C \quad OH \quad (?) \end{smallmatrix}$ [305°] Formed by potash fusion from (8) oxy (Py 1) phenyl quinoline [235°], the yield being 75 p c (Koenigs a Nef, B 20, 632) Needles, v sol alcohol Has no basic properties An isomeride, solid at 315°, appears to be formed from (a)-oxy (Py 1)-phenyl quinoline by potash fusion.

Di oxy-phenyl quinoline [114°] Formed in small quantity when p amido (Py 3) phenyl quinoline is acted upon by nitrous acid (Weidel, M 8, 127) Prisms, insol water Reduced by zinc dust to (Py 3) phenyl quinoline

Di-oxy phenyl-quinoline
 $C(OH) \quad CH \quad C \quad CH \quad CH$
 $CH = CH \quad C - N \quad C(C_6H_5OH)$ [247°] Formed from (B 2) oxy p amido (Py 3) phenyl quinoline by the diazo reaction (Weidel a Georgievitch, M 9, 138) Pale-yellow mass of hair like needles, sl sol hot water Yields (Py 3) phenyl quinoline when distilled with zinc dust

Di-oxy-(Py 1) phenyl-quinoline *Methyl derivative*
 $C(OH) \quad CH \quad C \quad C(C_6H_5OH) \quad CH$
 $CH \quad CH \quad \quad \quad C \quad N \quad \quad \quad CH$
 [188°] Got by boiling the diazo compound of m amido (Py 1)-phenyl (B 2) methoxy quinoline with dilute H_2SO_4 (Miller a Kinkel, B 20, 1922) Plates Its solution in dilute H_2SO_4 gives with chlorine water, a little NaOH, and ammonia, a dark green colour

Tetrahydride [111°] Tablets Gives a deep green with $FeCl_3$ and HCl $-B'HCl$ plates (Py 4)-**OXY (Py 2) PHENYL-ISOQUINOL-**

$C_6H_5 \cdot \begin{smallmatrix} CH \quad CPh \\ \backslash \quad \quad \quad / \\ N \quad \quad \quad CO \quad NH \end{smallmatrix}$ or *Isobenzylidene*

phthalimidine [197°] Formed by the action of ammonia on deoxybenzoin o carboxylic acid CH_2Bz $C_6H_5 \quad CO \quad H$ Prepared by heating crude isobenzylidene phthalide (15 g) with alcoholic NH_3 (30 cc) at 100° for 8-10 hours (Gabriel, B 18, 2449, 3470, 19, 830) Triclinic crystals, $abc = 8608 \text{ 1 ?}$, $\alpha = 88^\circ 28'$, $\beta = 111^\circ 24'$, $\gamma = 74^\circ 47'$ Sl sol cold alcohol By heating with $POCl_3$ or PCl_5 it is converted into (Py 4)-chloro (Py 2) phenyl isoquinoline, PCl_5 yields (Py 1, 4) di chloro (Py 2) phenyl quinoline Nitrous acid converts it into (Py 1) nitro-(Py 4) oxy (Py 2) phenyl isoquinoline

Ethylether $C_{11}H_{11}N(OEt)$ [46°] Formed by heating chloro phenyl isoquinoline with alcoholic sodium ethylate (Gabriel, B 19, 835) Flat colourless needles, v sol ordinary solvents, weak base $-B'_2H_2PtCl_4$, reddish pp.

Di oxy-phenyl-iso-quinoline
 $C_6H_5 \cdot \begin{smallmatrix} C(OH) \quad CPh \\ \backslash \quad \quad \quad / \\ CO \quad \quad \quad NH \end{smallmatrix}$ [257°] Formed from nitro benzylidene phthalide, P, and HI (Gabriel, B 18, 3471, 20, 2866) Crystals, sl sol boiling Aq
Methyl ethers Two isomeric methyl ethers [237°] and [121°] may be prepared Both are crystalline.

Reference — CHLORO OXY PHENYL ISOQUINOLINE

(Py 1, 3, 2)-OXY PHENYL-QUINOLINE CARBOXYLIC ACID $C_6H_4 \begin{smallmatrix} \diagup C(OH) \\ \diagdown N \end{smallmatrix} \begin{smallmatrix} \diagup CO_2H \\ \diagdown CPh \end{smallmatrix}$

[232°] Made from its ethyl ether [262°] which is formed by heating NPh CPh CH(CO₂Et), to 150°, alcohol being split off (Just, B 18, 2632, 19, 1462) Crystals (containing aq), v sl sol hot water —AgA' sl sol water

o Oxy (Py 3, 1)-phenyl quinoline carboxylic acid $C_6H_4 \begin{smallmatrix} \diagup C(CO_2H)CH \\ \diagdown N \end{smallmatrix} C_6H_4OH$ [238°] Made from salicylic aldehyde, aniline, and pyruvic acid (Doebner, A 249, 98) Needles, v sl alcohol —AgA'—H₂A'.H₂PtCl₆, golden yellow needles.

(B 2, Py 3, 1) Oxy phenyl quinoline carboxylic acid *Methyl derivative* C(OMe) CH C C(CO₂H) CH CH CH—C—N CPh *Phenyl quinic acid* [237°] Formed by heating together

equimolecular quantities of *p* amido anisole, benzene aldehyde, and pyruvic acid in an alcoholic solution (Doebner, A 249, 105) Needles, insol water —AgA'—H₂A'.H₂PtCl₆, orange red crystals

(B 4, Py 3, 1) Oxy phenyl quinoline carboxylic acid *Methyl derivative* CH CH—C C(CO₂H) CH CH C(OMe) C—N CPh [216°] Formed in like manner from [2 1]C₆H₄(NH₂)OMe, benzoic aldehyde, and pyruvic acid (D) Yellow needles —PbA'.aq —AgA'—H₂A'.H₂PtCl₆.

DI OXY DI-PHENYL QUINOXALINE *Diethyl derivative* CH C(OEt) C N CPh CH C(OEt) C N CPh

[163°] Formed by the action of benzil and NaOAc on an alcoholic solution of the diethyl ether of (a) di amido hydroquinone (Nietzki & Rechberg, B 23, 1212) Yellow needles

Di oxy di phenyl quinoxaline Dihydrate *Di methyl derivative*

$C_6H_4 \begin{smallmatrix} \diagup NCHC_6H_4OMe \\ \diagdown NCHC_6H_4OMe \end{smallmatrix}$ *Phenylanisaldehyde* [129°] Formed from anisic aldehyde and phenylene *o* diamine (Rugheimer a Ladenburg, B 11, 1660) Needles (from alcohol)

α-OXY PHENYL-SUCCINIC ACID

$C_6H_4C(OH)(CO_2H)CH_2CO_2H$ *Phenyl maleic acid* [188°] S 159 at 15° Formed by heating a bromo phenyl succinic acid with hot water (Alexander, A 258, 76) Needles (from chloroform) Yields, on heating, phenyl maleic anhydride [119°], phenyl fumaric acid [161°], and a sublimate [106°] probably atropic acid

β Oxy-phenyl succinic acid $C_6H_4CH(CO_2H)CH(OH)CO_2H$ [150°–160°] S 37 35 at 15° Got from C₆H₄CH(CHO)CO₂Et, potassium cyanide, and HCl (A) Crystalline powder, v sol water Yields phenyl maleic anhydride when heated to 160°

Di-oxy-di-phenyl succinic acid CO₂H CPh(OH) CPh(OH) CO₂H *The nitrile* of this acid CPh(OH)Cy CPh(OH)Cy [182°] is formed by union of benzil with HCN (Zimin, A 34, 189, Jacoby, B 19, 1519) It is converted in the cold by HOAc saturated with HBr into the amide C₆H₄N₂O [150°–230°] (Burton, B 16, 2282)

DI-OXY-DI-PHENYL SULPHIDE S(C₆H₄OH)₂ [151 5°] Formed from 'thio-

aniline' by the diazo reaction (Krafft, B 7, 1165, 22, 821), and also by the action of SOCl₂ on phenol in presence of CS₂ (Tassinari, G 17, 88) Silvery plates, sl sol cold water.

Di acetyl derivative [93°]

An isomeride [129°] may be obtained from S(C₆H₄Br OH)₂ [176°] by reduction with zinc-dust (T)

Reference — DI BROMO DI-OXY-DI PHENYL SULPHIDE

Di p-oxy-di-phenyl disulphide S₂(C₆H₄OH)₂ [151°] Formed by oxidising *p*-oxy phenyl mercaptan (thio hydroquinone) with dilute FeCl₃ (Leuckart, J pr [2] 41, 196) Yellow needles, v e sol alcohol

Di acetyl derivative [89°]. Plates

Di o oxy di-phenyl disulphide

S₂(C₆H₄OH)₂ Formed by heating sodium phenate (2 mols) with sulphur (1 at) at 190° (Hartinger, M 4, 165) Thick oil Yields *o* oxy-phenyl mercaptan on reduction —NaHA"6aq —KHA"5aq —PbA" yellow pp

Di methyl derivative Me₂A" [119° cor]. Yields anisole *o* sulphonic acid on oxidation

DI-OXY DI-PHENYL SULPHONE

SO₂(C₆H₄OH)₂ [239°] S G 15 1 366 Formed by heating phenol (2 pts) with fuming H₂SO₄ (1 pt) at 190° (Glutz, A 147, 52, Annaheim, J pr [2] 1, 14, 2, 385, B 8, 1059, 11, 1668, A 172, 28) Trimetric crystals, a b c = 1.283 1.0 534. V sol hot water and alcohol —K A" Sol. alcohol (Guarachi, G 8, 246) —NH₄HA" —NaHA"aq —AgHA" —AgA"

Di acetyl derivative Needles (Glutz)

Di methyl ether Me₂A" [130°]

Di ethyl ether [159°] Laminæ

Di isoamyl ether [98°] Plates

Di oxy di-phenyl sulphone SO₂(C₆H₄OH)₂

[187°] Got from its acetyl derivative which is made by oxidation of C₆H₄(OAc)(SH) [1.2 or 3] (Tassinari, C C 1888, 1354) Crystalline, sl sol water

References — DI BROMO, DI BROMO DI-NITRO-, DI CHLORO, TETRA IODO, DI IODO-DI-NITRO, and DI NITRO, DI OXY DI PHENYL SULPHONE

OXY PHENYL SULPHONIC ACID *o* PHENOL SULPHONIC ACID

Oxy diphenyl sulphonic acid C₆H₅(OH)SO₃H Made, together with the disulphonic acid, by heating oxydiphenyl (1 pt) with H₂SO₄ (3 pts) (Latschnoff, J R 5, 54) —KA' aq plates, sl sol water —BaA'₂aq —CaA'₂3aq —K₂CuA'₂ 6aq green laminæ

Oxy diphenyl disulphonic acid

C₆H₅(OH)(SO₃H)₂ Made as above —K₂A" 1½aq.

Di oxy diphenyl disulphonic acid

[4 2 1] C₆H₅(OH)(SO₃H) C₆H₅(SO₃H)(OH) [1 2 4] Made from benzidine disulphonic acid by the diazo reaction (Lamprecht, A 261, 334) Crystalline mass The K salt forms laminæ, v e sol. water —BaA"2aq —PbA"4aq rhombohedra.

Di p-oxy diphenyl disulphonic acid

C₆H₄(OH)₂(SO₃H)₂ Got by warming di-oxy-diphenyl with fuming H₂SO₄ (Doebner, B 9, 130). Probably identical with the preceding acid. —K₂A" prisms, sl sol cold water

Di p-oxy-diphenyl trisulphonic acid

C₆H₄(OH)₂(SO₃H)₃ Formed, together with the tetra-sulphonic acid, from benzidine by diazotising and warming with conc H₂SO₄ (Griess, J 1866, 462) —Ph₂C₆H₄S₃O₉ 2aq —Ph₂C₆H₄S₄O₁₀ 2PbO. 8 c 2

Di *p*-oxy-diphenyl tetrasulphonie acid
 $C_{12}H_8(OH)_2(SO_3H)_4$ Made as above—BaA^{iv} 54q
 prisms— Pb, C, H, S, O_2 amorphous pp

***o*-OXY-PHENYL-SULPHONO-180BUTYRIC ACID**
 $C_{12}H_8, SO_2, CH_2, OMe(OH) CO_2H$ [121°]
 Made by oxidising $CH_2(SPh) CMe(OH) CO_2H$
 with $KMnO_4$ in the cold (Delisle, *A* 260, 259)
 Prisms (from ether)—KAⁱ 2aq needles—BaAⁱ,
 —CaAⁱ, aq groups of needles

OXY-PHENYL SULPHURIC ACID

$C_6H_4(OH) O SO_3H$ The K salts of the *o*-, *m*-,
 and *p*-compounds are formed by the action of
 $K_2S_2O_8$ on potassium pyrocatechin, resorcin, and
 hydroquinone respectively (Baumann, *B* 11,
 1918) They are crystalline, sol water, and
 unstable Potassium pyrogallol yields, in like
 manner, $C_6H_3(OH)_3SO_3K$ crystallising in needles

OXY-PHENYL-THIAZOLE $\begin{matrix} S(OH) \\ CH OPh \end{matrix} \gg N$

[204°] Formed from bromo acetophenone by
 the action of xanthamide (Hubacher, *A* 259,
 249), or by successive treatment with barium
 sulphocyanide and dilute HCl (Arapides, *A* 249,
 14, of Dyckerhoff, *C J* 92, 327) Needles,
 insol water, sol cold $NaOH$ aq Converted by
 PCl_5 into chloro-oxy-phenyl thiazole [206°] and
 another body [98°] (Schatzmann, *A* 261, 18).

Sodium and EtI yield $\begin{matrix} S CO \\ CH OPh \end{matrix} > NEt$ [71°],
 which on heating with HCl at 220° gives $NEtH_2$.

Di-*exo-o*-oxy-phenyl-thiazole dihydride

Methyl derivative $\begin{matrix} S-CO \\ CH_2, CH_2 \end{matrix} > N C_6H_4OMe$

[116°] Formed by the action of boiling alco-
 holic chloro acetic acid on the product of the
 action of CS_2 on the substance formed from
 ethylene bromide and di anisyl thio-urea (Foer-
 ster, *B* 21, 1867) Plates, v sol alcohol

***p*-OXY-PHENYL THIOCARBIMIDE** *Acety-*
l derivative $C_6H_4(OAc) NCS$ [36°] Made
 by heating di *p*-oxy di phenyl urea with Ac_2O
 (Kalkhoff, *B* 16, 1831) Plates, sol alcohol

Methyl derivative $C_6H_4(OMe) NCS$
 [265°] Formed from anisidine and CS_2 (Hof-
 mann, *B* 20, 1796, of Salkowski, *B* 7, 1012)

Isomeride **OXY METHENYL AMIDO PHENYL-**
MERCAPTAN

Di-oxy phenyl-thiocarbimide *Di-methyl*
derivative $C_6H_4(OMe)_2 NCS$ Formed by heat-
 ing $CS(NH C_6H_4(OMe)_2)_2$ with conc HCl aq
 (Bechhold, *B* 22, 2381) Occurs in three crys-
 talline forms [57°], [120°], and [155°] Sol
 aniline and H_2SO_4

DI-OXY-DI-PHENYL-THIOHYDANTOIN
Di-methyl derivative $C_{12}H_{14}N_2SO_2$, i.e.
 $C_6H_4(OMe) N C N(C_6H_4OMe) S CH_2 CO$ [190°] Made

from di-anisyl thio urea and fused chloro acetic
 acid (Foerster, *B* 21, 1867) Needles

***p*-OXY-PHENYL-THIO-UREA** C_6H_4NH, SO i.e.
 $C_6H_4NH CS NH(OH)$ [108°] Formed from

phenyl-thiocarbimide and aqueous hydroxyl
 amine (Schiff, E Fischer, *B* 22, 1935, Tie-
 mann, *B* 22, 1939, Von der Kall, *A* 263, 260,
 Voltmer, *B* 24, 378) White plates, insol water,
 sl sol ether, v sol alcohol Fehling's solution
 ppt Os_2S Dilute acids and alkalis decompose
 it, yielding phenyl cyanamide

Methyl derivative $NHPh CS NH OMe$
 [116°] Formed from phenyl-thiocarbimide and
 methyl-hydroxylamine.

Ethyl derivative [103°] Prisms De-
 composed, by boiling its alcoholic solution, into
 phenyl cyanamide, alcohol, and S

Benzyl derivative [113°] Prisms

o-Oxy-phenyl-thio-urea

$C_6H_4(OH) NH CS NH_2$ [161°] Made from *o*-
 amido phenol hydrochloride and potassium sul-
 phocyanide (Bendix, *B* 11, 2263) Needles, sol
 hot water— B^i, H_2PtCl_6

Methyl derivative

$C_6H_4(OMe) NH CS NH_2$ [152°] Needles (from
 alcohol) (Muhlhauser, *B* 13, 923)

Ethyl derivative

[110°] Tablets, sol
 alkalis (Berlinerblau, *J pr* [2] 80, 106)

p-Oxy-phenyl thio-urea

[41] $C_6H_4(OH) NH CS NH_2$ [214°] Got by
 evaporating to dryness on the water-bath, a so-
 lution of potassium sulphocyanide and *p* amido
 phenol hydrochloride (Kalkhoff, *B* 16, 375).
 Tables, sl sol cold water

Ethyl derivative Crystalline (B)

o Oxy-diphenyl-thio-urea

$NHPh CS NH C_6H_4OH$ [146°] Formed by the
 action of phenyl thiocarbimide on *o*-amido-
 phenol (Kalkhoff, *B* 16, 1829) White pearly
 plates Heated by itself or with HgO it loses
 H_2S , yielding anilido carbamido phenol

Methyl derivative

$NHPh CS NH C_6H_4OMe$ [127°] Formed from
 anisyl thiocarbimide and aniline, and also from
 phenyl-thiocarbimide and anisidine (Foerster,
B 21, 1868) Prisms With ethylene bromide
 it forms crystals [143°] whence platine chloride
 yields $(C_6H_4N_2OS)_2, H_2PtCl_6$

p-Oxy-di-phenyl-thio urea

$NHPh CS NH C_6H_4OH$ *p* Oxy thiocarbimide
 [162°] Formed by the action of phenyl thio-
 carbimide on *p* amido phenol (K) V sol alcohol
 and alkalis, nearly insol water

Acetyl derivative

[137°] Insol alkalis
Di *o*-oxy-di-phenyl thio urea *Di-methyl*
derivative $CS(NH C_6H_4OMe)_2$ [135°] Formed
 from *o* anisidine, CS_2 , and alcoholic potash
 (Muhlhauser) Formed also from anisidine and
 thio urea in alcoholic solution (F) Needles, sl
 sol water With methyl iodide it forms
 $C_6H_4(OMe) N C(SMe) NH C_6H_4OMe$ [87°] EtI
 forms $C_{12}H_{18}N_2SO_2$ [82.5°], while propyl iodide
 gives $C_{18}H_{28}N_2SO_2$ [58°] Ethylene bromide gives
 $CH_2 < \begin{matrix} S \\ CS \end{matrix} CH_2$ crystallising in trimetric
 prisms [186°] and yielding $C_6H_4NH_2NS_2OMeI$
 [c 141°] and, on boiling with alcohol and chloro-
 acetic acid, $S < \begin{matrix} CO \\ CH_2 \end{matrix} > NC_6H_4OMe$ [116°]

Di-*p*-oxy-di-phenyl thio-urea [222°] Got
 from *p* amido phenol and CS_2 (K) Pearly
 plates, v sol alkalis

Di-methyl derivative [185°] Plates
 (Salkowski, *B* 7, 1012) Sl sol alcohol

Di-oxy-di-phenyl-thio urea

Di-methyl derivative $NHPh CS NH C_6H_4(OMe)_2$ [137°]
 Formed by the combination of phenyl thiocarb-
 imide with amido-di methyl-hydroquinone
 (Baessler, *B* 17, 2123) Minute needles, v sol.
 hot benzene and hot alcohol

Tetra-oxy-di phenyl thio urea

Tetra-
methyl derivative $CS(NH C_6H_4(OMe)_2)_2$,
 [160°] Formed from the di methyl derivative of
 amido resorcin and CS_2 (Bechhold, *B* 22, 2380).
 White mass (from alcohol-ether), insol alkalis

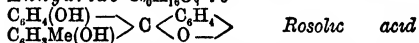
An isomeride [109°] is formed from the dimethyl derivative of amido-hydroquinone (Baessler)

Reference - DI CHLORO DI OXY-DI-PHENYL-THIO-UREA

OXY-PHENYL-TOLYLAMINE v TOLYL-AMIDO PHENOL

TRI OXY-DI-PHENYL-TOLYL-CARBINOL

Anhydride $C_{20}H_{16}O_4$, i.e.



Corallin Formed by the action of nitrous acid on rosaniline (Caro & Wanklyn, *J pr* 100, 49, A 179, 192) Got also by heating a mixture of phenol and cresol with H_2SO_4 and arsenic acid (Zulkowsky, *B* 10, 1201) Red plates with green lustre, nearly insol water, sol chloroform, alcohol, and acids Alkalis form a red solution Reduced by sodium amalgam to tri oxy di-phenyl tolyl methane Water at 240° yields phenol and $C_6H_4(OH)CO C_6H_4Me(OH)$ KCl followed by HCl forms crystalline $C_{20}H_{16}NO_4$, whence Ac_2O yields $C_{21}H_{14}Ac_2NO_4$ [142°]

p-OXY-PHENYL p-TOLYL-HYDRAZINE

Ethyl derivative

$C_6H_4(OEt)NH NH C_6H_4Me$ Formed by reducing $C_6H_4(OEt)N_2 C_6H_4Me$ with alcoholic ammonium sulphide (Noelting & Werner, *B* 23, 3258, *Bl* [3] 4, 794) Needles, v sol hot alcohol The isomerides $C_6H_4NH NHCO C_6H_4Me(OEt)$ [1 3 4] [76°] and $C_6H_4NH NHCO C_6H_4Me(OEt)$ [1 5 2] [105°] are formed in like manner

DI OXY-PHENYL-TOLYL KETONE

$C_6H_4(OH)CO C_6H_4Me(OH)$ [200°] Formed by heating rosolic acid with water at 220°-250° (Graebe & Caro, A 179, 196) Crystals, sol hot water

Di acetyl derivative [150°] Needles

Di oxy phenyl tolyl ketone Anhydride

$C_6H_4 \begin{array}{c} \diagup \\ O \\ \diagdown \end{array} C_6H_4Me$ [105°] Made by boiling p tolyl salicylate for a long time (Graebe & Feer, *B* 19, 2612) V sol hot alcohol

Tri oxy phenyl-tolyl ketone

$C_6H_4(OH)CO C_6H_4Me(OH)_2$ Anhydride $C_{18}H_{14}O_4$ [140°] Made by fusing salicylic acid with orcin and $ZnCl_2$ (Michael, *Am* 5, 95) Yellow needles (from alcohol) -NaA' 14aq

Acetyl derivative $C_{19}H_{14}AcO_4$ [152°]

DI OXY-PHENYL-DI-TOLYL METHANE

$CHPh(C_6H_4MeOH)_2$ [171°] Formed from o-cresol benzoin and boiling SO_2Aq (Schroeter, A 257, 70) Yellow needles Yields a di bromo-derivative [130°] and a di nitro-derivative [127°]

Di acetyl derivative $C_{22}H_{16}Ac_2O_4$ [94°]

Di benzoyl derivative [91 5°] Plates

Tri oxy-di-phenyl-tolyl methane

$C_6H_4Me(OH)CH(C_6H_4OH)_2$ Phenolphthalol [190°] Obtained by boiling phenol phthalin $CO_2H C_6H_4CH(O_2H)_2$ with $HOAc$ and sodium-amalgam (Baeyer, A 202, 87) Prisms (from dilute $HOAc$), v sol alcohol

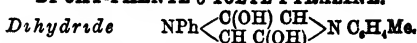
Tri-acetyl derivative $C_{22}H_{14}O_4$ [40°]

Tri-oxy di-phenyl-tolyl-methane $C_{20}H_{16}O_4$, i.e.

$C_6H_4Me(OH)CH(O_2H)_2$ Leucorosolic acid Got by reducing rosolic acid with potash and zinc-dust (Graebe & Caro, A 179, 198) Needles (from alcohol), v al sol water

Tri-acetyl derivative. [149°]. Needles.

DI OXY-PHENYL o TOLYL PYRAZINE.



[166°] Formed by heating aniline with chloro-acetyl o tolyl amido acetic acid at 160° (Abenius, *J pr* [2] 40, 443) Formed also by the action of o toluidine on chloro acetyl phenyl amido-acetic acid in presence of $NaOAc$ (Bischoff, *B* 23, 1996) White needles, v sol hot alcohol The isomeric compound from p-toluidine also crystallises in needles [221°]

OXY-PHENYL-p-TOLYL-PYRIMIDINE



from tolenyl amidine and benzoyl acetic ether (Pinner, *B* 23, 3826) Slender needles, sl sol hot alcohol

o OXY PHENYL o TOLYL-THIO-UREA

$CH_2(OH)C_6H_4NH OS NHPH$ [o 136°] Made from phenyl thiocarbimide and o amido benzyl alcohol (Söderbaum & Widman, *B* 22, 1871). Prisms Alcohol and HgO convert it into



o OXY-PHENYL TOLYL-UREA

$CH_2OH C_6H_4NH CO NHPH$ [191°] Made from o amido benzyl alcohol and $PhNCO$ (Söderbaum & Widman, *B* 22, 1670) Needles

v OXY-PHENYL UREA $C_6H_4NH CO NH OH$

[140°] Made from phenyl cyanate and aqueous hydroxylamine (E Fischer, *B* 22, 1935, von der Kall, A 263, 264) Crystals, v al sol water, sol alkalis Reduces hot Fehling's solution On boiling with alkalis it yields aniline, CO_2 , and hydroxylamine

Benzyl derivative

$NHPH CO NH OCH_2Ph$ [106°] Needles (Tiemann, *B* 22, 1940, Voltmer, *B* 24, 384)

o Oxy phenyl-urea $C_6H_4(OH)NH CO NH_2$

[154°] Made from o amido phenol hydrochloride and potassium cyanate (Kalkhoff, *B* 18, 374) Prisms, v sol water

Methyl derivative

$C_6H_4(OMe)NH CO NH$ [147°] Crystals (Muhlhauser, *B* 13, 922, A 207, 244)

p-Oxy phenyl-urea $C_6H_4N_2O_4$ [168°] (K).

Needles or tables, sol acids and alkalis

Ethyl derivative [160°] Made from

$C_6H_4(NH_2)OEt$ and $KCyO$ (Berlinerblau, *J pr* [2] 30, 103) Nitrous acid passed into its alcoholic solution forms brick-red crystals of $C_6H_4N_2O_4$

Di-o-oxy di-phenyl urea Di-methyl

derivative $CO(NH C_6H_4OMe)_2$ [182°] Made from $COCl_2$ and o anisidine (M., Conrad, *B* 21, 1649) Crystals, v sol alcohol

Di p oxy-di phenyl urea Di-methyl derivative

$C_{12}H_{10}N_2O_4$ [234°] Prisms or needles (Lossen, A 175, 295, Pieschel, A 175, 812) V sl sol hot alcohol

β-OXY PHENYL VALERIC ACID $C_{11}H_{14}O_4$, i.e.

$CHPh(OH)CMe_2CO_2H$ Phenyl-oxypivalic acid [184°] Formed from benzoic aldehyde, sodium isobutyrate, and isobutyric anhydride by Perkins's reaction (Fittig & Jayne, A 216, 119, Ott, A 227, 61) Needles, sl sol. cold water. Split up at 150° into CO_2 and phenyl-butylene. -BaA', 4aq -CaA', 4aq plates, m. sol. water

Acetyl derivative [187°]. Monoclinic crystals (from CS_2). Yields $Ba(C_{11}H_{13}O_4)_2$ 2aq and CaA' 2aq

Anhydride $C_{10}H_{16}O_4$, [155°] Prisms
Iso-butyryl derivative [65°] Needles,
 v. sl. sol water Yields $Ba(C_{11}H_{17}O_4)_2$, 2aq,
 CaA'_2 , 2aq, and AgA'

β -Oxy-phenyl-n-valeric acid
 $C_6H_5CH_2CH_2CH(OH)CH_2CO_2H$ [131°] Got
 by boiling β styryl-propionic acid with $NaOH$ aq
 (Fittig, *B* 24, 84) Prisms

β -Oxy-phenyl-valeric acid
 $CH_3CH(OH)CH(CH_2Ph)CO_2H$ [153°] Got
 by reducing benzyl aceto acetic ether (Ehrlich,
B 8, 1086, *A* 187, 26) Needles (from benzene)
 $-BaA'_2$, 2aq $-Zn(OH)A'$ $-Cu(OH)A'$

γ -Oxy- α -phenyl-valeric acid
 $CH_3CH(OH)CH_2CHPhCO_2H$ Formed from
 α -phenyl- β acetyl propionic acid by reduction
 with sodium amalgam (Weltner, *B* 17, 73)
 The free acid at once forms the only anhydride
 $C_{11}H_{18}O_4 - Ca(C_{11}H_{17}O_4)_2$ aq crystalline

γ -Oxy- γ -phenyl-isovaleric acid The only
 lactone $C_9H_{14}O_3$ $\begin{smallmatrix} CHMe \\ \diagup \quad \diagdown \\ O \quad CO \end{smallmatrix} CH_2$ is made by
 warming γ -bromo γ -phenyl isovaleric acid with
 aqueous Na_2CO_3 (Fittig a Liebmann, *A* 255,
 271) It yields $Ba(C_{11}H_{17}O_4)_2$

Oxy-phenyl-valeric acid **Methyl deriva-**
tive $C_6H_5(OMe)CH_2CH_2CH_2CO_2H$ Got
 by reducing $C_6H_5(OMe)C_2H_4CO_2H$ with sodium-
 amalgam (Perkin, *C* 39, 438) $-Oil - BaA'_2$

Di-oxy-phenyl-valeric acid **Methylene**
derivative $C_6H_5OCH_2CH_2CO_2H$ *Piperhy-*
droic acid [96°] Made by reducing (8) hydro-
 piperic acid with sodium amalgam (Fittig a
 Buri, *A* 216, 178) Thin tables (from alcohol) $-$
 CaA'_2 aq v sl sol cold water

Di-oxy-phenyl-valeric acid
 $C_6H_5CH(OH)CH_2CH(CO_2H)CH(OH)CH_3$ The
 only anhydride of this acid is got by reducing
 phenacyl-acetoacetic ether (Wellner, *B* 17, 69)

Tri-oxy-phenyl-valeric acid **Methylene**
derivative

$CH_3OCH_2C_6H_4CH_2CH(OH)CH_2CH_2CO_2H$ [95°]
 Made by reducing with sodium amalgam the acid
 $CH_3OCH_2C_6H_4CH_2COCH_2CH_2CO_2H$ (Weinstein,
A 227, 38) Crystals, m sol hot water Yields
 an oily lactone $-BaA'_2 - AgA'$ flocculent pp

Tri-ppr-oxy di phenyl-valeric acid **Di-**
methyl derivative

$C_6H_5(OMe)CH(OH)CH_2CH(CO_2H)CH_2C_6H_5OMe$
 The lactone [83°] is got by the action of sodium-
 amalgam on its bromo derivative [136°], which
 is formed by dissolving di bromo di p methoxy-
 di-phenyl-valeric acid in $HOAc$ (Fittig a Politis,
A 255, 305) It yields a gummy Ba salt

Tetra-oxy-phenyl-valeric acid **Methylene**
derivative

$CH_3OCH_2C_6H_4CH_2CH(OH)CH(OH)CH_2CO_2H$
 [128°] Made by oxidising (a) hydropiperic
 acid with $KMnO_4$ at 0° (Regel, *B* 20, 415)
 Slender crystals $-BaA'_2 - AgA'$ needles

Anhydride $C_{11}H_{18}O_5$ [104.5°] Crystals.
 An isomeric acid

$CH_3OCH_2C_6H_4CH_2CH_2CH(OH)CH(OH)CO_2H$
 [165°], formed by oxidising (8) hydropiperic
 acid, yields the salts CaA'_2 aq and AgA'

Reference $-Tetra-bromo-oxy-phenyl-val-$
eric acid

c OXY-PHTHALIC ACID $C_6H_5(OH)(CO_2H)_2$
 [123] [α 200°] *S* 20 at 17° Obtained by
 potash fusion from its methyl derivative (Jacob

sen, *B* 16, 1965), and from c sulpho phthalic
 acid (Stokes, *Am* 6, 282) Got also from c -
 amido-phthalic acid (Bernthsen, *B* 20, 937).
 Prisms (from water), v sol alcohol and ether
 Coloured red by $FeCl_3$ Forms a fluorescein
 when heated with resorcin $-KHA''$ needles $-$
 AgA'' pp, sl sol hot water

Anhydride $C_6H_5(OH) \begin{smallmatrix} CO \\ \diagup \quad \diagdown \\ CO \end{smallmatrix} O$ [α 148°]

Formed by heating the acid

Methyl derivative $C_6H_5(OMe)(CO_2H)_2$
 [160°] Formed by oxidation of c methoxy-
 toluic acid with $KMnO_4$ Minute prisms

Anhydride of the methyl derivative

$C_6H_5(OMe)C_2O_2$ [87°] Needles (by sublimation)
 α -Oxy-phthalic acid $C_6H_5(OH)(CO_2H)_2$ [1344]
 [185°] (Rée), [205°] (Claus) *S* 3 at 10°

Formation -1 From t -amido phthalic acid
 by the diazo reaction (Baeyer, *B* 10, 124, 1079)
 2 By oxidising α methoxy o toluic acid with
 potassium permanganate, and heating the re-
 sulting $C_6H_5(OMe)(CO_2H)_2$ with hydrochloric acid
 (Schall, *B* 12, 816) -3 By potash fusion from
 $C_6H_5(SO_2NH_2)(CO_2H)_2$, which is got by oxidising
 [341] and [521] $C_6H_5Me(SO_2NH_2)(CO_2H)$ (Ja-
 cobsen, *B* 14, 42) -4 By fusing i sulpho phtha-
 lic acid with $NaOH$ (Graebe a Rée, *C* 49, 524,
B 18, 1130, 1630) -5 By fusing di chloro-(a)
 naphthoquinone sulphonic acid with potash
 (Claus, *J* pr [2] 87, 194) -6 By fusing di
 nitro naphthol sulphonic acid with potash (Rée,
A 233, 230)

Properties $-$ Needles, sl sol cold water, sol
 alcohol and ether Gives a yellowish red colour
 with $FeCl_3$ Dilute HCl at 180° forms m oxy
 benzoic acid Gives the fluorescein reaction
 with resorcin at 200° $-AgA''$ tufts of needles

Methyl ether Me_2A'' [102°] Plates

Anhydride $C_6H_5O_4$ [165°] Made by
 heating the acid

Methyl derivative $C_6H_5(OMe)(CO_2H)_2$
 [138°-144°] Needles Forms on heating an
 anhydride [93°] $-Ag_2C_6H_5O_4$ curdy pp

α Oxy-isophthalic acid $C_6H_5(OH)(CO_2H)_2$

[135] [288° cor] *S* 0305 at 5°, 19 at 99°

Made by fusing s sulpho isophthalic acid with
 potash (Heine, *B* 13, 491, Lommies, *B* 13, 705)

Got also from rufigallie acid by potash fusion
 (Schreder, *M* 1, 437), and by the action of
 nitrous acid on s -amido isophthalic acid (Beyer,
J pr [2] 25, 515) Needles (containing 2aq), v

sol hot water, alcohol, and ether $-BaA''$ 3aq $-$
 $Cu_2(C_6H_5O_4)_2$ 4aq $-Ag_2A'' - AgHA''$ needles

Di-methyl ether Me_2A'' [160°] Needles

Di-ethyl ether Et_2A'' [103°] Prisms

c -Oxy-isophthalic acid $C_6H_5(OH)(CO_2H)_2$
 [126] [239°] *S* 8 at 100°, 14 at 24°

Formation -1 Obtained by potash fusion
 from (2,1,6) aldehyde oxy benzoic acid (Tiemann
 a Reimer, *B* 10, 1562), and also from the acid
 [126] $C_6H_5(SO_2NH_2)(CO_2H)_2$ (Jacobsen, *B* 11,
 902) -2 From (1,2,6) methoxy-toluic acid by
 oxidising with $KMnO_4$, and heating the resulting
 $C_6H_5(OMe)(CO_2H)_2$ with HCl at 160° (Schall, *B*
 12, 826) -3 In small quantity, in the prepara-

tion of the following isomeride, by the action of
 CCl_4 and KOH on salicylic acid (Hasse, *B* 10,
 2185) -4 By oxidising a naphthol with CrO_3
 and $HOAc$ (Miller, *A* 208, 247) -5 From c -
 amido isophthalic ether (*M*)

Properties $-$ Needles (containing aq), v sol.

alcohol and ether. Melts at 244° after drying at 100° (T & R). Coloured cherry red by FeCl₃. Forms salicylic acid when strongly heated. Its solutions exhibit bluish violet fluorescence -- Ag, A'' minute plates -- Et, A'' oil.

Mono-methyl ether

$\text{C}_6\text{H}_5(\text{OH})(\text{CO}_2\text{H})(\text{CO}_2\text{Me})$ [135°] Made from
 $[\text{1 2}]\text{C}_6\text{H}_5(\text{ONa})\text{CO}_2\text{CH}_3$ by heating with CO_2 at
 150° under pressure (Hahle, *J pr* [2] 44, 6)
 Grouped needles, which melt under water
 Coloured red by FeCl_3 Gives NaA' as colourless
 needles

Methyl derivative $C_6H_3(OMe)(CO_2H)_2$
[218°] Prisms, sol hot water

2 Oxy isophthalic acid $C_6H_4(OH)(CO_2H)_2$

[1 2 4] [306"] S 02 at 10°, 625 at 100°
Formation—1 By potash fusion from (2,1,4)- and (4,1,2) aldehyde oxy benzoic acid (Tiernann a Reimer, B 10, 1562), from *m* xyenol (Jacob sen, B 11, 377), from *m* xylene sulphonic acid (Remsen, B 11, 580), from $C_6H_5(SO_2NH_2)(CO_2H)$, (Remsen, *Am* 1, 131), and from benzoic acid (Barth, M 3, 803)—2 By the action of KOH, dilute alcohol, and CCl₄ on salicylic acid (Hasse) 3 By heating $C_6H_5(ONa)CO_2Na$ [1 2] or [1 4] in a current of CO (Ost, J pr [2] 14, 93, 15, 301. Kunferberger, J pr [2] 16, 428).

Preparation—A mixture of C_6H_5ONa (3 mols) and C_6H_5OK (1 mol) is heated in a current of CO_2 at 300° . The yield is 30 p.c. of the phenol used (Ost).

Properties—Needles, sl sol water, insol chloroform. Not volatile with steam. Its aqueous solution is coloured cherry red by FeCl_3 . Yields salicylic acid and phenol when distilled. Br $\text{C}_6\text{H}_4\text{Br}$ bromo phenol on heating. $-\text{Na A}^{\cdot\cdot}\text{aq} - \text{Ca}_3(\text{C}_6\text{H}_5\text{O}_3)_2 \cdot 5\text{aq} - \text{CdA}^{\cdot\cdot}\text{aq} - \text{Ag}_2\text{A}^{\cdot\cdot} - \text{AgHA}^{\cdot\cdot}$ needles

Methylether $\text{Me}_2\text{A}''$ [96°]

Mono ethylether

$C_6H_5(OH)(CO_2H)(CO_2Et)$ [4 3 1] [195°] Made from [4 1] $C_6H_5(ONa)CO_2Et$ and CO_2 at 170° under pressure (Hahle, *J pr* [2] 44, 12) Monoclinic leaflets

Diethyl ether Et₂A'' [52°]

Phenylethylether
 $C_6H_5(OH)(CO_2Ph)(CO Et)$ [4 3 1] [65°] Got
 from $HEtA''$, phenol, and $POCl_3$ (H)

Amide $C_6H_3(OH)(CONH_2)_2$ [250°]

Methyl derivative $C_9H_8O_3$ [261°]

Oxy isophthalic acid *Hexahydrate*

$\begin{array}{l} \text{CH}_2\text{CH}_2\text{C}(\text{OH})(\text{COH}) \\ \text{CH}_2\text{CH}(\text{CO}_2\text{H})\text{CH}_2- \end{array} >$ Formed by saponi-
 cation of its semi nitrile $[\text{130}^\circ\text{--}\text{140}^\circ]$ which is
 the product of the union of prussic acid with
 $\begin{array}{l} \text{CH}_2\text{CH}_2\text{CO} \\ \text{CH}_2\text{CH}(\text{CO}_2\text{H}) \end{array} > \text{CH}_2$ (Baeyer & Tutein, *B* 22,
 2186) Nodules (from alcohol) — $\text{Ag}_2\text{A}''$

Di oxy phthalic acid Methyl derivative
 $C_8H_6(O_2Me)(OH)(CO_2H)$ *Normethylhemipic acid*
 [2255] (Wegscheider, *M* 3, 378) Made by
 warming hemipic acid with conc HIAq (Liechti,
A Suppl 7, 151, Beckett & Wright, *C J* 29,
 281) Nodules (containing 2aq) Melts at 150°-
 155° when hydrated V sol water and alcohol
 Coloured blue by FeCl₃ Yields protocatechuic
 acid when fused with potash The anhydride
 $C_8H_4O_2$ 2aq forms crystals [1489]— KHA^+

D-methyl derivatives of Hemipic acid

Methylene derivative
CH₂O.C₆H₄(CO₂H)₂ [175°] *Hydrastic acid.*

Got by the action of boiling potash on the methylimide [233^o] which is obtained by oxidizing hydrastinic acid (Freund & Lachmann, *B* 22, 2325) Needles (from Aq) — $\text{NMeH.A}'$ [224^o]

Di-oxy phthalic acid *Hexahydrate*
 $C_8H_6O_8$ [180°] Got by the action of baryta on 'bromo malophthalic' acid (Baeyer, *A* 166, 355) Prisms (containing 2aq), v sol water — PbA" aq needles

Reference — OXYTEREPHTHALIC ACID

(a) OXY-ISOPHTHALIC ALDEHYDE

$\text{C}_6\text{H}_5(\text{OH})(\text{CHO})_2$ [431] [1087] Formed by boiling *p* oxy benzoic aldehyde with chloroform and NaOHAq , and also, together with the (β)-isomeride, from salicylic aldehyde in like manner (Voswinckel, B 15, 2021) Needles, sol. ether, sl sol water, almost insol ligroin

(β) Oxy-isophthalic aldehyde

$C_6H_3(OH)(CHO)_2$ [2 3 1] [88°] Made as above
Needles (from water), v sol ligroin Yields
c oxy isophthalic acid on fusion with potash

D1 oxy-isophthalic aldehyde

$C_6H_2(OH)_2(CHO)_2$ *Resorcylic dialdehyde*
[127*] Made by the action of chloroform and
NaOHaq on resorcin (Tiemann & Lewy, B 10,
2212, 13, 2368) Needles (from hot water)
With alcoholic aniline it yields yellow crystals
[199*] Phenyl hydrazine acetate forma
 $C_6H_2(OH)_2(CHO)_2$ [o 230*] (Rudolph, A 248, 105)
Yields a dioxim [209*] (Marcus, B 24, 3652)

Methyl derivatives

$C_6H_2(OMe)(OH)(CHO)_2$ Two isomerides, $[179^\circ]$ and $[89^\circ]$, are formed from $[1\ 3]$ $C_6H_4(OMe)(OH)$, chloroform, and $NaOHaq$

OXY-PHTHALIDE $C_8H_4O_4$, 16.

$C_6H_5(OH) \langle \begin{smallmatrix} CH_2 \\ CO \end{smallmatrix} \rangle O$ [222°] Got by reducing oxyphtalimide with tin and HCl, treating with nitrous acid, and heating the resulting nitroso oxyphtalidine with NaOH aq (Graebe & Rée, *C J* 49, 526) Prisms or needles, sl sol ether

OXY-PHTHALIMIDE $C_6H_4(OH) \begin{smallmatrix} \diagup CO \diagdown \\ \diagdown CO \diagup \end{smallmatrix} NH$

or $\text{C}_6\text{H}_3(\text{OH})\langle\frac{\text{C}(\text{NH})}{\text{CO}}\rangle\text{O}$ [290°] Formed by saturating α oxy phthalic acid with NH_3 (Réé, *A* 233, 232, *C J* 49, 525) Yellow crystals, sl. sol ether When reduced by tin and HCl it yields a product from which a nitroso compound [170°] may be prepared

OXY-DIPHTHALYL $C_{18}H_8O_5$, 2 e

$$\text{C}_6\text{H}_5 \begin{array}{c} \diagup \text{C} = \text{C} \diagdown \\ \diagdown \text{CO} \diagup \quad \diagdown \text{CO} \diagup \end{array} \text{C}_6\text{H}_4\text{OH} \quad [\text{above } 374^\circ]$$

Made by heating phthalide with 1-oxy phthalic anhydride and NaOAc at 200° (Graebe & Guye, *A* 233, 244) Needles (from HOAc)

A di-oxy-diphthalyl [250°] appears to be formed by dissolving di-chloro diphthalyl in alcoholic potash (Ador, *A* 164, 245)

DI-OXY-PROPANE v PROPYLENE GLYCOL

DI-OXY PROPANE TRI-CARBOXYLIC ACID

$C_6H_3(OH)_2(CO_2H)_2$. Formed by oxidation of isosaccharin with HNO_3 (Kiliani, *B* 18, 636). Colourless syrup. By HI it is reduced to glutaric acid $C_5H_7(CO_2H)_2$. — A''' . H_2Ca small prisms

An isomeric acid, formed by boiling chloro-
nitroic acid with lime, forms the salts $\text{Ca}_2\text{A}'''$, 9aq,
 $\text{Ba}_2\text{A}'''$, 5aq, and $\text{Cd}_2\text{C}_2\text{H}_3\text{O}_8$, 8aq (Pavloveck, *A.*
178, 157). The same acid, yielding $\text{Ca}_2\text{A}''$, 10aq,

occurs in beet root juice (Lippmann, *B* 16, 1078)

Tri-oxy-propane tri carboxylic acid $C_3H_4O_8$, *s.e.* $CO_2H \cdot CH(OH) \cdot CH(OH) \cdot C(OH)(CO_2H)_2$ Got by oxidising mannite with alkaline $KMnO_4$ (Pabst, *J* 1880, 611) Yields a soluble NH_4 salt

OXY-PROPANE PHOSPHONIC ACID
 $C_3H_5PO_4$, *s.e.* $CH_2(OH) \cdot PO(OH)_2$ [162°] Formed from propionic aldehyde and PCl_3 , followed by water (Fossek, *M* 7, 29) Plates, v sol water— CaA''

OXY-PROPANE SULPHONIC ACID
 $C_3H_5(OH)(SO_3H)$ Formed from *n* propyl alcohol and SO_3 , and also by boiling allyl alcohol with aqueous $KHSO_4$ (Max Müller, *B* 6, 1441) The same, or an isomeric acid, is got from propylene oxide and $NaHSO_4$ (Erlenmeyer, *A* 158, 280) The *K* salt crystallises in small needles

Oxy-propane disulphonic acid
 $CH(OH)(CH_2SO_3H)_2$ Formed by boiling glycerin dichlorhydrin with aqueous K_2SO_5 (Schauffelen, *A* 148, 111) and, in like manner, from epichlorhydrin (Pazschke, *J pr* [2] 1, 86, *Z* [2] 5, 613) Syrup.— K_2A'' 2aq trimetric octahedra $abc = 41 \cdot 48 \cdot 1$ — BaA' 2aq m sol water— PbA'' 2aq— AgA'' crystals, v sol alcohol

Reference—CHLORO-OXY-PROPANE SULPHONIC ACID

TRI-OXY-PROPENYL-BENZENE *Tri-methyl derivative*

[1 2 4 5] $C_9H_8(OMe)_3$, $CH \cdot CHMe$ This is the constitution of asaron (*q v*) (Eykmann, *B* 22, 3172).

OXY-PROPENYL-BENZOIC ACID
 $C_9H_8(OH)(OH)CO_2H$ [4 2 1] *Propenyl-salicylic acid* [146°] Formed from oxyisopropyl-salicylic acid $C_9H_8(CMe_2OH)(OH)CO_2H$, by elimination of H_2O by warming with dilute HCl White slender needles V sol. alcohol, ether, and CS_2 , sl sol water Sublimable Volatile with steam Gives a deep reddish-violet colouration with $FeCl_3$. By sodium amalgam it is reduced to oxy-cummic acid $C_9H_8 \cdot Pr(OH)CO_2H$ [4 2 1]

Salts— $A'Ag$ sparingly soluble white crystalline pp— A'_2Cu 2aq small green crystals, nearly insol water

Polymeride $(C_9H_8O)_x$ [230°] Formed by boiling oxy-propenyl benzoic acid with strong HCl Small white crystals M sol hot acetic acid, alcohol, and ether, insol water and CS_2 Gives a deep bluish-violet colouration with $FeCl_3$ Not volatile with steam— $(CuA') \frac{1}{2} aq$ — $(AgA')_2$ white pp (Heymann & Königs, *B* 19, 3318, 20, 2390)

OXY-PROPIONAMIDE

$CH_3 \cdot CH(OH) \cdot C(NH) \cdot NH_2$
The hydrochloride $B'HCl$ is formed from $(CH_3 \cdot CH(OH) \cdot C(NH) \cdot OC_2H_5) \cdot HCl$, and alcoholic NH_3 (Pinner, *B* 23, 2947) Needles, converted by $AgNO_3$ into $B'HNO_3$ [84°], v sol water

OXY-PROPIONIC ACID v LACTIC ACID AND HYDRAXYLIC ACID V. also BROMO-, CHLORO-, and IODO-, OXY-PROPIONIC ACIDS

Di-oxy-propionic acid v GLYCERIC, GLYXYLIC, PRUVIC, and GLYCIDIC ACIDS

***α*-OXY-PROPIONIC ORTHALDEHYDE** *Tri-methyl derivative* $C_3H_4O_5$, *s.e.*

$CH_3 \cdot CH(OMe) \cdot CH(OMe)_2$ (148°) *SG* 2 948 Formed by heating acrolein (2 vols) with methyl alcohol (6 vols) and $HOAc$ (1 vol) at 100° (Alsborg, *J* 1864, 495). Liquid, with pleasant smell, sl sol water.

Tri-ethyl derivative (186°) *SG* 13 90.

OXY-PROPIONIC IMIDO ISOAMYL ETHER
 $CH_3 \cdot CH(OH) \cdot C(NH) \cdot OC_4H_9$ The hydrochloride $B'HCl$ [69°], formed by the action of HCl gas on a solution of aldehyde cyanhydrin in isoamyl alcohol, crystallises in thin needles The homologous $CH_3 \cdot CH(OH) \cdot C(NH \cdot Cl) \cdot OC_4H_9$ also melts at 69° and gradually decomposes forming lactamide and ammonium chloride (Pinner, *B* 23, 2947)

OXYPROPYL-AMIDO BENZOIC ACID

$C_9H_8(CMe_2OH)(NH_2)CO_2H$ [4 2 1] [158°] Made by reduction of nitro oxypropyl-benzoic acid with $FeSO_4$ and ammonia (Widman, *B* 19, 271) Prisms Exhibits blue fluorescence in ethereal solution

Acetyl derivative

$C_9H_8(C_2H_5OH)(NHAc)CO_2H$ [174°] Trimetric tables, v sol hot alcohol

Oxypropyl amido-benzoic acid

$C_9H_8(CMe_2OH)(NH_2)(CO_2H)$ [4 3 1] Made by reducing the corresponding nitro acid (Widman, *B* 16, 2570, 17, 1305) Prisms, v sol alcohol, v sl sol ether With $ClCO_2Et$ it forms $C_9H_8(CMe_2OH)(NH \cdot CO_2Et) \cdot CO_2H$ [167°] and $C_9H_8 \cdot N_2O_2$ (above 300°) which is converted by H_2SO_4 into $CO(NH \cdot C_6H_5)(CMe_2OH)(CO_2H)_2$

Acetyl derivative Crystalline

***α* OXY PROPYLAMINE**

$CH_3 \cdot CH(OH) \cdot CH \cdot NH_2$ *Amido iso propyl alcohol* [30°] (176°) Formed from allylamine and H_2SO_4 , followed by water (Liebermann a Paal, *B* 16, 531) Needles

Benzoyl derivative

$CH_3 \cdot CH(OBz) \cdot CH_2 \cdot NH_2$ Got by boiling bromopropyl benzamide with $HBrAq$ Oil, v sol water Converted into $CH_3 \cdot CH(OBz) \cdot CH \cdot OH$ by nitrous acid Boiling $NaOHAq$ converts it into the isomeric $CH_3 \cdot CH(OH) \cdot CH_2 \cdot NHBz$ [93°] (Hirsch, *B* 23, 970)— $C_{10}H_{15}NO \cdot HBr$ [133°] Needles— $B'C_6H_5N_2O$, [189°] Needles— $B'_2H_2PtCl_6$, yellow needles (from water)

***β*-Oxy-propylamine** $CH_2(OH) \cdot CH_2 \cdot CH_2 \cdot NH_2$ Formed by heating bromo propyl phthalimide with dilute (1 2) H_2SO_4 for 5 hours at 200° (Gabriel, *B* 21, 2672)— $B'_2H_2PtCl_6$, yellow plates— $B'HAuCl_4$ plates

Sulphuric acid derivative

$CH_2(O \cdot SO_3H) \cdot CH_2 \cdot CH_2 \cdot NH_2$ [221°] Formed from *β* bromo propylamine hydrobromide and aqueous Ag_2SO_4 , at 100° (Gabriel a Lauer, *B* 23, 91) Prisms (from warm water) Indifferent body

Benzoyl derivative $NH_2 \cdot C_6H_5 \cdot OBz$

Liquid, v sol water Formed from phenyl-pentoxazoline and HBr (Gabriel a Elfeldt, *B* 24, 3216) Yields $B'HBz$ [135°], $B'_2H_2PtCl_6$, [205°] and $B'O_2C_6H_5N_2O$, [178°]

Phenyl derivative $PhO \cdot C_6H_5 \cdot NH_2$

(242°) Got from phenoxy propyl phthalimide acid and HCl (Lohmann, *B* 24, 2684) Yields $B'HCl$ [168°] crystallising in shining plates, and $PhO \cdot C_6H_5 \cdot NHBz$ [118°] Cyanic acid forms $PhO \cdot C_6H_5 \cdot NH \cdot CO \cdot NH_2$, [114°]

Oxy-propyl amine $(C_3H_7OH)(C_3H_7) \cdot NH$ (175°) [80°] *SG* 13 9018 Prepared by heating propyl-allyl amine with H_2SO_4 , and pouring the product into water (Liebermann a Paal, *B* 16, 581) Needles $B'_2H_2PtCl_6$ 2aq efflorescent

Oxy-tri-propyl amine $(C_3H_7OH)_3 \cdot N(C_3H_7)_2$ Got by heating di-propyl-allyl-amine with

H_2SO_4 , and pouring the product into water (L. a. P.) — $\text{B}^+\text{H}_2\text{PtCl}_6$

OXY PROPYL AMYL AMINE $\text{C}_9\text{H}_{19}\text{NO}$ (c. 200°) [c 2°] Formed by heating allyl amyl amine with H_2SO_4 , and pouring the product into water (Liebmann a. Baal, B 16, 531) Solidifies to long fine needles

Oxy propyl di isomyl-amine $(\text{C}_5\text{H}_{11})_2\text{N}(\text{C}_5\text{H}_9\text{OH})$ (243°) Formed from propylene chlorhydrin and di isomylamine (Lomse, A Ch [6] 13, 433) Oil, sl sol water Inactive to light — $\text{B}^+\text{H}_2\text{PtCl}_6$, orange crystals The acetyl and benzoyl derivatives form crystalline oxalates, the latter being $\text{C}_{20}\text{H}_{35}\text{NO}_8\text{H}_2\text{C}_2\text{O}_4$

OXY PROPYL-BENZENE v. **PROPYL PHENOL** and **PHENYL PROPYL ALCOHOL**.

Di-oxy propyl benzene $\text{CH}_2\text{CH}(\text{OH})\text{CH}(\text{OH})\text{C}_6\text{H}_5$, [53°] Made from phenyl propylene bromide by successive treatment with KOAc and alcoholic potash (Zincke, B 17, 709) Tables (from ether ligron), v e sol water A more sparingly soluble isomeride [93°], made from phenyl propylene bromide by boiling with aqueous K_2CO_3 , crystallises from ether in monoclinic tables

Tri-oxy propyl-benzene v. **PROPYL PYRO-GALLOL**

Tetra-oxy-propyl-benzene *Methylene derivative* $\text{CH}_2(\text{OH})\text{CH}(\text{OH})\text{CH}_2\text{C}_6\text{H}_4\text{CH}_2$, [83°] Formed from safrol and dilute KMnO_4 , at 75° (Tiemann, B 24, 2881) White needles, v. sol boiling water and ether Yields piperonal, piperonyl acid, and CH_2O , $\text{C}_6\text{H}_5\text{CH}$, CO_2H on further oxidation Phenyl cyanate forms CH_2O_2 , $\text{C}_6\text{H}_5\text{CH}$, $\text{C}_6\text{H}_5(\text{O CO NHPh})_2$ [127°]

Acetyl derivative CH_2O , $\text{C}_6\text{H}_5\text{CH}$, $\text{C}_6\text{H}_5(\text{OAc})_2$, (240° at 18 mm) **OXY PROPYL-BENZENE SULPHONIC ACID** Me $\text{C}(\text{OH})\text{C}_6\text{H}_4\text{SO}_3\text{H}$ Got from cumene p-sulphonic acid, KOH and KMnO_4 (R Meyer, A 219, 302)

Salts — KA' — BAa' , Splits off H_2O at 140° — PbA' , Splits off $2\text{H}_2\text{O}$ at 110°, probably forming lead propenyl benzene sulphonate

Reaction — PCl_5 followed by ammonia forms the amide of propenyl benzene sulphonic acid [152°]

OXY o-ISOPROPYL BENZOIC ACID $\text{OMe}_2(\text{OH})\text{C}_6\text{H}_4\text{CO}_2\text{H}$ The salt KA' [197°] is formed by the action of conc KOH aq on dimethyl-phthalide (Wischneus, A 248, 59) The free acid is unstable, at once forming di methyl-phthalide [68°]

Oxy-p isopropyl-benzoic acid $\text{OMe}_2(\text{OH})\text{C}_6\text{H}_4\text{CO}_2\text{H}$ [156°] Formed by oxidising cumic acid, or cymene, with alkaline KMnO_4 (R Meyer, B 11, 1283, 1790, A 219, 248, Remsen & Emerson, A C J 1, 267, Widman, B 19, 583) Thin triclinic prisms (from water), v sol alcohol and ether Gives no colour with FeCl_3 Yields terephthalic and acetyl benzoic acids on oxidation by CrO_3 Boiling HCl aq forms two isomeric propenyl-benzoic acids. — BAa' , aq — CaA' , $2\frac{1}{2}$ aq — CuA' , 3 aq — AgA' , 4 aq crystalline pp

An isomeric or identical acid is got by boiling bromo-propyl benzoic acid with alcoholic potash (Ozumpelk, B. 8, 478)

Isomerides v Oxy-CUMINIC ACID.

Di-oxy-isopropyl-benzoic acid $\text{C}_6\text{H}_4(\text{OMe}_2\text{OH})(\text{OH})\text{CO}_2\text{H}$ [1 2 4] [173°].

Formed by the action of nitrous acid on oxy-amido isopropyl benzoic acid (Widman, B 17, 722) Crystals (from water), v sol alcohol and ether Coloured dark brown by FeCl_3

Di oxy-isopropyl benzoic acid $\text{C}_6\text{H}_4(\text{OMe}_2\text{OH})(\text{OH})\text{CO}_2\text{H}$ [1 3 4] [130°-185°]. Formed by oxidising carvacryl sulphuric acid with alkaline KMnO_4 (Heymann a. Konigs, B. 19, 3310) Flat needles (from water), v e sol alcohol — CuA' , aq — AgA' needles

DI-OXY-DI-ISOPROPYL-DI CARBOXY DI-PHENYL-ALLOPHANIC ETHER $\text{C}_{24}\text{H}_{28}\text{N}_2\text{O}_8$, i.e. $\text{N}(\text{CO}_2\text{Et})\text{C}_6\text{H}_4(\text{OMe}_2\text{OH})\text{CO}_2\text{H}$ [above 300°]. $\text{CO NH C}_6\text{H}_4(\text{OMe}_2\text{OH})\text{CO}_2\text{H}$

Formed, with $\text{CO}(\text{NH C}_6\text{H}_4(\text{OMe}_2\text{OH})\text{CO}_2\text{H})_2$, and the action of ClCO_2Et on oxy amido-isopropyl benzoic acid (Widman, B 17, 1306) Tables (from HOAc), almost insol water

OXY-PROPYLENE-DIAMINE $\text{C}_3\text{H}_6\text{N}_2\text{O}$ + a. $\text{CH}(\text{OH})(\text{CH}_2\text{NH}_2)_2$ Formed from glycerin dichlorhydrin and alcoholic NH_3 (Claus, A 168, 36) — $\text{B}^+\text{H}_2\text{PtCl}_6$

OXY-PROPYLENE-TETRA METHYL-DI-AMINE $\text{C}_3\text{H}_8(\text{OH})(\text{NMe}_2)_2$, (170°-185°) Formed by heating s dichlorhydrin with NMe_2H (Berend, B 17, 510) Liquid, v sol water

Benzoyl derivative Crystalline

OXY PROPYL-ETHYL-AMINE v. **ETHYL-OXY-PROPYL AMINE**

OXY-PROPYL-MALONIC ACID

$\text{CO}_2\text{HCH}(\text{CH}_2\text{CHMeOH})\text{CO}_2\text{H}$ The free acid at once splits off water, leaving the lactonic acid The salts $\text{BaC}_4\text{H}_6\text{O}_5$, CaA' , and $\text{Ag}_2\text{A}'$ may, however, be prepared

Lactonic acid $\text{C}_4\text{H}_6\text{O}_5$ Got by combining allyl malonic acid with HBr , and boiling the product with water (Hjelt, B 15, 621, A 216, 53) Syrup, v sol water, sl sol ether Yields $\text{Ba}(\text{C}_4\text{H}_6\text{O}_5)_2$ crystallising in soluble plates

Di-oxy-propyl-malonic acid

$(\text{CO}_2\text{H})_2\text{CHCH}_2\text{CH}(\text{OH})\text{CH}_2\text{OH}$ The free acid in aqueous solution remains unchanged at 15°, but at 100° it splits off water, forming a lactonic acid whose barium salt is $(\text{C}_6\text{H}_6\text{O}_5)_2\text{Ba}$

Salts — BAa'' Got by boiling di-bromo-propyl malonic acid with baryta (Hjelt, A 216, 58) — $\text{Ag}_2\text{A}''$ flocculent pp

Di oxy di propyl-malonic acid

$(\text{CH}_2\text{CH}(\text{OH})\text{CH}_2)_2\text{C}(\text{CO}_2\text{H})_2$
Dilactone $\text{C}_8\text{H}_{10}\text{O}$, [106°] Obtained from di allyl malonic acid by evaporating with conc HBr aq (Hjelt) Thin plates (from alcohol), long needles (from water), or trimetric crystals (from conc HBr aq) $a:b:c = 61:1:94$ Warm baryta water forms $(\text{C}_8\text{H}_8\text{O}_4)_2\text{Ba}$, which on heating splits up into BaCO_3 , and the neutral lactone $\text{C}_8\text{H}_8\text{O}_4$ $\text{CH} < \text{CH}_2\text{CHMe} >$

Reference — **DI-BROMO-DI-OXY-DI-PROPYL-MALONIC ACID**

DI-OXY-ISOPROPYL-TRIMETHYLENE so-called $\text{CHPr} < \text{CH OH} > \text{CH OH}$? (126°) S 10 Formed

from isobutyric aldehyde and ethylene glycol by heating in a sealed tube (Lochert, B! [2] 46, 716) Liquid, lighter than water, v sol alcohol and ether Decomposed by water at 100° into its constituents. Bromine yields a heavy oil $\text{C}_8\text{H}_8\text{BrCH}(\text{CH OH})_2$ (c 187°).

α OXY-*p*-PROPYL-PHENYL-ACETIC ACID $C_9H_9O_3$, $\approx C_6H_4(C_2H_5)CH(OH)CO_2H$ [158°]. S 19 at 21°. Formed from cuminnic aldehyde, HCl, and HCl (Raab, *B* 8, 1148, Plöchl, *B* 14, 1816) Small needles (from water) — BaA_2 , aq — PbA_2 , — AgA_2 stellate needles

OXY ISOPROPYL-DIPHENYLENE-KETONE

CARBOXYLIC ACID $C_{11}H_{11}O_4$, \approx
 $CO < \begin{smallmatrix} C_6H_5 \\ C_6H_2(CMe_2OH) \end{smallmatrix} CO_2H$ [5 2]. [190°]

Formed by oxidising retene quinone with alkaline $KMnO_4$ (Bamberger & Hooker, *B* 18, 1080, 1750, A 229, 150) Yellow plates, sl sol cold water and ether, m sol alcohol — BaA_2 , aq — CuA_2 , — AgA_2 yellow flocculent pp

Oxam Not melted at 270°

TRI OXY-TRI-PROPYL-PHOSPHINE

Hydrate $PH(OH)(CH(OH)C_2H_5)_3$. Formed by the action of conc KOHAq upon $(C_2H_5O)_3PHI$, which is got, together with the crystalline oxypropyl iodide $(C_2H_5O)_3PI$ by dissolving PHI in propionic aldehyde at 0° (De Girard, *A Ch* [6] 2, 24) Syrup, sl sol water

OXY PROPYL PHTHALIMIDE $C_{11}H_9NO$, $\approx C_6H_4C_2O, NCH_2CH_2CH_2OH$ Formed from bromo propyl phthalimide and hot conc KOHAq (Gabriel & Lauer, *B* 23, 87) Needles (from Aq)

Phenyl derivative $C_{15}H_{11}O_2$, $C_6H_5NCH_2CH_2CH_2OPh$ [88°] Formed from the bromo propyl derivative of phenol and potassium phthalimide at 220° (Lohmann, *B* 24, 2633) Needles, converted by KOH into $PhO C_2H_5NH CO C_2H_5CO_2Et$ [134°] a white crystalline powder

OXY PROPYL-PIPERIDINE

$C_9H_{19}N$, C_2H_5OH So called '*piperpropylalkine*' (194°) SG \approx 947, Δ 936 VD 4 79 (obs) Made from piperidine and propylene chlorhydrin (Ladenburg, *B* 14, 1880, 2407, 15, 1144, Laun, *B* 17, 680) Liquid, sol water — B^+HAuCl_4 — $B^+H_2PtCl_6$ — Mandelate dilute HCl forms $C_9H_{19}NO_3$, which gives B^+HAuCl_4

Acetyl derivative $C_9H_{17}NO$ — B^+HAuCl_4

Benzoyl derivative $C_{11}H_{13}NO_2$, OBz — B^+HAuCl_4 — $B^+C_2H_5N_2O$, yellow powder

Oxy-propyl-piperidine

$C_9H_{19}N$, $CH_2CH(OH)CH_2NH$ So called '*(α)-pipercolyl methylalkine*' [47°] (225°) Got by reducing the corresponding oxypropyl pyridine with sodium and alcohol (Ladenburg, *B* 22, 2588) Crystalline, v sol water, alcohol, and ether Yields an oily nitrosamine — $B^+H_2PtCl_6$, [149°] Small crystals (from alcohol)

Oxy-propyl-piperidine $C_9H_{19}N$, $CH_2CH(OH)EtNH$ [100°] Got by reducing (α) pyridyl ethyl ketone in amyl alcohol with sodium (Engler & Bauer, *B* 24, 2533) Needles Probably identical with the ψ conhydrin in *Conium maculatum*

Dioxypropyl-piperidine $C_8H_{11}NO_2$ Made by heating piperidine with glycerin chlorhydrin (Roth, *B* 15, 1150) Silky plates — B^+HBr — B^+HAuCl_4 , yellow needles

OXY-PROPYL-PYRIDINE C_8H_9NO , \approx

$C_6H_4(CH_2CH_2CH_2OH)N$ So called '*(α) isovalalkine*' (\approx 130° at 17 mm) Formed from (α) ethyl pyridine and formic aldehyde (Ladenburg & Adam, *B* 24, 1878) V sol water. — B^+HAuCl_4 , [71°] — $B^+H_2PtCl_6$, [142°]

Hexahydrate $C_8H_{11}NO$ — B^+HBr

Oxypropyl-pyridine

C_8H_9N , $CH_2CH(OH)CH_2N$ '*(α)-procolylmethylalkine*' (176°–181° at 18 mm.) Made from

(α) methyl-pyridine and acetic aldehyde (*L*) Liquid, sl sol water — $B^+H_2PtCl_6$, [189°] Small tables, sl sol water

Oxy propyl-pyridine C_8H_9N , $CH_2CH(OH)EtN$ (215°) Got, together with corvine and another body [69°], by reducing (α)-pyridyl ethyl ketone with sodium amalgam at 30°–40° (Engler & Bauer, *B* 24, 2532) — $B^+H_2PtCl_6$

Oxy-propyl-pyridine Tetrahydrate
 $CH_2 < \begin{smallmatrix} CH_2(C_6H_5) \\ CH_2 \end{smallmatrix} CO > NH$ [57°] (274°)

Made by heating δ amido- α propyl valeric acid to 200° (Aschan, *B* 23, 3701) Flat snowy needles

OXY - PROPYL - PYRIDINE DIHYDRIDE

CARBOXYLIC ACID ν MORRHUIC ACID

(*Py* 3) **OXY-(B 3)-ISOPROPYL-QUINOLINE**

$CH CH C CH CH$ Cumostyryl [169°] Made

by boiling the hydrochloride of phenyl o amido cumyl acrylic acid with water and a few drops of HCl for 5 hours (Widman, *B* 19, 264) Needles, v sol hot alcohol, sl sol hot water

Oxy - n - propyl - quinoline Dihydrate

$CH CH C CH_2CH_2$ [134°] Formed, by intra

molecular change, by reducing the preceding body, and also from nitro n cumyl propionic acid (Widman, *B* 19, 2778) Prisms (from benzene ligroin), insol water

OXY-PROPYL SUCCINIC ACID Lactonic

$acid CH_2CH < \begin{smallmatrix} CH_2CH_2CO_2H \\ O-CO \end{smallmatrix}$ [69°]

(\approx 260°) Formed from allyl succinic acid and conc HBrAq (Hjelt, *B* 16, 334) Crystals (from alcohol)

OXY ISOPROPYL-SULPHOBENZOIC ACID

$CMe_2(OH)C_6H_4(SO_3H)CO_2H$ [13 4] Made by oxidising cymene sulphonic acid, an intramolecular change taking place Got also from iso cymene sulphonic acid and $KMnO_4$ (R Meyer & Boner, *A* 220, 8, 30, *B* 13, 1495, 14, 2391, Remsen, *Am* 8, 262) — KA'' 5aq Triclinic crystals, $a b c = 675.1 \ 542$, $\alpha = 131^\circ 14'$, $\beta = 104^\circ 26'$, $\gamma = 66^\circ 27'$ — KA'' 2aq — BaA'' aq An isomeric acid, got from m isocymene sulphonic acid, yields BaA'' and PbA''

OXYPROPYL *p*-TOLUIDINE $C_{10}H_{11}NO$, \approx

$C_6H_4MeNHCH_2CH_2OH$ [74°] (293° cor) Formed from propylene oxide and *p* toluidine (Morley, *C J* 41, 387, *B* 15, 179) Needles (from light petroleum), insol water, v sol benzene and ether — $B^+H_2C_2O_4$, [151°] Pearly plates

(α) **OXY-PYRIDINE** C_8H_9NO , \approx

$N < \begin{smallmatrix} (OH)CH \\ CH=CH \end{smallmatrix} CH$ [107°] VD 2 8 (obs)

Formed by distilling its carboxylic acids (Königs, *B* 16, 2160, 17, 590, 2391, 19, 2463, Pechmann, *B* 17, 2384, Weidel & Strache, *M* 7, 297) Dimetric needles, v e sol water and alcohol Coloured red by $FeCl_3$ Yields a di bromo-oxy pyridine [207°]

Tetrahydrate C_8H_9NO , \approx

$NH < \begin{smallmatrix} COCH_2 \\ CH_2CH_2 \end{smallmatrix} CH_2$ [40°] (255°) Made by

distilling δ amido valeric acid (Schotten, *B* 21, 2285, Gabriel, *B* 23, 1770) Crystalline mass. Yields an acetyl derivative (298°)

(β) **Oxy-pyridine** C_8H_9NO , \approx

$N < \begin{smallmatrix} CH(OH) \\ CH=CH \end{smallmatrix} CH$ [124 5°] Formed by

potash fusion from pyridine sulphonie acid (Fischer a Renout, *B* 17, 763, 1896) Needles, v sol water and alcohol Coloured red by FeCl₃. —Oxalate [175°] White needles

Acetyl derivative (210° uncor) Oil

Ethyl derivative Made by ethylation, and also from bromo pyridine and alcoholic potash (Weidel a Blau, *M* 6, 664) Oil — B₂H₂PtCl₆ [192°] Prisms

Methyl iodide B'MeI Needles

Methylo-chloride B'MeCl Needles — B₂Me₂PtCl₆ Orange prisms, sl sol alcohol

(γ) *Oxy pyridine* NH $\begin{smallmatrix} \text{CH} \\ \text{CH} \end{smallmatrix}$ CO *Pyrid.*

one [148°] (above 350°) S 100 at 15° Formed by heating chelidamic acid under reduced pressure at 230° or with water at 196° (Lerch, *M* 5, 402, Hartinger a Lieben, *M* 6, 300) Formed also by distilling its carboxylic acid (Ost, *J pr* [2] 29, 65) Small efflorescent grains (from alcohol), prisms (containing aq), or trimetric tables, *abc* = 55 1 15 V sol water and alcohol Melts below 100° when hydrated Converted by treatment with MeI and KOH into NMe $\begin{smallmatrix} \text{CH} \\ \text{CH} \end{smallmatrix}$ CO, a deliquescent mass [89°]

which gives (C₅H₅NO).H₂PtCl₆ aq [176°], B'MeI, and B Me₂PtCl₆ Does not yield an acetyl derivative

Salts — B₂H₂PtCl₆ aq — B' H₂PtCl₆ 2aq Efflorescent monoclinic needles [200°], converted by boiling water into B'₂H₂PtCl₆ aq — B'HNO₃AgNO₃ tables — B'HgCl₂ — B'HHg₂Cl₄

Methyl derivative N $\begin{smallmatrix} \text{CH} \\ \text{CH} \end{smallmatrix}$ C OMe

(191° cor) at 738 mm Formed from (γ) chloro-pyridine and NaOMe (H a L) Liquid, sol water Alkaline in reaction Changes at 220° into the crystalline isomeride (*v supra*) Gives a deep blue liquid with aqueous CuSO₄ Conc HIAq converts it into (γ) oxy pyridine — B'₂H₂PtCl₆ crystals, sl sol water

Di-oxy-pyridine C₅H₂(OH)₂N [c 255°] (K a G), [239°] (W a B) Formed by potash fusion from its ethyl derivatives and also from pyridine disulphonic acid (Königs a Geigy, *B* 17, 1835, Weidel a Blau, *M* 6, 651) Needles (containing 4aq), v sol water, sl sol alcohol Coloured red by FeCl₃ — B'HCl needles

Mono ethyl derivative C₅H₄NO [128°] Formed, together with the di ethyl derivative, by heating dibromopyridine [111°] with alcoholic potash Tables, sl sol cold water — B'HNO₃ — B'₂H₂PtCl₆ red triclinic needles

Di ethyl derivative C₅H₄(OEt)₂N (c 244°) Made as above Liquid, nearly insol water — B'₂H₂PtCl₆ yellow needles — B'HHg₂Cl₄ [106°] Crystals (from HClAq)

Di-oxy-pyridine *Di oxim of the dihydroxide* NH $\begin{smallmatrix} \text{C(OH)CH}_2 \\ \text{C(OH)CH}_2 \end{smallmatrix}$ CH₂ [193°] A product of the action of hydroxylamine on trimethylene cyanide (Biedermann, *B* 22, 2967) Yields a di acetyl derivative [127°] and a dibenzoyl derivative [180°]

Picrate [175°] Needles

Tri-oxy-pyridine C₅H₂NO₃ a.s

N $\begin{smallmatrix} \text{C(OH)CH}_2 \\ \text{C(OH)CH}_2 \end{smallmatrix}$ C OH [220°-230°]. Formed by heating di-oxy amido pyridine (glutazine)

with conc HClAq (Stokes a Von Pechmann, *Am* 8, 384, *B* 19, 2701) Yellowish sandy powder, v sol hot water On evaporation of its solution it is partly converted into its anhydride. FeCl₃ gives a red colour Forms with bromine CBr₄ CO CBr₂ CONH₂ NH₄OAc at 140° converts it into glutazine — BaA' — AgA' — B'HCl

Oxim NH $\begin{smallmatrix} \text{CO CH}_2 \\ \text{CO CH}_2 \end{smallmatrix}$ C NOH [196°].

Made by boiling tri oxy pyridine or glutazine with hydroxylamine hydrochloride Minute hexagonal plates (containing aq), m sol hot water — B'HCl plates

Phenyl-hydrazide

NH $\begin{smallmatrix} \text{CO CH}_2 \\ \text{CO CH}_2 \end{smallmatrix}$ C N₂HPh [230°] Tables

Anhydride C₅H₂N₂O₂ Made by boiling glutazine with dilute H₂SO₄ Minute flesh coloured prisms, sl sol water — BaA'₂ 4aq yellow prisms — AgHA' — B'H₂SO₄ — B'HCl

References — Di bromo, Di chloro, and Di iodo, oxy PYRIDINE

OXY-PYRIDINE CARBOXYLIC ACID

C₅H₃N(OH)CO₂H (a) *Oxypicolinic acid* [267°] Made by heating di chloro oxy pyridine carb oxylie acid [282°] with HI in HOAc at 210° (Ost, *J pr* [2] 27, 289) Long needles (containing aq) or short anhydrous needles, v sol hot water and alcohol, insol ether Coloured reddish brown by FeCl₃ AgNO₃ is not reduced, but gives a white pp — BaA'₂ aq — CaA'₂ — C₅H₃N(OK)CO₂K aq groups of needles

Oxy pyridine carboxylic acid

C₅H₃N(OH)CO₂H (b) *Piccolinic acid* [250°] Made by the action of HI in HOAc on chloro oxy pyridine carboxylic acid [257°] at 200° (Ost) Formed also without by products by boiling comanic acid C₅H₄O₂(CO H) with NH₄Aq (Ost, *J pr* [2] 29, 64) Glittering plates Yields oxypyridine [148°] on heating strongly — Salt BaA'₂ 2aq small needles, m sol water

Oxy pyridine carboxylic acid

C₅H₃N(OH)CO₂H (γ) *Oxy picolinic acid* [258°] Formed by the action of tin and HClAq on chloro oxy pyridine carboxylic acid [224°] (Bellmann, *J pr* [2] 29, 7) Small trimetric pyramids (containing aq), sl sol water, sol conc HClAq Coloured brown by FeCl₃ — BaA'₂ prisms — CaA'₂ 4aq needles

Oxy-pyridine carboxylic acid

C₅H₃N(OH)CO₂H ϵ C(OH) $\begin{smallmatrix} \text{N} \\ \text{CH} \end{smallmatrix}$ C CO₂H *Oxy nicotinic acid* [302°] Formed by heating oxy-pyridine di carboxylic (oxyquynolinic) acid with water at 195° (Königs a Geigy, *B* 17, 589) Formed also by the action of NH₄Aq on the methyl ether of coumalic acid (*v vol u p* 264), the product being saponified (Pechmann a Welsh, *B* 17, 2384, *C J* 47, 145) Needles, sl sol hot water May be sublimed PCl₅ yields chloro pyridine carboxylic acid, whence tin and HCl form nicotinic acid FeCl₃ gives a light yellow colour — PbA'₂ 2½aq needles, sl hot water

Methyl derivative C₅H₃N(OMe)CO₂H [238°] Formed by methylating the acid, and also from methyl coumalate and methylamine, the product being saponified Needles (containing aq), nearly insol cold water

Phenyl derivative C₅H₃N(OPh)CO₂H.

[280°] Formed by the action of boiling NaOHAq on methyl coumal anilide (*loc cit*) Needles

Oxy-pyridine carboxylic acid

$\text{CH} \begin{smallmatrix} \text{N} \text{C(OH)} \\ \text{CH} \text{CH} \end{smallmatrix} > \text{C CO}_2\text{H}$ (a) *Oxy-nicotinic acid*

[256°] Formed by heating (a)-oxy isocinchomeronic acid with HOAc and a little Ac₂O at 210° (Weidel a Strache, *M* 7, 295) Slender needles (from water) Yields (a) oxy pyridine when heated —AgA' silky needles

Oxy-pyridine dicarboxylic acid C₈H₅NO₄, *z* e

$\text{N} \begin{smallmatrix} \text{C(CO}_2\text{H)} \text{CH} \\ \text{C(OH)} \text{C(CO}_2\text{H)} \end{smallmatrix} > \text{CH}$ (a) *Oxyisocinchomeronic acid*

Formed by oxidising (a) diquinolyl with KMnO₄ (Weidel a Strache, *M* 7, 293) Vitreous crystals Gives no colour with FeCl₃ —BaA' —AgA'' needles, nearly insol water

Oxy-pyridine dicarboxylic acid

C₈H₅N(OH)(CO₂H)₂, *z* e

$\text{N} \begin{smallmatrix} \text{C(CO}_2\text{H)} \text{C(CO}_2\text{H)} \\ \text{C(OH)} \text{CH} \end{smallmatrix} > \text{CH}$ *Oxyquinolinic acid*

Formed from quinolinic acid by potash-fusion (Königs a Körner, *B* 16, 2158) Small crystals (from dilute H₂SO₄), blackening at 254° FeCl₃ colours its aqueous solution red Its Ag salt yields (a) oxy-pyridine on heating —BaA' 4aq needles (from hot water)

Methyl derivative C₈H₅N(OMe)(CO₂H)₂

[140°] Got by adding KMnO₄ to an aqueous solution of the methyl derivative of (γ) amidocarboxystyryl (Feer a Königs, *B* 18, 2398) Needles, *v* e sol water —AgH₂A''₂ needles (from water)

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Di-oxy-pyridine carboxylic acid C₈H₅NO₄, *z* e

$\text{N} \begin{smallmatrix} \text{C(OH)} \text{CH} \\ \text{C(OH)} \text{CH} \end{smallmatrix} > \text{C CO}_2\text{H}$ *Citraconic acid* Formed by heating the mono-, di-, or tri- amide of citric acid with HCl or H₂SO₄ (Behrmann a Hofmann, *B* 17, 2687) Crystalline powder, nearly insol water, sl sol hot HClAq Carbonises at 300° PCl₅ converts it into di chloro-pyridine carboxylic acid [210°] Gives a deep blue colour with NaNO₂. Tin and HCl reduce it to tricarballic acid —BaA', 2aq

Di-acetyl derivative Crystalline

Methyl ether MeA' Plates, decomposing above 220°

Ethyl ether EtA' Plates

Amide C₈H₅N(OH)₂CONH₂ Formed by the action of conc NH₃Aq on ethyl acetyl citrate and on acetic ether (Ruhemann, *O* J 51, 405, *B* 20, 8368) Small grey crystals (from water)

Di-oxy-pyridine carboxylic acid

C₈H₅N(OH)₂CO₂H *Comenamic acid* *Di oxy-pyridine carboxylic acid* Formed by heating hydrogen ammonium comenamate at 190°, or by boiling comenic acid with NH₃Aq The yield is about 45 p c of the comenic acid used (How, *T* *E* 20 [2] 255, *A* 80, 65, 83, 350, Ost, *J* pr [2] 27, 269) Tables (containing 2aq), m sol hot water and alcohol Not decomposed by boiling NaOHAq Gives a purple colour with FeCl₃

Reactions —1 Yields pyridine on distillation with zinc-dust (Lieben a Hattinger, *B* 16, 1263) 2 Conc HIAq at 200° does not attack it, but when heated with it for two days at 270° pyrocomenic acid (probably a di oxy-pyridine) C₈H₅NO₄ is formed This body crystallises in needles (containing aq), gives a violet colour with FeCl₃, and forms B' HBr —8 PCl₅ (8 mols) acting on the acid (1 mol) at 100° forms a product

which on treatment with tin and HClAq yields a di-oxy-methyl-pyridine C₈H₅NO₄, crystallising in trimetric prisms (containing aq), and forming the salts B' HCl and B' I₃PO₄ PCl₅ and POCl₃ at 200° convert this tri-oxy methyl pyridine at 200° into hexa chloro methyl pyridine and C₈H₅Cl₆N(CCl₃), which is converted by boiling water into chloro (γ) oxy pyridine (a) carboxylic acid (Bellmann, *J* pr [2] 29, 19) Excess of PCl₅ (5 mols) at 220° acting on comenamic acid in presence of POCl₃ forms penta and hexa chloro-methyl pyridine, and other bodies, whence water produces chloro (γ)-oxy pyridine (a) carboxylic acid and chloro cyamic acid C₈H₅NO₄, which crystallises in needles [186°], and gives a blue colour with FeCl₃ Chlorocymamic acid yields the salts AgA' and BaA', aq —4 Ammonium comenamate forms, among other products, on distillation, a very poisonous base called 'Oxycomazine' C₈H₅N₂O (Krippendorff, *J* pr [2] 32, 163) The base crystallises from alcohol in four sided prisms, *S* 0035 at 20° Its solutions in dilute acids show green fluorescence, and in strong acids a blue fluorescence Tin and HCl reduce it to oxy amido pyridine Oxycomazine forms the following salts B'H₂Cl₂ [c 265°], B'H₂PtCl₆, B'H₂SO₄, 8aq, [c 295°], and C₈H₅AgN₂O —5 KMnO₄ oxidises comenamic acid to tri oxy pyridine carboxylic acid

Salts —NH₄A' very small grains —BaC₈H₅NO₄, aq pp —BaA', 2aq crystalline

Ethyl ether EtA' [205°] Needles (containing aq), sol hot water (Reibstein, *J* pr [2] 24, 284) Yields Ba(C₈H₅NO₄)₂, 2aq and EtA' HClAq, both crystalline When heated with AcCl it yields an anhydride C₈H₅NO₃ [261°] and two derivatives, C₈H₅N(OH)(OAc)CO₂Et [152°] and C₈H₅N(OAc)₂CO₂Et [38°] BrCl forms C₈H₅N(OBz)₂CO₂Et [102°]

Di-oxy-pyridine carboxylic acid C₈H₅NO₄ *Oxymido comenic acid* Made from comenic acid and hydroxylamine (Ost, *J* pr [2] 29, 378) Small needles, decomposing at 200° Reduced by tin and HCl to (8) oxy picolinic acid

Di oxy pyridine dicarboxylic acid *Ethyl derivative* $\text{N} \begin{smallmatrix} \text{C(OEt)} \text{C(CO}_2\text{Et)} \\ \text{C(OH)} \text{C(CO}_2\text{H)} \end{smallmatrix} > \text{CH}$ [182°]

Formed by the action of NaOHAq on the mono-ethyl ether EtHA'' [160°], which is made by treating ethoxy (a) pyrone dicarboxylic ether with NH₃Aq (Guthzeit a Dressel, *B* 22, 1427, *A* 262, 104) Needles (containing aq) Conc HClAq at 140° forms glutamic acid [184°] PCl₅ in POCl₃ at 250° gives di chloro pyridine dicarboxylic acid [230°] whence EtA'' [76°] and, by treatment with HI, pyridine dicarboxylic acid [322°] may be prepared —AgA''

Mono-ethyl ether

$\text{OH} \begin{smallmatrix} \text{C(OEt)} \text{C(CO}_2\text{Et)} \\ \text{C(OH)} \text{C(CO}_2\text{H)} \end{smallmatrix} > \text{N}$ [160°]. Needles Yields AgA' and an acetyl derivative C₈H₅NO₄ [100°].

Di-ethyl ether EtA' [81°]

Tri-oxy-pyridine carboxylic acid C₈H₅NO₄ *Tri-oxy-picolinic acid* *Oxy comenamic acid*. Formed, in small quantity, by oxidising comenic acid with potassium permanganate and H₂SO₄ in the cold Prepared by heating oxycomenic acid C₈H₅O₄(OH)₂CO₂H with conc NH₃Aq at 180° (Reibstein, *J* pr [2] 24, 290; Ost, *J* pr [2] 27, 265) Small needles (contain-

ing aq) (from water). Does not form a hydrochloride. Gives an azure pp with BaCl_2 and NH_4 . Alcoholic potash gives, in alcoholic solutions, a blue flocculent pp, forming a colourless solution in water. FeCl_3 gives an indigo blue colour. AgNO_3 gives a white pp, soon turning black. Br forms bromo tri oxy pyridine carb oxylic acid, crystallising with 2aq. Nitric acid added to the ethereal solution forms oxypyridoquinone carboxylic acid $\text{C}_8\text{HNO}_2(\text{OH})\text{CO}_2\text{H}$, crystallising in orange tables (containing 2aq).

References.—Bromo, Chloro, and Bromonitro-, OXY PYRIDINE CARBOXYLIC ACID

OXY PYRIDYL ETHYL FURFURANE

$\text{CH} \begin{smallmatrix} \text{CH CH} \\ \text{CH N} \end{smallmatrix} > \text{C CH}_2 \text{CH}(\text{OH}) \text{C} \begin{smallmatrix} \text{CH CH} \\ \text{OH} \end{smallmatrix} [\text{43}^\circ]$. (164° at 20 mm). Formed from (a) methylpyridine, furfuraldehyde, and some water at 150° (Klein, B 23, 2693). Greenish mass, v sol alcohol.— $\text{B}^+\text{H}_2\text{PtCl}_6$ [162°].— B^+HHgCl_4 [c 150°].— B^+HCl_4 .— $\text{B}^+\text{C}_6\text{H}_5\text{N}_3\text{O}$, [c 160°]

Acetyl derivative Oil

$(\text{C}_{11}\text{H}_{10}\text{AcNO}_2)_2 \cdot \text{H}_2\text{PtCl}_6$ [165°].— B^+HHgCl_4 [c 155°]. Small needles

Benzoyl derivative $\text{C}_8\text{H}_{10}\text{BzNO}_2$ [49°].— $\text{B}^+\text{H}_2\text{PtCl}_6$ [140°–145°].— B^+HHgCl_4

OXY PYRIDYL-MALONIC ACID

$\text{C}_8\text{H}_8\text{N C}(\text{OH})(\text{CO}_2\text{H})_2$. Formed by oxidising pilocarpine with KMnO_4 (Hardy & Calmels, Bl [2] 48, 228). Syrup. Yields pyridine (β) carb oxylic acid on further oxidation.— BaA'' 3aq.— $\text{Cu}_2\text{A}''(\text{OH})$ aq.— $\text{Ag}_2\text{A}''$ pp

OXY-PYRIDYL-PHENYL-PROPIONIC ACID

Hexahydrate $\text{C}_8\text{H}_{10}\text{N CHPh CH}(\text{OH}) \text{CO}_2\text{H}$ [244°]. Formed from piperidine and sodium phenyl glycidate (Erlenmeyer, B 22, 1482)

α -OXY (β)-PYRIDYL PROPIONIC ACID

$\text{C}_8\text{H}_8\text{N CH}_2\text{CH}(\text{OH})\text{CO}_2\text{H}$. Formed by boiling pilocarpine with water for 12 hours (Hardy & Calmels, Bl [2] 48, 227). Gummy mass.— B^+HCl — B^+PtCl_6 .— B^+AuCl_3 [154°]. Prismatic needles

a Oxy-(α)-pyridyl propionic acid $\text{C}_8\text{H}_8\text{N CH}_2\text{CH}(\text{OH})\text{CO}_2\text{H}$ [125°]. Formed by the action of hot dilute caustic soda on $\text{C}_8\text{H}_8\text{N CH}_2\text{CH}(\text{OH})\text{CCl}_2$, the product of the union of chloral with (a) picoline (Einhorn, B 23, 219, A 265, 211). Prisms.— $\text{Cu}_2\text{A}''\text{O}$ needles.— $(\text{HA}')\text{H}_2\text{PtCl}_6$ [204°].— $\text{HA}'\text{HAuCl}_4$ [174°]. Orange prisms.— AgA' white needles.— $\text{HA}'\text{HCl}$ [86°].— $\text{HA}'\text{HBr}$

Benzoyl derivative [145°]. Needles.— $\text{B}^+\text{H}_2\text{PtCl}_6$ [179°]. Yellow prisms

Methylether MeA' — $\text{MeA}'\text{HAuCl}_4$ [119°]

—Benzoyl derivative of the ether $\text{C}_8\text{H}_8\text{N CH}_2\text{CH}(\text{OBz})\text{CO}_2\text{Me}$ [c 41°]. Yields $\text{B}^+\text{H}_2\text{PtCl}_6$ [193°], crystallising in yellow needles

β Oxy-(α) pyridyl propionic acid

$\text{C}_8\text{H}_8\text{N CH}_2\text{CH}(\text{OH})\text{CH}_2\text{CO}_2\text{H}$ [86°]. Formed by warming β bromo pyridyl propionic acid with NaOH aq (Einhorn, B 23, 221). White needles, v e sol water.— $\text{Cu}_2\text{A}''\text{O}$ blue crystals.— B^+HCl [147°]. Prisms.— $\text{B}^+\text{H}_2\text{PtCl}_6$ [191°].

Benzoyl derivative [185.5°]. Prisms

Methylethyl MeA' Yields $(\text{MeA}')\text{H}_2\text{PtCl}_6$ [178.5°] and a benzoyl derivative [79°] crystallising in prisms

Ethyl ether EtA' Yields $(\text{EtA}')\text{H}_2\text{PtCl}_6$

Di-oxy-pyridyl-propionic acid

$\text{C}_8\text{H}_8\text{N CH}(\text{OH})\text{CH}(\text{OH})\text{CO}_2\text{H}$ [190°]. Formed by oxidising pyridyl-acrylic acid with alkaline KMnO_4 (Einhorn, B 23, 223). White crystals

Ethyl ether EtA' [96°]. Tables. Yields a benzoyl derivative [122°] crystallising in needles

DI-OXY-PYRIMIDINE o HYDROQUINONE TETRACARBOXYLIC ACID

OXY-PYRONE DICARBOXYLIC ACID

Ethyl derivative of the ethyl ether

$\text{CH} \begin{smallmatrix} \text{C}(\text{CO}_2\text{Et}) \\ \text{C}(\text{CO}_2\text{Et}) \end{smallmatrix} > \text{CO}$ Anhydride of tri-

ethyl propylene-tetracarboxylate [94°]. Formed by distilling di carboxy-glutaconic ether at 210° under 15 mm pressure (Guthzeit & Dressel, B 22, 1415). Needles, insol water and alcohol. Converted by HClAq into glutaconic acid

OXY-PYROTARTARIC ACID $\text{C}_8\text{H}_6\text{O}_4$ t.s

$\text{CH}_2\text{C}(\text{OH})(\text{CO}_2\text{H})\text{CH}_2\text{CO}_2\text{H}$ Mol w 148 [108°]. Formed by the action of boiling dilute HCl upon $\text{CH}_2\text{C}(\text{OH})(\text{CN})\text{CH}_2\text{CO}_2\text{Et}$, which is got by heating acetoacetic ether for three days with dry HCy at 100° (Morris, C J 37, 7, c Demarçay, Bl [2] 27, 120). Made also by oxidation of isovaleric acid by long boiling with dilute HNO_3 (Bredt, B 14, 1782, 15, 2318). Deliquescent, star like groups of needles, sol water, alcohol, and ether. On dry distillation it splits up into water and citraconic anhydride— BaA'' 2aq. Not decomposed by boiling with water— CaA'' 1½aq.— AgA'' ½aq needles

Oxy-pyrotartaric acid

$\text{CH}_2\text{CH}(\text{CO}_2\text{H})\text{CH}(\text{OH})\text{CO}_2\text{H}$ Citramalic acid [119°]. Formed by the action of zinc on a dilute solution of chloro citramalic acid (which melts at 130° according to Melikoff, A 253, 88). HCl is added towards the end of the reaction (Carius, A 129, 160, Morawski, Sitz W 76 [2] 670, J pr [2] 10, 69). Large hygroscopic crystals, yielding citraconic anhydride and water on distillation—Salts KA'' 2aq.— BaHA'' 2aq.— MgA'' — CaA'' 2aq.— CaA'' 1½aq.— CaHA'' ½aq.— ZnA'' 2aq.— PbA'' 3½aq.— AgA''

Oxy pyrotartaric acid

$\text{CH}(\text{OH})\text{CH}(\text{CO}_2\text{H})\text{CH}_2\text{CO}_2\text{H}$ Itamalic acid

Formed from itaconic acid by successive treatment with HBr and hot water. Formed also by boiling ita chloro pyrotartaric acid with aqueous NaCO_3 (Swarts, Bull Acad Belg [2] 24, 25, Bl [2] 9, 317, Fittig, A 188, 76, Morris, C J 37, 14). The free acid, liberated from its Ca salt by oxalic acid, or from its Ag salt by H_2S , changes on evaporation, even at 15°, into its lactone [58°].— NaA'' — $(\text{NH}_4)\text{HA}''$ — CaA'' aq.— CaA'' 3aq.— PbA'' — CuA'' — $\text{Cu}_2\text{A}''\text{O}$ — AgA'' — EtA' oil

Lactone $\text{CH}(\text{CO}_2\text{H}) \begin{smallmatrix} \text{CH}_2 \\ \text{CO} \end{smallmatrix}$ Paraconic

acid [58°]. Formed as above, and also by boiling ita bromo pyrotartaric acid (1 pt) with water (10 pts) (Beer, A 216, 90). Crystalline. Yields citraconic anhydride on distillation. With bases it yields salts of itamalic acid— $\text{NaC}_8\text{H}_6\text{O}_4$ — CaA'' 3aq. Small needles. Yields calcium itamale on boiling with CaCO_3 — AgA''

Chloro-itamalic acid $\text{C}_8\text{H}_6\text{ClO}_4$ [150°]. Made by passing chlorine into a solution of sodium itaconate. Crystals, v e sol water

Oxypyrotartaric acid [c 135°] described by Maxwell Simpson (Pr 13, 44) as got from glycerin diolrhhydrin by successive treatments with KCy and KOH , is probably β oxy glutaric acid. It yields $\text{Ag}_2\text{A}''$ and EtA' (298°). An isomeric or identical acid, made by fusing

sulpho-pyrotartaric acid with potash, yields $\text{Ag}_2\text{A}'$ aq (Wieland, *A* 157, 41) A liquid isomeride, got by potash fusion from bromo-eyano-butyric acid, yields $\text{Ag}_2\text{A}'$

Di-oxy pyrotartaric acid $\text{C}_4\text{H}_4\text{O}_6$ *Citratar-taric acid* Formed by the action of boiling baryta water on chloro citramalic acid prepared from barium citraconate and HOCl (Carius, *A* 129, 159) and by heating oxycitraconates with water at 120° (Morawski, *J pr* [2] 11, 432) Amorphous, deliquescent mass — $\text{Pb}_2\text{A}'$ aq

Isomeride v ITATARTARIC ACID

DI-OXY-DI-PYRRYL-BUTANE $\text{C}_{12}\text{H}_{16}\text{N}_4\text{O}_2$ $\text{C}_6\text{H}_7\text{N CMe(OH) CMe(OH) C}_6\text{H}_7\text{N}$ [120°] Formed from pyrrol methyl ketone, water, and sodium amalgam (Dennstedt a Zimmermann, *B* 19, 2204) Monoclinic prisms (containing 2aq) Melts at 98° when hydrated v e sol alcohol

OXY PYRUVIC ACID $\text{C}_3\text{H}_4\text{O}_5$ $\text{CH}_2\text{(OH) CO CO}_2\text{H}$ Formed by dissolving 'nitro cellulose' (collodion) in dilute NaOHAq and allowing the solution to stand at 20° (Will, *B* 24, 405) Amorphous, v sol water, but ppd by alcohol Reduces Fehling's solution and ammoniacal AgNO_3 Its solutions, and those of its salts, are slightly laevogyrate Phenyl hydrazine forms the compound $\text{C}_{11}\text{H}_{11}\text{N}_3\text{O}_2$ or $(\text{N.HPh})\text{CH C(N.HPh) CO}_2\text{H}$, [205°], whence NaA' [231°], KA' [233°], $\text{NH}_4\text{A}'$ [200°], CaA' , and EtA' [149°]

Salts — CaA' , 8aq — SrA' , 4aq — CdA' , 4aq

OXY QUINALDINE v OXY METHYL QUIN-

OLINE

OXY QUINAZOLINE *Dihydrate*

$\text{C}_8\text{H}_6\text{N}_2\text{O}_2$ CH_2NH NH CO 'Phenylidihydroacimazine' [160°] Formed from oxy tolyl urea and HCl (Soderbaum a Widman, *B* 22, 1669) Scales, insol cold conc KOH aq — B'HCl — $\text{B}'_2\text{H}_2\text{PtCl}_2$, 2aq [205°] — B'H aq , [179°]

Oxy-quinazoline $\text{C}_8\text{H}_6\text{N}_2\text{O}$ C(OH)N N=CH [212°]

Made by heating formyl o amido benzamide [123°] for two hours at 180° (Knappe, *J pr* [2] 43, 214) Thin needles — $\text{B}'_2\text{H}_2\text{PtCl}_2$ aq [above 250°]

Methyl derivative [71°].

Di oxy quinazoline

$\text{C}_8\text{H}_6\text{N}_2\text{O}_2$ C(OH)NH N=COH 'Uramido benzoyl' [above 350°] Formed by passing cyanogen into an alcoholic solution of o amido benzoic acid and boiling the product with HCl aq Formed also by fusing o amido benzoic acid or o amido benzamide with urea, and by heating the product of the action of ClCO_2Et on o amido benzamide (Griess, *B* 2, 415, 11, 1985, *Abt*, *J pr* [2] 89, 140) Needles, sl sol hot water With PCl_5 it yields di chloro quinazoline [115°] — $\text{C}_8\text{H}_6\text{N}_2\text{O}_2\text{EtOH}$ needles

Di-methyl derivative $\text{C}_8\text{H}_8\text{(OMe)}_2\text{N}_2$, [66°] Made from di chloro quinazoline and NaOMe Needles, v e sol alcohol

OXY p-QUINAZOLYL-BENZOIC ACID

$\text{C}_{12}\text{H}_8\text{N}_2\text{O}_4$ N CH $\text{CO N C}_6\text{H}_4\text{ CO}_2\text{H}$ Made by oxidising p-tolyl quinazoline dihydrate with KMnO_4 (Paal a Busch, *B* 22, 2699) Small needles — AgA'

OXY-QUINHYDRONE $\text{C}_8\text{H}_8\text{N}_2\text{O}_2$ Formed from oxyhydroquinone and HNO_3 (Barth a Schreder, *M* 5, 595) Dark greyish-blue crystals.

(*Py* 1)-**OXY-QUINOLINE** $\text{C}_8\text{H}_7\text{N}$ CH CO CH NH CH

[235°] Made by heating phenyl- β amido acrylic acid at 200° or oxanilic acid at 170° (Reissert, *B* 20, 3109, *B* 21, 1876) Lup needles (from alcohol) Yields quinoline when distilled with zinc dust

Acetyl derivative [228°] Needles

Phenyl hydrazide $\text{C}_{15}\text{H}_{11}\text{N}_3$, [168°]

(*Py* 2)-**Oxy-quinoline** This is probably the constitution of cynurine v *infra*

(*Py* 3) **Oxy-quinoline** $\text{C}_8\text{H}_7\text{N}$ CH CH N=COH

Carbostyryl [199°]

Formation — 1 By reducing o nitro cinnamic acid (Chiozza, *A* 83, 118, Tiemann, *B* 13, 2070, Friedlander, *B* 14, 1916) — 2 By heating o amido cinnamic acid with HCl aq (T) or dilute H_2SO_4 (Feer a Königs, *B* 18, 2395) — 3 By reducing tri chloro-oxy quinoline with HI — 4 By heating (*Py* 3) chloro quinoline with water at 120° (Friedlander a Ostermaier, *B* 15, 335) — 5 By the action of aqueous HOCl upon quinoline (Erlenmeyer a Rosenhek, *B* 18, 3295) — 6 By heating quinoline on the water bath with a conc solution of bleaching powder (E a R, *B* 19, 489, Roos, *B* 21, 619)

Properties — Long thin feathery crystals (containing aq) (from water) or thick anhydrous prisms (from alcohol), v sl sol cold water, insol NH_4Aq

Salts — $\text{Ba(C}_2\text{H}_3\text{NO)}_2$ plates — AgA' pp

Methyl ether MeA' (247°) Oil

Ethyl ether EtA' (256°) Formed from (*Py* 3) chloro quinoline and KOEt Formed also by heating o amido cinnamic ether with alcohol and ZnCl_2 at 90° (Friedlander a Weinberg, *B* 15, 1424, 2103) and by ethylation of carbostyryl Pungent oil, solidifying below 0° Yields a dihydrate [199°] when reduced by sodium amalgam

Phenyl ether [69°] Plates

Dihydrate v AMIDO PHENYL PROPIONIC ACID

(*B* 1)-**Oxy-quinoline** CH C(OH) C CH CH $\text{CH CH} - \text{C N CH}$

[224°] Formed by potash fusion from quinoline (*Py* 1) sulphonic acid (Riemerschmied, *B* 16, 721, Lellmann, *B* 20, 2174) Formed also from (*Py* 1)-amido-quinoline by the diazo reaction (Skraup, *M* 5, 533) Silky needles or plates, sol alcohol and aqueous Na_2CO_3 , v sl sol water — Salts BHCl yellow needles — $\text{B}'_2\text{H}_2\text{PtCl}_2$, 4aq orange tables

Tetrahydrate $\text{C}_8\text{H}_8\text{(OH)}_4\text{N}_2$ CH_2 NH CH_2

[117°] Made by reducing with tin and HCl Needles, sol water, alcohol, and ether Yields a nitrosamine crystallising in tables, sol alcohol

(*B* 2) **Oxy-quinoline** C(OH) CH C CH CH $\text{CH CH} - \text{C N CH}$

[193°] (above 360°) Formed by heating a mixture of p amido-phenol, p nitro phenol, glycerin, and H_2SO_4 (Skraup, *B* 15, 893, *M* 3, 545) Formed also by heating its carboxylic acids (Weidel, *M* 2, 575, Skraup, *M* 4, 696) and by potash-fusion from its sulphonic acid (Fischer, *B* 17, 440) Small prisms (from alcohol) Not coloured by ferric chloride solution $\text{B}'_2\text{H}_2\text{PtCl}_2$, 2aq [236°] — $\text{B}'_2\text{Cu(OAc)}_2$ — $\text{B}'_2(\text{H}_2\text{SO}_4)_2$, 11aq — B'HCl aq prisms, v e sol.

water — $B'MeIaq$ Crystalline (Claus & Howitz, *J pr* [2] 42, 232, 43, 520) — $B'MeCl$ [c 272°] — $B'Me_2PtCl_2$ — $B'Me_2SO, 5aq$ — $B'MeOH$ [c 200°] — $B'EtBr$ [c 242°] — $B'C_2H_5Cl 1\frac{1}{2}aq$ [237°] — $(B'C_2H_5Cl)_2PtCl_2$

Methyl ether MeA' p Quinansole (305°) $SG \pm 1.665$ Got by methylation, and also from p-amisidine, nitro anisole, glycerin, and H_2SO_4 (Skraup, *M* 6, 762) Oil Solutions of its salts show blue fluorescence Gives a green colour with chlorine water and ammonia — $B'HCl 2aq$ — $B'_2H_2PtCl_4, 4aq$ — $B'_2H_2SO_4$ — $B'_2H_2Cr_2O_7$ — $B'MeI$ [235°] Prisms — **Picrate** [204°]

Acetyl derivative $C_6H_5(OAc)N$ [38°] (298°) Crystals — $B'_2H_2PtCl_4$ Prisms

Benzoyl derivative [231°] Needles

Tetrahydride of the methyl ether $C_6H_5(OMe)N$ **Thallin** [43°] (283°) at 735 mm Got by reducing the methyl ether with tin and conc $HClAq$ Prisms $FeCl_3$ gives a golden colour, changing to emerald green — Chlorine water gives a green colour turned yellow by ammonia — $B'HCl$ — $B'_2H_2SO_4, 2aq$ — $B'HI$ [155°] — $B'C_2H_5O_2$ four sided prisms S 10 at 15° — **Picrate** [162°]

Acetyl derivative $C_6H_5Ac(OMe)N$ [47°]

(B 3) Oxy quinoline $\begin{matrix} CH & CH & - & C & CH & CH \\ & C(OH) & CH & C & - & N & CH \end{matrix}$ [c 238°] Formed by heating *m*-nitro phenol with *m* amido phenol, glycerin, and H_2SO_4 (Skraup, *B* 15, 893, *M* 3, 559) Formed also by potash fusion from quinoline (B 3) sulphonic acid (Fischer, *B* 15, 1979) Silky needles, sol alcohol, not volatile with steam Its solutions show green fluorescence $FeCl_3$ gives a brownish red colour — $B'_2H_2PtCl_4, 2aq$ — $B'HCl 1\frac{1}{2}aq$ prisms — $B'Cu(OAc)_2$ — **Picrate** [244°]

Benzoyl derivative [86°] Prisms

Methyl ether C_6H_5NO (275° at 720 mm) Oil, volatile with steam

(B 4) Oxy quinoline $\begin{matrix} CH & CH & - & C & CH & CH \\ & CH & C(OH) & C & - & N & CH \end{matrix}$

[75°] (267° cor)

Formation — 1 By distilling its carboxylic acid (Weidel & Cobenzl, *M* 1, 862) — 2 By soda fusion from its sulphonic acid (Bedall & Fischer, *B* 14, 443, 1366) — 3 By heating *o* amido phenol with *o* nitro phenol, glycerin, and H_2SO_4 (Skraup, *B* 15, 893, *M* 3, 536)

Properties — Prisms, sl sol water May be distilled with steam $FeCl_3$ gives a green colour Gives quinolinic acid on oxidation with $KMnO_4$ (Fischer & Renouf, *B* 17, 756) Chlorine in $HOAc$ forms mono, di-, and tri, chloro derivatives (Zincke & Hebebrand, *A* 264, 198) Ethylene chlorhydrin forms crystalline $B'(C_2H_5OH)Cl$, whence $(C_6H_4NO_2Cl)_2PtCl_2$ (Wurtz, *C R* 96, 1269) $ClCO_2Et$ forms C_2H_5NO , [105°] whence $(C_6H_4NO_2)_2H_2PtCl_4$ (Lippmann, *M* 8, 439) According to Lippmann (*M* 10, 667), MeI in $MeOH$ at 100° forms $C_6H_5(OMe)N(C_6H_5(OH)NMeI)HI 2aq$, whence $C_6H_5ClN_2O_2HCl 5aq$ and $C_{20}H_{12}Cl_4N_2O_2PtCl_4 2aq$ These bodies may perhaps be more simply formulated, as below Chloroform and Na yield $CH(C_6H_5(OH)N)$. (Lippmann, *B* 19, 2471)

Salts — $B'HCl aq$ — $B'_2H_2PtCl_4, 2aq$ golden needles — $B'_2H_2SO_4, 2aq$ — $B'_2C_2H_5N_2O_2$ [204°] $Cu(C_6H_5NO)_2$ canary-yellow pp — $B'MeI aq$ [c 170°] — $B'MeCl 2aq$ [c 260°] — $B'_2Me_2PtCl_4, 2aq$

Acetyl derivative $C_6H_5(OAc)N$ (280°).

Oil — $B'_2H_2PtCl_4, 2aq$ yellow plates

Benzoyl derivative [120°] Crystals

Methyl ether $C_6H_5(OMe)N$ (268°) Got by methylation, and also from *o* amido anisole by Skraup's reaction Oil — $B'_2H_2PtCl_4, 2aq$ — $B'HCl$ — $B'_2C_2H_5N_2O_2$, yellow needles or plates — $B'MeI aq$ [160°] Leaflets (Claus & Howitz, *J pr* [2] 42, 229)

Ethyl ether $C_6H_5(OEt)N$ (286°) at 718 mm Needles (Fischer & Renouf, *B* 17, 759) — **Picrate** [181°] Yellow needles

Tetrahydride $C_6H_5(OH)C_6H_5N$ [122°]

Made by reducing (B 4)-oxy-quinoline with $SnCl_4$ (Fischer, *B* 14, 1368, 14, 2571, 16, 713, 17, 759) Needles or prisms, sol hot water Yields a nitrosamine [68°] The methyl ether $C_6H_5(OMe)N$ is oily and yields a crystalline hydrochloride and a nitrosamine [80°] The ethyl ether is also liquid (275°) at 715 mm, and forms a crystalline nitrosamine [113°] and an oily acetyl derivative (307°)

Oxy quinoline C_6H_5NO **Cynurine** [201°] above 300° S 477 at 15° Probably (Py 1)- or (Py 2) oxy quinoline Formed by heating its carboxylic acid (cynurenic acid) (Schmiedeberg & Schultzen, *A* 164, 158, Kretschy, *M* 2, 68) Formed also by oxidising cinchonine or cinchonine acid (Skraup, *M* 9, 821, 10, 729) Monoclinic prisms (containing 3aq), m sol hot water Melts at 52° when hydrated Tastes bitter $KMnO_4$ oxidises it to cynuric acid ICl gives a brownish pp [275°] (Dittmar, *B* 18, 1618) Distillation with zinc dust forms quinoline Ac_2O on heating forms an indigo blue dye Yields a tetrahydride — $B'_2H_2Cl 2aq$ monoclinic prisms — $B'HCl aq$ — $B'_2H_2PtCl_4, 2aq$ orange needles

(Py 2, 3) **Di oxy-quinoline**

$C_6H_5 \begin{matrix} < CH & C(OH) \\ & N=C(OH) \end{matrix}$ **β -Oxy-carbostyryl** [above

300°] Prepared by heating (Py 2, 3) chloro oxy quinoline (8 chloro carbostyryl) with fused KOH at 200° (Friedlander & Weinberg, *B* 15, 2681) Fine colourless needles Maybe sublimed Is a very weak base but a strong acid, it dissolves in concentrated HCl , but is reprecipitated on dilution By PCl_5 it is converted into the di chloro-quinoline [104°] — **A'Ag** crystalline

Di-oxy-quinoline $C_6H_5 \begin{matrix} < C(OH) & CH \\ & N=C & OH \end{matrix}$ [above

320°] **Formation** — 1 By the action of conc H_2SO_4 on *o* amido phenyl propionic acid (Baeyer & Bloem, *B* 15, 2151) — 2 By potash fusion from (Py 1, 3) bromo oxy quinoline (Friedlander & Weinberg, *B* 15, 2683) — 3 By reducing *o* nitro benzoyl malonic ether with tin and HCl (Bischoff, *B* 22, 387, *A* 251, 377) — 4 By boiling its carboxylic acid with conc $HClAq$ (B)

Properties — Needles, sol Na_2CO_3 and in a mixture of alcohol and $HClAq$, insol ordinary menstrua Its ammoniacal solution turns blue in air PCl_5 yields di-chloro-quinoline [67°]

Salt — $C_6H_5AgNO_2$ needles

Ethyl ether $C_6H_5 \begin{matrix} < C(OH) & CH \\ & N=C(OEt) \end{matrix}$ [228°].

Formed by reducing *o* nitro benzoyl malonic ether with tin and HCl (B) Slender needles

Dihydride $C_6H_5 \begin{matrix} < CH(OH) & CH \\ & N=C(OH) \end{matrix}$ [149°]

Formed by reducing *o*-nitro- β -oxy phenyl propi-

(containing aq), sl sol hot water, m sol hot alcohol. Gives a green colour with FeCl_3 . Yields (B 4) oxy quinoline on distillation, and pyridine (a) tri carboxylic acid on oxidation— BaA' — $\text{BaC}_{10}\text{H}_7\text{NO}_3$, aq— HgA' , aq— AgA' — $\text{HA}'\text{HCl}$ monoclinic needles— $\text{B}_2\text{H}_2\text{PtCl}_6$, 2aq needles

(B 2) Oxy quinoline carboxylic acid $\text{C}_{10}\text{H}_7\text{NO}_3$, (B) Oxy cinchonic acid [c 320°]. Made by potash fusion from (B) sulpho cinchonic acid (Weidel, *M* 2, 571). Tables (containing aq), sl sol water. Yields (B 2) oxy quinoline on distillation, and a pyridine tricarboxylic acid on oxidation— BaA' — $\text{HA}'\text{HCl}$ aq needles—(H) H_2PtCl_6 , 2aq monoclinic tables, decomposed by water

(B 3) Oxy quinoline (Py 1?) carboxylic acid $\text{C}_{10}\text{H}_7\text{NO}_3$, Xanthoquinic acid [above 300°]. Possibly identical with the preceding acid. Got by heating quinic acid with conc HCl aq at 225° (Skraup, *M* 2, 601, 4, 695). Yellow grains. Yields (B 2) oxy quinoline on distillation—Salts BaA' , 6aq— CaA' , 10aq— CuA' , aq— AgA' , 2aq— $\text{HA}'\text{HCl}$ 2aq— HA' , H_2PtCl_6 , 6aq— H H H SO_3 , 3aq golden prisms

Methyl derivative $\text{C}_{10}\text{H}_7(\text{OMe})(\text{CO}_2\text{H})\text{N}$ Quinic acid [280°]. Made by oxidising quinine or cinchonine with chromic acid (Skraup, *M* 2, 589). Thin yellowish prisms, sl sol hot water and hot alcohol, nearly insol ether. Its alcoholic solution shows blue fluorescence, destroyed by H_2SO_4 . KMnO_4 oxidises it to pyridine tri carboxylic acid—Salts BaA' , 4aq— CaA' , 2aq— CuA' , 1 $\frac{1}{2}$ aq— AgA' pulverulent pp $\text{HA}'\text{HCl}$ 2aq triclinic tables— $\text{H}_2\text{A}'$, H PtCl_6 , 4aq yellow crystals

(B 4) Oxy quinoline carboxylic acid $\text{C}_{10}\text{H}_7\text{NO}_3$, [280°]. Made by boiling o oxy quinoline with CCl_4 , water, KOH and alcohol (Lippmann a Fleissner, *B* 19, 2467, *M* 8, 318). Minute prisms, v sl sol hot water. Yields (B 4) oxy quinoline on distillation, and pyridine dicarboxylic (quinolime) acid [235°] on oxidation. FeCl_3 gives a green colour. Yields a di bromo derivative [193°]— $\text{BaC}_{10}\text{H}_7\text{NO}_3$, aq needles— AgHA' , (dried at 105°). Minute needles

Tetrahydride $\text{C}_{10}\text{H}_7\text{NO}_3$, [265°]. Got by reduction with tin and HCl . Prisms, sl sol water, almost insol ether. Reduces AgNO_3 in the cold. Gives a red colour with FeCl_3 . PtI yields $\text{C}_{10}\text{H}_7\text{NO}_3\text{EtNO}_3\text{HI}$, whence $\text{C}_{10}\text{H}_7\text{NO}_3$, [220°] may be got. Nitrous acid forms a nitrosamine [195°]— $(\text{C}_{10}\text{H}_7\text{NO}_3)\text{HCl}$ aq needles— B' H_2SO_4 , 3aq— $\text{B}'\text{HOAc}$ pp

(B 4) Oxy quinoline carboxylic acid $\text{C}_{10}\text{H}_7\text{NO}_3$, aq [250°]. Got from o oxy quinoline dithiocarboxylic acid $\text{C}_{10}\text{H}_7(\text{OH})(\text{CSH})\text{N}$ by warming with lead acetate and KOH aq (Lippmann a Fleissner, *M* 9, 300). Silky needles, sol water. Coloured red by FeCl_3 . Yields o oxy quinoline on distillation— KA' — BaA' , (dried at 180°)— AgA' — HgA' , Cl_2 — HA' , H_2PtCl_6 , 4aq— $\text{HA}'\text{HCl}$ 2 $\frac{1}{2}$ aq trimetric crystals

Tetrahydride $\text{C}_{10}\text{H}_7\text{NO}_3$, [222°]. Crystalline powder— $\text{B}'\text{HCl}$ needles, v sol water

(B 4) Oxy quinoline carboxylic acid $\text{C}_{10}\text{H}_7(\text{OH})(\text{CO}_2\text{H})\text{N}$. Formed by heating sodium o-oxy quinoline with liquid CO_2 in a closed vessel at 150° (Schmitt a Engelmann, *B* 20, 1217, 2690). Small yellow prisms (containing aq), m sol hot water and hot alcohol. Coloured red

by FeCl_3 . At 150° it splits up into CO_2 and o-oxy quinoline. Yields $\text{C}_{10}\text{H}_7\text{BrNO}_3$, [235°]

Salts— $\text{B}'\text{HCl}$ needles— $\text{B}'\text{HNO}_3$ — $\text{NH}_4\text{A}'$ aq— BaA' , 2aq needles, sl sol water— $\text{BaC}_{10}\text{H}_7\text{NO}_3$, amorphous, v sl sol water— AgA' amorphous powder

Phenyl ether PhA' [226°]. Made by heating the acid with phenol and POCl_3 at 170°. Prisms

Tetrahydride $\text{C}_{10}\text{H}_7(\text{OH})(\text{CO}_2\text{H})\text{N}$. Colourless prisms— $\text{B}'\text{HCl}$ prisms. MeI and MeOH at 100° form $\text{C}_{10}\text{H}_7\text{Me}(\text{OH})(\text{CO}_2\text{H})\text{N}$ [211°], which crystallises with 2aq

(B 2) Oxy quinoline carboxylic acid $\text{C}_{10}\text{H}_7(\text{OH})(\text{CO}_2\text{H})\text{N}$ [204°]. Formed by boiling p oxy quinoline with NaOH , CCl_4 , water, and alcohol (Lippmann a Fleissner, *M* 8, 324). Made also by heating potassium (not sodium) p oxy quinoline with liquid CO at 170° (Schmitt a Altschul, *B* 20, 2695). Minute prisms, v sl sol hot water and alcohol. Splits up at 200° into CO_2 and p oxy quinoline. Yields quinolime acid on oxidation— $\text{HA}'\text{HCl}$ — $\text{HA}'\text{HNO}_3$ needles— H A' , H_2PtCl_6 , 2aq— $\text{NH}_4\text{A}'$, $\frac{1}{2}$ aq needles, v sol hot water— BaA' , 2aq— CaA' , 6aq— $\text{PbC}_{10}\text{H}_7\text{NO}_3$, aq— CaA' , 6aq small needles— AgA'

Oxy quinoline carboxylic acid $\text{C}_{10}\text{H}_7\text{NO}_3$, Cynurenic acid [258°]. S 9 at 100°. Occurs in the urine of dogs after a fat diet (Liebig, *A* 8b, 125, 108, 354. Voit a Richter, *J* 1865, 676, Schmiedeberg a Schultzen, *A* 164, 155, Hofmeister, *H* 5, 70), or a diet of flesh only (Kietzschy, *M* 2, 57, 5, 16). Prisms (containing aq) insol cold water. Decomposed by heat into CO and oxy quinoline. Yields quinoline on distillation with zinc dust. Evaporation with KClO_4 and HCl leaves a residue which is turned emerald green by ammonia (Jaffé, *H* 7, 399)— $\text{HA}'\text{HCl}$. Decomposed by water (Bruegel, *H* 4, 92)— $\text{NH}_4\text{A}'$ — KA' , 2aq— BaA' , 4 $\frac{1}{2}$ aq— BaA' , 3aq— CaA' , 2aq— CuA' , 2aq— AgA' aq thick white pp

(Py 3) Oxy quinoline (B 3) carboxylic acid. Dihydride $\begin{array}{c} \text{CH CH} \\ | \quad | \\ \text{C}(\text{CO}_2\text{H})\text{CH C NH CO} \end{array}$ [above 280°]. Prepared by the reduction of [3 1 4] $\text{C}_{10}\text{H}_7(\text{NO}_2)(\text{CO}_2\text{H})\text{CH CH}_2\text{CO}_2\text{H}$ with ammonia and FeSO_4 (Widman, *B* 22, 2274). Yellow plates (from water), v sl sol alcohol. Yields a methyl ether MeA' [192°] crystallising in tables

Di-oxy-quinoline carboxylic acid Ethyl derivative of the ethyl ether

$\text{C}_6\text{H}_5\text{N} \begin{array}{c} \text{C}(\text{OH})\text{C CO}_2\text{Et} \\ \text{C OEt} \end{array}$ [107°]. Got by the action of zinc, alcohol, and gaseous HCl on o-nitro benzoyl malonic ether (Bischoff, *B* 22, 386). Small needles. Coloured violet by FeCl_3 .

Tetra oxy quinoline carboxylic acid Lactone of the di methyl derivative of the dihydride $\text{C}_{12}\text{H}_{11}\text{NO}_5$, etc

$\text{C}_6\text{H}(\text{OMe}) \begin{array}{c} \text{CH CH}_2 \\ | \quad | \\ \text{CO O} \end{array}$ NH CO [c 256°]. Prepared by

the reduction of o nitro meconin acetic acid

$\text{C}_6\text{H}(\text{NO}_2)(\text{OMe}) \begin{array}{c} \text{CO O} \\ | \quad | \\ \text{CH CH}_2 \text{CO}_2\text{H} \end{array}$ with tin and

HCl (Liebermann a Kleemann, *B* 19, 2296). Colourless needles (from water), v sol alcohol. Boiling baryta water yields $\text{Ba}(\text{C}_{10}\text{H}_7\text{NO}_3)_2$, 6aq.

oxidation by KMnO_4 . — NaA ' 8aq — AgA '. yellow pp — H_2A , H_2PtCl_6 5aq

B Oxy-(Py 8)-quinolyl-propionic acid

$\text{C}_6\text{H}_4 \begin{smallmatrix} \text{CH} & \text{CH} \\ \diagdown & \diagup \\ \text{N}=\text{O} & \text{CF}(\text{OH})\text{CH}_2\text{CO}_2\text{H} \end{smallmatrix}$ [176°]. Got from its amide, or by the action of Na_2CO_3 aq on bromo quinolyl propionic acid in the cold (Einhorn, *A* 246, 176) Colourless prisms, v sol alcohol and HOAc, insol chloroform. — NaA ' — AgA ' — HA ' HCl [188°] White prisms — H_2A , H_2PtCl_6 [218°] Yellowish red prisms
Methyl ether MeA' [62°] Prisms
Amide [152°] Made by dissolving the hydrobromide of bromo quinolyl propionic acid in ammonia in the cold White crystals (from alcohol)

Lactone $(\text{C}_6\text{H}_4\text{N}) \text{CH} \begin{smallmatrix} \diagup \text{O} \\ \diagdown \text{CH}_2 \end{smallmatrix} \text{CO}$ [82°]

Made by adding an equivalent quantity of Na_2CO_3 to the hydrobromide of bromo quinolyl propionic acid suspended in water (Einhorn, *A* 246, 169) Needles — $\text{C}_6\text{H}_4\text{NO}_2\text{HCl}$ [138°] — $\text{B C}_6\text{H}_4\text{N}_2\text{O}$, Golden plates (from alcohol)

OXY-QUINONE *Methyl ether*

$\text{C}_6\text{H}_4\text{O}_2(\text{OMe})$ [140°] Prepared by oxidation of *o*-anisidine with $\text{K}_2\text{Cr}_2\text{O}_7$ and dilute H_2SO_4 (Muhlhaner, *B* 13, 323, *A* 207, 251, Will, *B* 21, 605) Got in like manner from the methyl ether of amido resorcin (Bechhold, *B* 22, 2381) Yellow needles, with pleasant smell, sol alcohol, m sol ether and water Conc H_2SO_4 forms a deep blue solution The vapour colours filter paper red Reduced by SO_2 to $\text{C}_6\text{H}_4(\text{OH})_2(\text{OMe})$ With aniline it forms $\text{C}_6\text{H}_4(\text{NHPh})_2(\text{OMe})_2$, crystallising in coppery needles, and forming a dark blue solution in H_2SO_4 (Schweitzer, *C* 1888, 1434) *o*-Toluidine, *o*-xyldine, and diphenylamine form corresponding bodies melting at 239°, 228°, and 120° respectively

Ethyl ether $\text{C}_6\text{H}_4\text{O}_2(\text{OEt})$ [117°] Made by oxidation of $\text{C}_6\text{H}_4(\text{NH}_2)(\text{OEt})$, with $\text{K}_2\text{Cr}_2\text{O}_7$ and dilute H_2SO_4 at 15° (Will a Pukall, *B* 20, 1128) Yellow needles (by sublimation), m sol warm water, decomposed by hot water

Dioxy-quinone $\text{C}_6\text{H}_4(\text{OH})_2\text{O}$ [5 2 4 1]

Formation —1 By boiling the basic sodium salt of dioxyquinone dicarboxylic acid with HCl or H_2SO_4 (Loewy, *B* 19, 2387) —2 From di-amido-resorcin by oxidation to di-imido resorcin and treatment of this body with dilute (10 p c) KOH at 70° (Nietzki, *B* 21, 2374, Böninger, *B* 22, 1298) —3 By heating $\text{C}_6\text{H}_4(\text{NHPH})(\text{OH})_2$ or tetra-methyl di-amido-quinone with KOHAq (Kehrmann, *B* 23, 904)

Properties —Dark yellow needles, almost insol cold water, v sol alcohol. Its alkaline solutions are red. Not melted at 180° May be sublimed Forms a dioxim Conc HNO_3 forms nitranilic acid — $\text{Na}_2\text{C}_6\text{H}_4\text{O}_4$ — $\text{BaC}_6\text{H}_4\text{O}_4$ aq bluish black needles

Di-methyl ether MeA' [c 220°] Obtained by methylation Prepared also from $\text{C}_6\text{H}_4(\text{NO}_2)_2(\text{OMe})$, by reduction followed by oxidation with FeCl_3 (Nietzki a Reehberg, *B* 23, 1216) Got also by oxidation of acetyl-di methyl pyrogallol $\text{C}_6\text{H}_4(\text{OMe})(\text{OAc})$ (Hofmann, *B* 11, 332) Yellow needles Yields, on reduction, colourless $\text{C}_6\text{H}_4\text{O}_2$ [166°]

Di-ethyl ether $\text{C}_6\text{H}_4(\text{OEt})_2$ [183°] Got by oxidising the di ethyl ether of di-amido-hydroquinone with FeCl_3 (N a. B.) Sulphur-

yellow plates (from water) Yields a crystalline dioxim $\text{C}_6\text{H}_4(\text{OEt})_2(\text{NOH})_2$, which may be reduced by SnCl_2 to $\text{C}_6\text{H}_4(\text{OEt})_2(\text{NH}_2)_2$

Di-oxy-quinone *Di-methyl ether* $\text{C}_6\text{H}_4(\text{OMe})_2$ [249°] A product of the oxidation of $\text{C}_6\text{H}_4(\text{OMe})_2$ [1 2 3] by nitric acid (Will, *B* 21, 608) Prisms, v sol hot HOAc May be sublimed Reduced by SnCl_2 to $\text{C}_6\text{H}_4(\text{OH})_2(\text{OMe})_2$ [158°] Gives $\text{C}_6\text{Br}_2(\text{OMe})_2$ [176°]

Tri-oxy-quinone $\text{C}_6\text{H}_4(\text{OH})_3$ Made from tri amido resorcin by the action of FeCl_3 , the resulting amido-di-imido-resorcin being heated with HClAq at 150° (Merz a Zetter, *B* 12, 2035) Nearly black powder, insol water, sl sol alcohol Yields a crystalline tri acetyl derivative. — BaA''' , nearly black pp — PhA''' , — AgA''' , black pp

Tetra-oxy-quinone $\text{C}_6(\text{OH})_4\text{O}_2$ *'Dehydro-carboxylic acid'* Formed by the action of alcohol, air, and HCl on the black mass containing $\text{C}_6(\text{OK})_2$ got by combination of potassium with CO (Lerch, *A* 124, 20) Formed also by atmospheric oxidation of a solution of hexa-oxy benzene (Nietzki a Benckser, *B* 18, 507, 1836, 1855) It is also a product of the action of HNO_3 on mosite (Maquenne, *A Ch* [6] 12, 112) Steel-blue monoclinic needles and plates, v sol alcohol and hot water, sl sol ether Oxidised in alkaline solution by the air to croconic acid. Nitric acid forms C_6O . Aniline forms the compound $\text{C}_6(\text{OH})_4\text{O}_2(\text{NPh})(\text{NHPh})$, crystallising in red needles, with green lustre Phenylene-*o*-diamine yields $\text{C}_6\text{H}_4\text{N}_2\text{O}_2$, sol HClAq (Kehrmann, *B* 23, 2448) *o*-Tolylene diamine forms black crystals of $\text{C}_6\text{O}(\text{OH})_2 \begin{smallmatrix} \diagup \text{N} \\ \diagdown \text{NH} \end{smallmatrix} \text{C}_6\text{H}_4$ (Nietzki a Kehrmann, *B* 20, 3150) — $\text{K}_2\text{C}_6\text{H}_4\text{O}_2$ stable in the air — $\text{K}_2\text{C}_6\text{O}_2$ Readily oxidised by air to $\text{C}_6(\text{OK})_2\text{O}_2$ — $\text{NaC}_6\text{H}_4\text{O}_2$ dark needles with metallic lustre Sl sol water, forming a dark-yellow solution — $\text{BaC}_6\text{H}_4\text{O}_2$ (dried at 100°): dark red pp

Di-acetyl derivative $\text{C}_6(\text{OH})_2(\text{OAc})_2$ [5 3 6 4 1] [205°]. Yellow plates, sl sol water

Tetra benzoyl derivative $\text{C}_6(\text{OBz})_4\text{O}_2$ Yellow needles, sol hot BaCl (Maquenne, *Bl* [2] 48, 64, *C R* 104, 1719)

Anilide $\text{C}_6(\text{OH})_4\text{O}(\text{NPh})$ Made from $\text{C}_6(\text{OH})_4$, aniline, and alcohol Red plates with golden lustre, v sl sol ordinary solvents

References —DI-BROMO and DI-CHLORO, *de-oxy quinone*

OXY-QUINONE OXIM v NITROSO RESORCIN
DI-OXY-QUINONE DICARBOXYLIC ETHER

$\text{C}_6\text{O}_2(\text{OH})_2(\text{CO}_2\text{Et})_2$ [151°] Formed by passing dry nitrous acid gas into an ethereal solution of di-oxy terephthalic ether (Hantzsch a Loewy, *B* 19, 26, 2393, 20, 1306, 1311) Got also by dissolving $\text{C}_6\text{Cl}_2\text{O}_2(\text{CO}_2\text{Et})_2$ in NaOHAq, and by atmospheric oxidation of tetra-oxy-terephthalic ether in presence of NaOHAq (Böninger, *B* 22, 1284) Greenish-yellow monoclinic prisms (from alcohol) or yellow triclinic plates (from xylene), sl sol cold water Acid to litmus Its solutions are yellow Has no action on phenyl cyanate (Goldschmidt, *B* 23, 265) SO_2 yields $\text{C}_6(\text{OH})_2(\text{CO}_2\text{Et})_2$ Hydroxylamine produces $\text{C}_6\text{O}_2(\text{ONH}_2\text{OH})_2(\text{CO}_2\text{Et})_2$ [170°] Phenyl-hydrazine forms $\text{C}_6\text{O}_2(\text{ONH}_2\text{HPh})_2(\text{CO}_2\text{Et})_2$ [184°] Hot NaOHAq yields an amorphous sodium salt

$\text{Na}_2\text{C}_6\text{H}_4\text{O}_2$, 2aq, decomposed by hot HCl aq yielding $\text{C}_6\text{H}_4(\text{OH})_2\text{O} - \text{Na}_2\text{C}_6\text{H}_4\text{O}_2$, 2aq heavy yellow powder — $\text{Na}_2\text{C}_6\text{H}_4\text{O}_2 \cdot \text{EtOH} - \text{MgC}_6\text{H}_4\text{O}_2$, 3aq orange pp — $\text{MnCl}_2 \cdot \text{H}_2\text{O}$, 1aq — $\text{Ag}_2\text{C}_6\text{H}_4\text{O}_2$, 1aq
Acetyl derivative $\text{C}_6\text{H}_4(\text{OAc})_2(\text{CO}_2\text{Et})_2$, [174°] Needles (from HOAc) (Böninger, *B* 22, 1284)

OXY-QUINOXALINE Dihydrate

$\text{C}_6\text{H}_4 \begin{smallmatrix} \text{NH} \text{CH}_3 \\ \text{N}=\text{COH} \end{smallmatrix}$ [o 130°] Made by reduction of *o* nitro phenyl amido acetic acid with tin and HCl (Plöchl, *B* 19, 8) Prisms (containing aq), sol alcohol, ether, acids, and alkalis Melts at 94° when hydrated

Di-oxy-quinoxaline $\text{C}_8\text{H}_6\text{N}_2\text{O}_2$ *1e*

$\text{C}_8\text{H}_6 \begin{smallmatrix} \text{N} \text{COH} \\ \text{N} \text{COH} \end{smallmatrix}$ Formed by heating with HCl at 150° the compound of phenylene *o* diamine and cyanogen (Bladin, *B* 18, 674, *Bl* [2] 42, 104) Needles (containing aq), sl sol water Not melted at 290°

DI-OXY DIQUINOYL $\text{C}_8\text{H}_6\text{O}_4$ *1e* $\text{C}_6(\text{OH})_4\text{O}_2$

Rhodronic acid 'Carboxylic acid' Formed by treatment of $\text{C}_6(\text{OK})_2$ with dilute alcohol and air (Heller, *A* 24, 1, 34, 232, Lerch, *A* 124, 32, Will, *A* 118, 189) Formed also by atmospheric oxidation of $\text{C}_6\text{O}_2(\text{OK})_2$ and by reduction of C_6O_4 with SO_2 (Nietzki, *a* Benckiser, *B* 18, 513, 1838, 20, 323, 23, 3136) The hydroxyls are probably in the *o*- position Colourless crystals, forming a colourless aqueous solution

Reactions — 1 Oxidised by HNO_3 to triquinoyl C_9O_2 — 2 Air and Na_2CO_3 aq yield oronic acid — 3 *Phenylene o* diamine forms reddish-brown needles of the azine $\text{C}_8\text{H}_4\text{N}_2\text{C}_6(\text{OH})_4\text{O}_2$ (Nietzki, *a* Schmidt, *B* 21, 1227) — 4 *Tolylene-o*-diamine forms $\text{C}_8(\text{OH})_4\text{O}_2 \cdot \text{N}_2\text{C}_6\text{H}_4$, which crystallises from HOAc in yellowish brown needles
Salts — $\text{Na}_2\text{C}_8\text{O}_4$ violet needles, forming an orange aqueous solution — $\text{K}_2\text{C}_8\text{O}_4$ red powder, or small blue black needles

DI-OXY RICINOLEIC ACID $\text{C}_{18}\text{H}_{34}\text{O}_4$ *Trioxyleic acid* [64°] Made from ricinoleic acid and H_2SO_4 (Liechti, *a* Suida, *B* 16, 2455) Insol water, v e sol alcohol

OXY-SALICYLIC ACID *v* **DI OXY BENZOIC ACID**

OXY-SALTS This name is sometimes used to distinguish salts which contain O from those which do not, *1e* it is applied to salts which are not haloid salts (including cyanides), thio (or sulpho-) salts, nor salts of acids composed of H, halogen, and metal or non metal (*v* **SALTS** in vol 14) M M P M

OXY SEBACIC ACID $\text{C}_{16}\text{H}_{32}\text{O}_4$ [143°] Made by boiling di bromo sebaccic acid with water (Claus, *a* Steinkauler, *B* 20, 2886) Granules, m sol cold water — $\text{Na}_2\text{A}'$ crystalline powder, v e sol water

Di-oxy-sebacic acid $\text{C}_{16}\text{H}_{32}\text{O}_4$ [130°] Made from di bromo sebaccic acid, water, and Ag_2O (C a S) Nodules, v e sol water — $\text{Na}_2\text{A}'$ v e sol water

DI-OXY-SHIKIMIC ACID Dihydrate

$\text{CH}(\text{OH}) \begin{smallmatrix} \text{CH}(\text{OH}) \text{CH}(\text{OH}) \\ \text{CH}_2 - \text{CH}(\text{OH}) \end{smallmatrix} \text{C}(\text{OH}) \text{CO}_2\text{H}$ [156°] $[\alpha]_D = -28^\circ$. Formed from bromo shikimolactone and baryta (Eykman, *B* 24, 1294) Long needles, m sol cold water

OXY-SORBIC ACID $\text{C}_{10}\text{H}_{16}\text{O}_4$ [85°] Made from pyridine (*a*) carboxylic acid by treatment

with sodium amalgam (Weidel, *B* 12, 2001). Very deliquescent needles Reduces Fehling's solution — BaA' , — CdA' ,

***o*-OXY-STEARIC ACID** $\text{C}_{18}\text{H}_{34}\text{O}_4$ *1e* $\text{C}_{18}\text{H}_{32}\text{O}_4$ $\text{CH}(\text{OH}) \text{CO}_2\text{H}$ [81°] (G a S) [85°] (S) S (alcohol) 9.63 at 20°, 2.3 at 20° (S)

Formation — 1 A mixture of H_2SO_4 (1 mol) with oleic acid (1 mol) at 0° yields $\text{C}_{18}\text{H}_{32}\text{O}_4$ $\text{CH}(\text{SO}_3\text{H}) \text{CO}_2\text{H}$, a liquid acid soluble in ether, water, and alcohol, and forming the salts $\text{K}_2\text{A}'$, $\text{Na}_2\text{A}'$, $(\text{NH}_4)_2\text{A}'$, $\text{Ba}(\text{HA}')_2$, and $\text{Cu}(\text{HA}')_2$. This acid is decomposed by boiling dilute acids into H_2SO_4 and *o* oxy stearic acid (Geitel, *J pr* [2] 37, 74, cf Frey, *A* 33, 15, Ssabanejeff, *B* 19, 239 Ref, Saytzeff, *J p* [4] 35, 369) — 2 By the action of moist Ag_2O on iodo stearic acid prepared from oleic acid, P, and I (Saytzeff, *J pr* [2] 33, 310, 35, 378, *Bl* [2] 47, 169)

Preparation — H_2SO_4 (32g) is gradually added to olive oil (88 pts) in the cold, and the mixture boiled with alcoholic potash. The product is acidified by hydrogen chloride, and the separated acids crystallised from ether (Geitel). In this preparation there is also formed the acid $(\text{C}_{18}\text{H}_{32}\text{O}_4) \text{CH}(\text{CO}_2\text{H})_2\text{SO}_3$ [24°], which yields *o* oxy stearic acid on boiling with dilute acids

Properties — White six sided plates, m sol alcohol, v sol ether. At 200° it forms a syrupy anhydride, which is also formed by heating with fuming HCl aq at 100° for 12 hours. Does not unite with Br. Reduced by HI to stearic acid

Salts — NaA' — CaA' , small crystals (G) — CaA'_2 aq (C a S) — BaA'_2 , — ZnA'_2 , — CuA'_2 , green powder — PbA'_2 , — AgA'_2

o Oxy-stearic acid

$\text{C}_{18}\text{H}_{32}\text{O}_4$ $\text{CH}(\text{OH}) \text{CH}_2 \text{CO}_2\text{H}$ The lactone $\text{C}_{18}\text{H}_{34}\text{O}_4$ [48°] is one of the products of the action of H_2SO_4 on oleic acid. After boiling with potash it yields $\text{Ca}(\text{C}_{18}\text{H}_{31}\text{O}_3)_2$ and $\text{Pb}(\text{C}_{18}\text{H}_{31}\text{O}_3)_2$, whence acids at once set free the lactone

Di oxy-stearic acid $\text{C}_{18}\text{H}_{34}\text{O}_4$ [136°] S (alcohol) 6 at 19° S (ether) 19 at 18° (Spiridonoff, *J pr* [2] 40, 243) Formed from dibromo stearic acid (oleic acid dibromide) by treatment with moist Ag_2O (Overbeck, *A* 140, 72) Got also by boiling oxyoleic acid with potash (O) Prepared by oxidising oleic acid with alkaline KMnO_4 (Saytzeff, *J pr* [2] 31, 541, 33, 304, *Bl* [2] 45, 255) Probably identical with *o* oxy stearic acid [181°] got by oxidising tallow with KMnO_4 (Groger, *B* 22, 620) Tables, insol water, v sol hot alcohol, v sl sol ether. Converted by HI into iodo stearic acid. Reduced by alcohol and zinc to stearic acid. Distilled under 100 mm it gives an acid [c 79°], which may be its anhydride $\text{C}_{18}\text{H}_{32}\text{O}_4$. This acid forms the salts $\text{C}_{18}\text{H}_{30}\text{AgO}_4$ and $\text{C}_{18}\text{H}_{30}\text{NaO}_4$. Yields octoic, sebaccic, and azelaic acids on oxidation by KMnO_4

Salts — NaA' — KA' — CaA'_2 aq — BaA'_2 , ZnA'_2 , — AgA'_2

Di-acetyl derivative Syrup, sol ether *Methyl ether* [106°] S (alcohol) 3.45 at 18° S (ether) 1.04 at 19° Plates

Ethyl ether [100°] S (alcohol) 4.8 at 16°, 4.95 at 18° S (ether) 1.78 at 18° Plates

Di-oxy-stearic acid $\text{C}_{18}\text{H}_{34}\text{O}_4$ [100°] Formed by oxidising elaidic acid with alkaline KMnO_4 (Saytzeff, *J pr* [2] 33, 316) More sol alcohol

and ether than the preceding isomeride —NaA' —AgA'

Di oxy stearic acid $C_{18}H_{34}O_4$ [78°] Got by the action of A_2O on the dibromide of iso-oleic acid [45°], which is formed from iodo stearic acid and alcoholic potash (Saytzeff, *J pr* [2] 37, 276) Crystalline powder, v sol alcohol and ether HI yields an iodo stearic acid reduced by tin and HCl to stearic acid

Tri-oxy stearic acid $C_{18}H_{32}O_5$ 'Ricinoic acid' [141°] Occurs among the products of the oxidation of castor oil with alkaline MnO_2 (Hazura a Grussner, *M* 9, 476) —KA' —NaA' 3ag

Tri oxy stearic acid $C_{18}H_{32}O_5$ 'Ricinoic acid' [111°] Occurs together with the preceding isomeride, among the products of oxidation of castor oil by $KMnO_4$ (H a G)

Tri oxy stearic acid $C_{18}H_{32}O_5$ [115°] Formed by oxidising ricinoleic acid with alkaline $KMnO_4$ (Hazura a Grussner, *M* 10, 199) Tri metric prisms, insol cold water, v sol HOAc and ether

Tetra oxy stearic acid $C_{18}H_{30}O_6$, i.e. $C_{18}H_{30}(OH)_4O_6$ 'Sativic acid' [173°] A product of the oxidation of linoleic acid by alkaline $KMnO_4$ (Hazura, *M* 9, 190) Long prisms Converted by HI into $C_{18}H_{34}I_2O_6$ and finally into stearic acid

Hexa-oxy stearic acid $C_{18}H_{26}O_8$ 'Linusic acid' [203°] Got by oxidising linoleic acid with alkaline $KMnO_4$, being derived from linoleic acid, which is present in linoleic acid (Hazura, *M* 7, 637, 8, 155, 267) Minute needles (from water) More sol water and less sol alcohol than sativic acid Yields a hexa acetyl derivative

Hexa oxy stearic acid $C_{18}H_{26}O_8$ 'Isolinusic acid' [175°] Occurs in small quantity, together with sativic and linusic acids, in the product of the oxidation of linoleic acid by alkaline $KMnO_4$ Needles, insol ether, v sol hot water Forms a hexa acetyl derivative, sl sol ether

OXY STILBENE v OXY DI PHENYL ETHYL ENI

DI-o OXY-DI STYRYL DIKETONE $C_{18}H_{16}O_4$, i.e. $CO(CH=CH-C_6H_4-OH)_2$ [160°] Got by heating the glucoside with dilute H_2SO_4 Brownish powder, sol alcohol

Glucoside $CO(CH=CH-C_6H_4-OC_6H_5)_2$ [257°] Formed, together with the compound $CH_3COCH=CH-C_6H_4-OC_6H_5$, by condensation of helicin with acetone in presence of a little alkali (Tiemann a Kees, *B* 18, 1907) Colourless crystals (containing aq), sl sol alcohol, nearly insol water, insol ether

Tetra-oxy di styryl ketone *Di methylene derivative* $CO(CH=CH-C_6H_4-O-CH_2)_2$ [185°] Made from piperonal, acetone, and NaOHAq (Haber, *B* 24, 617) Yellow needles, insol water, m sol alcohol Colours H_2SO_4 a deep blue

o OXY STYRYL METHYL KETONE $C_{18}H_{16}O_4$, i.e. $C_6H_4(OH)CH=CHCOCH_3$ [139°] Got by hydrolysis of its glucoside by emulsion (Tiemann a Klees, *B* 18, 1904) Formed also from salicylic aldehyde, acetone, and dilute NaOH (Harries, *B* 24, 8180) Long needles (from alcohol), sl sol water $FeCl_3$ colours its aqueous solution blue Sodium amalgam reduces it to $C_6H_4(OH)CH=CHCH(OH)CH_3$ [48°].

Yields a benzoyl derivative $C_6H_4(OBz)COCH_3$ [88°], an oxim $C_6H_4(OH)C(ONH)CH_3$ [85°], and a phenyl hydrazide [160°]

Glucoside $CH_3COCH=CH-C_6H_4-O(C_6H_5)_2$ [192°] Made by adding a few drops of caustic soda solution to a mixture of acetone and helicin $C_6H_4(OC_6H_5)_2CHO$ Slender needles (containing aq), v sol hot water Lævorotatory Yields an oxim [173°]

p-Oxy-styryl methyl ketone *Methyl derivative* $C_6H_4(OMe)CH=CHCOCH_3$ [73°] Made by allowing anisic aldehyde, acetone, and aqueous NaOH to stand in the cold (Einhorn a Grabfield, *A* 243, 863) Plates, v sol alcohol

Di-oxy styryl methyl ketone *Methyl derivative* $CH_3COCH=CH-C_6H_4(OH)(OMe)$ [143°] Got from its glucoside by the action of emulsion Yellow needles, v sol alcohol

Glucoside $CH_3COCH=CH-C_6H_4(OMe)(OC_6H_5)_2$ [207°] Got by heating the glucoside of vanillin with acetone and NaOHAq (Tiemann, *B* 18, 3491) Pale yellow needles (containing 2aq), m. sol water Lævorotatory

Methylene derivative $CH_3COCH=CH-C_6H_4-O-CH_2$ [107°] Made from piperonal, acetone, and NaOHAq (Haber, *B* 24, 618) Prisms, sl sol warm water Changed by steam into an isomeride (?) [111°] Yields a yellow oxim [186°] and phenyl hydrazide [160°] The isomeride [111°] yields a colourless oxim [183°] and phenyl hydrazide [163°]

DI-OXY-STYRYL m PYRAZOLE $C_{11}H_8N_2O_4$, i.e. $CO<NHCO<NHCH=CHCHPh$ *Di oxy styryl-glyoxaline Styryl hydantoin* [172°] Made by the action of boiling dilute hydrogen chloride upon $CHPhCH=CHC_6H_5NHCONH_2$, which is got from cinnamic aldehyde cyanhydrin, and urea (Pinner a Lifschutz, *B* 20, 2353, 22, 685) White plates, changing on fusion to an isomeride [195°]

Reactions —1 On heating with alcoholic potash it yields the isomeride $CO<NHCO<NHCH=CHCHPh$ which decomposes at 300° and, with KOH and EtBr, yields $C(NH)<NEtCO<NHCH=CHCHPh$ [280°]

2 Alcoholic potash and EtBr at 100° from $CO<NEtCO<NHCH=CHCHPh$ [162°] —3 **Bromine** in $CHCl_3$ forms $C_{11}H_8N_2O_4Br_2$ [200°] **Bromine** water produces $CO<NHCO<NHCH=CHCH(OH)CHBrPh$ [220°], whence NaOHAq forms tri oxy styryl-metapyrazole $CO<NHCO<NHCH=CHCOCH_2Ph$ a crystalline powder [185°]

Acetyl derivative $CO<NACCO<NHCH=CHCHPh$ [185°] Small white prisms **o-OXY-STYRYL-PYRIDINE** $C_{13}H_{11}NO$, i.e. $C_6H_4(OH)CH=CHC<NHCH=CH$ [132°]

Formed by heating (a) methyl pyridine (10 g) with salicylic aldehyde (13 g) and water (7 g) at 140° (Butter, *B* 23, 2697) Small crystals (from dilute alcohol) Yields a dibromide Reduced by Na and alcohol to $C_6H_4(OH)C_2H_4C_6H_5N$ [94°] Salts — B_2H_3PtCl [188°] — $B'HgSO_4$ [e 170°] Salts of the ethyl derivative

$(C_6H_5)_2(OEt)_2N_2 \cdot H_2HgCl_4$ [92°] Light-yellow needles — $(C_6H_5)_2(OEt)_2N_2 \cdot H_2PtCl_4$ [183°]

Ethyl-o-tolide of the ethyl derivative
 $C_6H_5(OEt)NEtI$ [218°] Golden crystals

Oxy-styryl-pyridine Methyl derivative
 $C_6H_5(OMe)CH(OH)C_6H_4N$ [97°] Formed from anisic aldehyde and methyl pyridine (Schuffen, B 23, 2719) Plates — $B_2H_2PtCl_4$ [184°]. Yellow plates

p-OXY (Py 3)-STYRYL QUINOLINE

$C_{11}H_{11}NO$ *s.s.* $C_6H_4 \cdot N \equiv C \cdot OH \cdot CH \cdot C_6H_4 \cdot OH$ [258°] Formed from *p*-amido-styryl-quinoline by the diazo-reaction (Eulach, B 22, 286) Formed also from *p*-oxybenzoic aldehyde, quinaldine, and $ZnCl_2$ (Wallach, B 16, 2009) Yellow plates

o-Oxy-(Py 1)-styryl-quinoline [215°] Made from salicylic aldehyde, lepidine, and $KHSO_4$ (Heymann & Königs, B 21, 1429, 2172) Yellow crystals

m-Oxy-(Py 1)-styryl-quinoline. [255°] Got from *m*-amido-styryl quinoline Brownish-yellow crystals (from alcohol)

p-Oxy-(Py 1)-styryl quinoline [249°] Got from *p*-oxy benzoic aldehyde, lepidine, and $KHSO_4$ (H. & K) Yellow crystals (from alcohol)

OXYSUBERANIC ACID $C_8H_8O_4$ *s.s.* $C_6H_5(OH)CO_2H$ [90°] Made from suberone cyanhydrin and conc $HClAq$ (Spiegel, A 211, 118) Tables (containing 2aq) Melts at 50° when hydrated V sol alcohol and hot water

OXYSUBERIC ACID $C_8H_8O_4$ [112°] Made by boiling bromo-suberic acid with $NaOHAq$ (Hell & Rempel, B 15, 149, 18, 817) Nodules (from ether) HNO_3 oxidises it to adipic and oxalic acids — MgA'' aq — CuA'' — ZnA'' 2½aq — Ag_2A'' crystalline pp

Ethyl derivative $C_8H_{11}(OEt)(CO_2H)_2$ Made from bromo suberic acid and alcoholic potash. Syrup, v sol water and alcohol — BaA'' — ZnA'' — PbA'' — Ag_2A'' amorphous pp

Oxy-isosuberic acid $C_8H_8O_4$ A sticky mass got by saponification of its ether, which is a product of the action of silver on bromo butyric ether (Hell & Muhlhauser, B 18, 477) $HIAq$ at 160° reduces it to β isosuberic acid — Ag_2A''

Di-oxy-suberic acid $C_8H_8O_6$ Made from di-bromo-suberic acid and KOH (Gay & Gay-Lussac, A. 155, 251) Amorphous

Di-ethyl derivative $C_8H_{10}(OEt)_2(CO_2H)_2$ Made from di bromo suberic acid and alcoholic potash (H. & R.) Syrup, v sol water, alcohol, and ether — Ag_2A'' amorphous pp

OXY-SUCCINIC ACID *v* **MALIC ACID**

Di-oxy-succinic acid v **TARTARIC ACID**

Tetra-oxy-succinic acid

$C_4H_4O_8$ *s.s.* $CO_2H \cdot C(OH) \cdot C(OH) \cdot CO_2H$ or $CO_2H \cdot CO \cdot CO \cdot CO_2H$ 2aq *Di-oxy tartaric acid*
Carboxytartaric acid [98°] Formed by passing nitrous acid into an ethereal solution of protocatechuic acid (Grüber, B 12, 514), of pyrocatechin (Barth, M. 1, 869), of guaiacol (Hering, M. 8, 825), or of 'nitro'-tartaric acid (Kekulé, A 221, 240). The free acid is got by decomposing the dry sodium salt in dry ether with HCl gas (W. L. Miller, B 22, 2015). White crystals, v sol. water The salt $Na_2C_4H_4O_8 \cdot 2aq$ is nearly insol water It splits up on heating into CO_2 and sodium tartronate. With $NaHSO_4$,

at 90° it yields glyoxal (Hinsberg, B 24, 3235). The salt $Ba_2(C_4H_4O_8)_2 \cdot aq$ is a crystalline pp. $HClAq$ and zinc reduce it to a mixture of racemic and inactive tartaric acids Yields hydantoin when the Na salt is rubbed up with urea and dilute $HClAq$ at 55° (Anschütz, A 254, 258)

Ethyl ether Thick liquid (Anschütz, A 261, 130). Converted by urea into the ureide $C_4H_4N_4O_6$, which crystallises in needles, sl sol cold water, decomposing at 245°

(*B*)-*Oxim* $CO_2H \cdot C(NOH) \cdot C(NOH) \cdot CO_2H$ [145°–150°] Got from the Na salt, hydroxyamine and HCl Colourless prisms, v sol water and alcohol, insol benzene (Müller, B 16, 2985, Söderbaum, B 24, 1223) After crystallisation from water it melts at 70°–75° It yields the salts CaA'' aq and Ag_2A'' and a crystalline diacetyl derivative, v sol water Conc $HClAq$ converts the oxim into an isomeride [145°–150°] crystallising in nodules, which are v sol water and yield the salts CaA'' 8aq and Ag_2A'' aq Ag_2O decomposes this (a) oxim into cyanogen and CO_2

Phenyl hydrazide $C_{10}H_8N_2O_4$ *s.s.* $CO_2H \cdot CO \cdot C(N_2HPh) \cdot CO_2H$ *Monophenylisone dioxytartaric acid* [218°] Made by mixing a solution of the acid (1 mol) with phenyl hydrazine (1 mol) dissolved in $HClAq$ (Ziegler & Locher, B 20, 835) Feathery needles, insol cold water, v sol hot alcohol Coloured red by $FeCl_3$ — $BaC_{10}H_8N_2O_8 \cdot 3aq$ — Na_2A'' — Ag_2A'' orange yellow pp

Di-phenyl di-hydrazide
 $CO_2H \cdot C(N_2HPh) \cdot C(N_2HPh) \cdot CO_2H$ *Diphenylisone dioxytartaric acid* [above 200°] Made from tetra-oxy succinic acid (1 mol) and phenyl hydrazine (2 mols) Orange-yellow powder, v al sol water, v sol warm alcohol On heating with Ag_2O it yields $O < CO \cdot C \cdot N_2HPh$
 $CO \cdot C \cdot N_2HPh$ [c 234°], which

crystallises in red needles, v sol Ag_2O — $(NH_4)_2A''$ colourless plates — $(NH_4)HA''$ brick red needles With silver nitrate it gives

$AgN < CO \cdot C \cdot N_2HPh$ of a vermilion colour

Na_2A'' lemon-yellow plates — $NaHA''$ brick red needles — The ethyl ether $C_{10}H_{12}N_2O_4$ [121°] is got from phenyl hydrazine and tetra oxy succinic ether (A & G), it forms canary yellow crystals

Di-p-sulpho di-phenyl-di-hydrazide
 $CO_2H \cdot O(NH_2CH_2SO_3H) \cdot C(N_2HPh) \cdot CO_2H$
 Formed from sodium tetra-oxy succinate, sodium phenyl-hydrazine *p*-sulphonate, and $HClAq$ The Na salt is an orange-yellow powder, v sol water, insol alcohol, and is used as a yellow dye (tartrazine)

Di-phenyl-hydrazide [158°] Made from the acid and N_2H_5Ph Yellow pp, turning blue in the air

Tetra phenyl-di hydrazide
 $CO_2H \cdot C(N_2Ph) \cdot C(N_2Ph) \cdot CO_2H$ [177°] Made from the acid (1 mol.) and diphenylhydrazine (2 mols) (Ziegler & Locher, B 20, 841). Yellowish plates (from hot alcohol), insol. water. Conc H_2SO_4 forms a red solution, becoming green On fusion with resorcin it gives a crimson mass, coloured blue by $NaOH$, hydroquinone and pyrogallol give green and blue masses respectively. Bromine gives a crystalline bromo-

derivative Ac_2O forms $\text{O} \begin{smallmatrix} \text{CO} \text{C} \text{N} \text{Ph}_2 \\ \text{CO} \text{C} \text{N} \text{Ph}_2 \end{smallmatrix}$ [222°] crystallising in ruby red prisms with green lustre
 Converted into the imide $\text{NH} \begin{smallmatrix} \text{CO} \text{C} \text{N} \text{Ph}_2 \\ \text{CO} \text{C} \text{N} \text{Ph}_2 \end{smallmatrix}$ [192°] by NH_3 — $\text{Na}_2\text{A}''$ rosettes of needles (from water) or plates (from alcohol)— CuA'' — PbA'' . white pp

m-Nitro-phenyl-hydrazide [175°] Made by using *m* nitro phenyl hydrazine (Bischler a Brodsky, *B* 22, 2814) Sl sol hot water and alcohol

Di-m nitro di-phenyl-dihydrazide [α 22°] Made from *m* nitro-phenyl-hydrazine (2 mols) and tetra oxy-succinic acid (1 mol). Sl sol hot water

OXY-SULPHIDES Compounds of an element, or positive compound radicle, with O and S The term is generally restricted to compounds of metals with O and S Oxy-sulphides of metals are not numerous, and they have not been much studied They are produced in various reactions, of which the following are the chief by acting on the sulphide with water, *eg* oxy-sulphide of Ba, by heating the oxide with S, *eg* Bi oxy-sulphide, by boiling the oxide and sulphide with water, *eg* Ca oxy-sulphide, by partially reducing the sulphate by H, *eg* oxy-sulphides of Co and Mn, by heating the oxide in H_2S , or by passing H_2S into an aqueous solution of the oxide, *eg* oxy-sulphides of Fe and Os, by adding a little alkali sulphide to the solution of a salt, *eg* Cu oxy-sulphide **M M P M**

OXY-SULPHO-BENZOIC ACID $\text{C}_6\text{H}_4\text{SO}_3 \cdot e$ $\text{C}_6\text{H}_3(\text{OH})(\text{SO}_3\text{H})\text{CO}_2\text{H}$ *Sulphosalicylic acid* [120°] Made by sulphonation of salicylic acid (Mendius, *A* 103, 45, Remsen, *A* 179, 107) Long thin needles, v e sol water and alcohol Gives a reddish-violet colour with FeCl_3 Yields phenol and salicylic acid on fusion with potash— NaHA'' 2aq— $\text{Na}_2\text{A}''$ 3aq— KHA'' 2aq— $\text{K}_2\text{A}''$ 2aq— $\text{K}_2\text{HA}''$ 1aq— KNaA'' 4aq— CaA'' 1aq— $\text{Ba(EA}'')$ 4aq— MgA'' 3aq— ZnA'' 8aq— PbA'' — CuA'' — $\text{Cu}_2\text{A}''(\text{OH})$ 1aq— $\text{Ag}_2\text{A}''$ 1aq crystalline powder, v sol hot water

Ethyl ether EtA'' [58°]. Crystals

Oxy-sulpho benzoic acid

$\text{C}_6\text{H}_3(\text{OH})(\text{SO}_3\text{H})(\text{CO}_2\text{H})$ [4 3 1] Made from *p*-oxy benzoic acid by treatment with SO_2 (Kölle, *A* 164, 150) or by digesting with H_2SO_4 at 100° (Klepl, *J pr* [2] 28, 196) Deliquescent needles, v e sol water and alcohol, insol ether FeCl_3 gives a red colour Potash-fusion yields protocatechuic acid— KHA'' 1aq crystals, sl sol water (Klepl)— $\text{K}_2\text{A}''$ 1aq (Kölle)— $\text{K}_2\text{C}_6\text{H}_3\text{SO}_3$ 2aq (Kölle)— BaA'' 4aq— $\text{Ba}_2(\text{C}_6\text{H}_3\text{SO}_3)_2$ amorphous (Kölle)— CdA'' 8aq— $\text{Ag}_2\text{A}''$ small prisms

Methyl derivative

$\text{C}_6\text{H}_3(\text{OMe})(\text{SO}_3\text{H})(\text{CO}_2\text{H})$ Made from fuming H_2SO_4 and anisic acid at 160° (Zervas, *A* 108, 388, Lmprecht, *Gm* 13, 128) Needles, insol ether— BaA'' 1aq— PbA'' 1aq: needles, sl sol Aq

Oxy-sulpho-benzoic acid

$\text{C}_6\text{H}_3(\text{OH})(\text{SO}_3\text{H})(\text{CO}_2\text{H})$ [4 2 1] Formed by the diazo-reaction from (4,2,1)-amido sulpho-benzoic acid (Hedrick, *Am* 9, 416) Crystalline, sol water, alcohol, and ether— $\text{Ba(HA}'')$, insol dilute HOAc — BaA'' — CaA'' 5aq triclinic prisms, sol hot water.— CoA'' 7aq— CdA'' 8aq— ZnA'' 8aq (Pisanello).

Imide of the ethyl derivative

$\text{C}_6\text{H}_3(\text{OEt}) \begin{smallmatrix} \text{CO} \\ \text{SO} \end{smallmatrix} \text{NH}$ [268°] Made by oxidising $\text{C}_6\text{H}_4\text{Me}(\text{OEt})\text{SO}_2\text{NH}_2$ with KMnO_4 and ppg with HCl (Remsen a Palmer, *Am* 8, 227) Concentric groups of needles— $\text{KC}_6\text{H}_4\text{NSO}_4$ — $\text{AgC}_6\text{H}_4\text{NSO}_4$ lustrous needles, sl sol water.

Oxy-sulpho-benzoic acid

$\text{C}_6\text{H}_3(\text{OH})(\text{SO}_3\text{H})(\text{CO}_2\text{H})$ [3 4 ? 1] Formed from *m* oxy-benzoic acid and SO_3 (Barth, *A* 148, 38, Senhofer, *A* 152, 102) Yellowish green needles (containing 1/2 aq), v sol alcohol Coloured red by FeCl_3 Yields protocatechuic acid and an acid [189°] when fused with potash— BaA'' 4/3 aq— $\text{Cd(HA}'')$ 2aq— $\text{Pb}_2(\text{C}_6\text{H}_3\text{SO}_3)_2$ reddish amorphous mass

Oxy-sulpho benzoic acid

$\text{C}_6\text{H}_3(\text{OH})(\text{SO}_3\text{H})\text{CO}_2\text{H}$ Got by dissolving *m* diazo benzoic acid in warm H_2SO_4 (Griess, *Z* 1864, 538) White laminae (from water) Perhaps identical with the preceding acid— BaA'' .

Oxy-disulpho-benzoic acid

$\text{C}_6\text{H}_3(\text{OH})(\text{SO}_3\text{H})_2\text{CO}_2\text{H}$ [146°] Made from salicylic acid and ClSO_3H at 180° (Pisanello, *G* 18, 346) Deliquescent needles (containing 4/3 aq) Gives a red colour with FeCl_3 — $\text{Na}_2\text{A}''$ 3aq— $\text{K}_2\text{A}''$ 3aq— $\text{Ca}_2\text{A}''$ 12aq— $\text{Ba}_2\text{A}''$ 6 1/2 aq prisms, sl sol water— $\text{Pb}_2\text{A}''$ 10aq— $\text{Cd}_2\text{A}''$ 18aq— $\text{Cu}_2\text{A}''$ 12aq— $\text{Zn}_2\text{A}''$ 15aq plates

Oxy-disulpho-benzoic acid $\text{C}_6\text{H}_3\text{S}_2\text{O}_6$ Got by boiling trisulpho *m* oxy benzoic acid with BaCO_3 (Kretschy, *B* 11, 862)— BaA'' 8aq

Oxy-trisulpho-benzoic acid

$\text{C}_6\text{H}_3(\text{OH})(\text{SO}_3\text{H})_3\text{CO}_2\text{H}$ Made by heating *m*-oxy-benzoic acid with H_2SO_4 , SO_3 , and P_2O_5 at 250° (Kretschy, *B* 11, 858) Hygroscopic syrup (containing 4/3 aq at 100°) FeCl_3 gives a red colour— $\text{K}_2\text{C}_6\text{H}_3\text{S}_3\text{O}_6$ 2aq monoclinic prisms— $\text{K}_2\text{A}''$ 2aq $\text{Pb}_2(\text{C}_6\text{H}_3\text{S}_3\text{O}_6)_2$ 6aq— $\text{Pb}_2\text{A}''$ 8aq— $\text{Cd}_2\text{A}''$ 3aq— $\text{Ba}_2\text{A}''$ 4aq colourless plates

Di-oxy-sulpho-benzoic acid

$\text{C}_6\text{H}_3(\text{OH})_2(\text{SO}_3\text{H})\text{CO}_2\text{H}$ Formed from (4, 2, 1)-di oxy benzoic acid and H_2SO_4 (Zehenter, *M* 2, 468) Hygroscopic needles (containing 2aq) FeCl_3 colours its aqueous solution red—**Salts**— $\text{K}_2\text{A}''$ 8 1/2 aq— BaA'' 2aq— PbA'' 2aq— $\text{Cu}_2(\text{C}_6\text{H}_3\text{SO}_3)_2$ 5aq— $\text{Ag}_2\text{A}''$ 2aq needles

Di oxy-sulpho-benzoic acid

$\text{C}_6\text{H}_3(\text{OH})_2(\text{SO}_3\text{H})(\text{CO}_2\text{H})$ [5 2 x 1] Made by heating (5, 2, 1)-di oxy-benzoic acid with H_2SO_4 and P_2O_5 at 180° (Senhofer a Sarlay, *M* 2, 454) Needles— $\text{K}_2\text{A}''$ 1aq— $\text{Ba(HA}'')$ 8 1/2 aq— BaA'' 2aq— PbA'' 2aq crystalline powder

OXY SULPHO-ISOCUMINIC ACID

$\text{CMe}_2(\text{OH})\text{C}_6\text{H}_3(\text{SO}_3\text{H})(\text{CO}_2\text{H})$ Made by oxidising the sulphonic acids of cymene and *m*-isocymene by KMnO_4 (R Meyer, *A* 220, 7, 29)— $\text{K}_2\text{A}''$ 5aq— BaA'' 1aq—minute plates— PbA''

OXY - SULPHOCYANO - ACETOACETIC

ETHER. *Anhydride* $\text{O}(\text{C}(\text{Ac}(\text{SCy})\text{CO}_2\text{Et}))_2$ [α 163°] Made by heating equivalent weights of barium sulphocyanide and dichloro-acetoacetic ether (207°) (Zürcher, *A* 250, 293) Yellow crystalline powder (from alcohol), insol ether.

OXY-SULPHO-NAPHTHOIC ACID

$\text{C}_{10}\text{H}_6(\text{OH})(\text{SO}_3\text{H})\text{CO}_2\text{H}$ Made from (α)-oxy-naphthoic acid and H_2SO_4 at 60° (König, *B* 22, 787, 28, 806) Needles (containing 5aq), v sol water and alcohol Its alkaline solutions fluoresce blue HNO_3 yields di-nitro-naphthol [188°]. Diazobenzene chloride forms the azo-compound

$C_{10}H_7(OH)(N_2Ph)SO_3H - NaHA''$ needles, v sol water $- Na_2A''$ zaq $- Ba(HA'')_2$

Oxy-di-naphtho-naphthoic acid

$C_{10}H_7(OH)(SO_3H)_2CO_2H$ Formed from (o) oxynaphthoic acid and fuming H_2SO_4 (K) Stellate groups of needles (containing 4aq) $- K_2A'' - Ba_2(C_{10}H_7S_2O_6)_2$ needles, m sol water

OXY-SULPHYDRO-ALLYL-QUINAZOLINE

$C_8H_7 \begin{smallmatrix} \diagup CO \ N \ C_2H_5 \\ \diagdown NH \ CS \end{smallmatrix}$ *Allylthio benzoylurea* [199°]

Formed from o amido benzamide and allyl thio carbimide (Stewart, *J pr* [2] 44, 416) Needles Its benzene solution fluoresces blue

OXY SULPHYDRO DI-METHYL-GLYOXALINE *Di methyl derivative*

$NMe \begin{smallmatrix} \diagup C(OMe) \ CMe \\ \diagdown C(SMe) \ N \end{smallmatrix}$ Formed from di methylthiohydantoin [167°], alcoholic potash, and MeI (Marckwald, *B* 24, 3293) Syrup $- B'HCl$ syrup $- B'H_2SO_4$ crystalline $- B'H_2PtCl_6$

The isomeride $NMe \begin{smallmatrix} \diagup CO \ CMe_2 \\ \diagdown C(SMe) \ N \end{smallmatrix}$ formed from tri methyl thiohydantoin, alcoholic potash, and MeI yields the crystalline salts $B'H_2SO_4$ [138°] and $B'H_2PtCl_6$ [150°]

OXY-SULPHYDRO-PHENYL-METHYL-GLYOXALINE *Di-methyl derivative*

$NPh \begin{smallmatrix} \diagup C(OMe) \ CMe \\ \diagdown C(SMe) \ N \end{smallmatrix}$ [90°] Formed from phenyl methyl thiohydantoin, alcoholic KOH, and MeI (Marckwald, *B* 24, 3290) Crystals $- B'HCl$ [140°] $- B'H_2PtCl_6$ [213°] $- B'C_2H_5N_3O_6$ [192°]

The isomeric compound $NPh \begin{smallmatrix} \diagup CO \ CMe_2 \\ \diagdown C(SMe) \ N \end{smallmatrix}$ [223°] from phenyl-di methyl thiohydantoin [67°] forms the salts $B'HCl$, $B'H_2PtCl_6$ [132°] and $B'C_2H_5N_3O_6$ [174°]

OXY-SULPHYDRO-PHENYL-QUINAZOLINE

$C_8H_7 \begin{smallmatrix} \diagup CO \ NPh \\ \diagdown NH \ CS \end{smallmatrix}$ [199°] Formed from o-amido benzamide and phenyl thiocarbimide (Stewart, *J pr* [2] 44, 416) Satiny tables

OXY-SULPHYDRO-QUINAZOLINE

$C_8H_7 \begin{smallmatrix} \diagup CO \ NH \\ \diagdown NH \ CS \end{smallmatrix}$ [281°] Formed from o amido benzamide and thio urea (Stewart, *J pr* [2] 44, 416) Nodules (from alcohol)

OXY-SULPHYDRO-THIAZOLE $C_8H_7NS_2O_2$

$NH \begin{smallmatrix} \diagup OS \ S \\ \diagdown CO \ CH_3 \end{smallmatrix}$ [167°] Formed by heating

$NH \begin{smallmatrix} \diagup C(NH) \ S \\ \diagdown CO \ OH_2 \end{smallmatrix}$ with OS_2 in alcohol at 160° (Miolati, *A* 262, 84) Formed also from chloro acetic ether, ammonium dithiocarbamate, and alcoholic HCl, and from sulphocyanosacetic ether by successive treatment with H_2S and HCl

OXY-SULPHYDRO-o-TOLYL-METHYL GLY-

OXALINE *Di methyl derivative*

$C_8H_7MeN \begin{smallmatrix} \diagup C(OMe) \ CMe \\ \diagdown C(SMe) \ N \end{smallmatrix}$ [120°] Got from o-tolyl-methyl-thiohydantoin, alcoholic potash, and MeI (Marckwald, *B* 24, 3292) Plates, v sol alcohol Yields the following salts $B'HCl$ [120°], $B'H_2PtCl_6$, $B'H_2SO_4$ [205°], $B'HNO_3$, and the picrate $B'C_2H_5N_3O_6$ [200°]

The isomeric $C_8H_7MeN \begin{smallmatrix} \diagup C(SMe) \ N \\ \diagdown CO \ CMe_2 \end{smallmatrix}$ from o-tolyl di methyl thiohydantoin yields the salts $B'HCl$ [118°], $B'H_2PtCl_6$, $B'H_2SO_4$ [208°], and $B'C_2H_5N_3O_6$ [212°].

Oxy-sulphydro-p-tolyl-methyl-glyoxaline *Di-methyl derivative* [109°] Plates Yields $B'HCl$ [123°], $B'H_2PtCl_6$, and $B'C_2H_5N_3O_6$ [180°]

The isomeric $[4 \ 1] C_8H_7MeN \begin{smallmatrix} \diagup C(SMe) \ N \\ \diagdown CO \ CMe_2 \end{smallmatrix}$ yields $B'H_2SO_4$ [210°], $B'H_2PtCl_6$ [162°], and $B'C_2H_5N_3O_6$, decomposing at 190°

DI-OXY TARTARIC ACID v **TETRA OXY-SUCCINIC ACID**

OXY TEREBIC ACID $C_8H_7O_3$ [100°-120°] Made by boiling chloro terebic acid with water and $CaCO_3$ (W Roser, *A* 220, 244) Syrup, crystallising with difficulty, v sol water $- CaA'$, $- AgA'$ needles, v sol water

OXY TEREPHTHALIC ACID $C_8H_7O_3$ $is C_8H_7(OH)(CO_2H)_2$ Formed by the diazo reaction from amido terephthalic acid (De la Rue a Müller, Burkhardt, *B* 10, 144, 1273) Prepared also by potash fusion from bromo terephthalic acid (Fischli, *B* 12, 621), from oxy aldehyde benzoic acid (Tiemann a Landshoff, *B* 12, 1335), from p xyleneol, from carvacrol, and thymol (Jacobsen, *B* 11, 570), from oxy p toluic acid (Hall a Remsen, *B* 12, 1433), and from rufigallic acid (Schreder, *M* 1, 439)

Properties—Crystalline powder, sl sol water, v sol alcohol Not melted at 300° $FeCl_3$ colours its solution reddish violet Yields CO_2 and phenol when strongly heated On fusion with NaOH it gives salicylic acid and some p-oxy benzoic acid (Barth a Schreder, *B* 12, 1260) Yields m oxy benzoic acid on heating with $HClAq$ at 120° Forms a di nitro derivative [179°]

Salts $- BaA''$ 3½ aq lamina (from water) $- Ag_2A''$ white pp, insol water

Methyl ether Me_2A'' [94°] Leaflets Yields an acetyl derivative $C_8H_7(OAc)(CO_2Me)_2$ [76°]

Methyl derivative $C_8H_7(OMe)(CO_2H)_2$ [279°] Got by oxidation of methoxy toluic acid and of methyl thymol Small prisms

Methylether of the methyl derivative $C_8H_7(OMe)(CO_2Me)_2$ [65°] Concentric needles

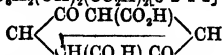
Ethyl derivative $C_8H_7(OEt)(CO_2H)_2$ [254°] Got by oxidation of the ethyl ether of thymol (Paterno a Canzoneri, *G* 9, 460) Stellate groups of minute crystals, insol water

Benzyl derivative $C_8H_7(OCH_2Ph)(CO_2H)_2$ [230°-240°] Made from $C_8H_7(ONa)(CO_2Me)_2$ and benzyl chloride, the product being saponified (Bayer a Tutein, *B* 22, 2188) Slender needles

Tetrahydride $C_8H_7(OH)(CO_2H)_2$ Got by reducing the acid with sodium amalgam (Bayer a Tutein, *B* 22, 2180) Yellowish white needles, sl sol cold water Coloured bluish violet by $FeCl_3$ Its methyl ether forms feathery needles [77°] Hydroxylamine converts the tetrahydride into $C_8H_7O_2(NOH)$ [170°], while phenyl hydrazine forms $C_8H_7O_2(N_2HPH)$ [125°] converted by $HClAq$ into $NH \begin{smallmatrix} \diagup C_8H_7 \\ \diagdown \end{smallmatrix} CO_2H$, a crystalline

body [230°] giving off a faecal odour when heated The tetrahydride, heated at 120°, loses CO_2 and forms $C_8H_7O(CO_2H)$, whence NaA' and $C_8H_7(OH)CO_2H$ [180°-140°] may be prepared, the latter body yielding the crystalline acid $C_8H_7(OH)(CO_2H)_2$ on saponification

Di-oxy-terephthalic acid $C_8H_6O_4$ \pm $C_8H_4(OH)_2(CO_2H)_2$ [5 2 4 1] or



Quinone hydro dicarboxylic acid Hydroquinone dicarboxylic acid [above 300°]

Formation—1 By passing a current of air through a solution of the dihydride of its ether (succinyl succinic ether) containing excess of alkali (Herrmann, *B* 10, 107, *A* 211, 335)—2 By the oxidising action of KMnO_4 on $C_8H_4Me_2(OPO_4K)_2$, a derivative of thymoquinone, and on $C_8H_4Me_2(OPO_4K)_2$, a derivative of hydro *p* xyloquinone (Heymann *a* Königs, *B* 20, 2392)—3 By the action of PCl_5 on succinylsuccinic ether (Levy *a* Curchod, *B* 22, 2108)—4 By saponifying its ether

Properties—Interlaced needles (containing 2aq) (from water) or yellow plates (from alcohol), sl sol alcohol and ether, v sl sol water. Its alcoholic solution shows blue fluorescence. FeCl_3 gives a deep blue colour. When distilled it yields hydroquinone. Bromine water yields tetrabromquinone. Chlorine passed into its alcoholic solution forms tetra chloro quinone (Loewy, *B* 19, 2394)

Salts— $\text{K}_2\text{A}''$ yellow needles, forming a solution with green fluorescence— KHA'' — $\text{Na}_2\text{A}''$ 2aq— NaHA'' 2aq— $\text{Na}_2\text{A}''(\text{OH})_{10}\text{aq}$ crystals ppd by conc NaOHaq — $(\text{NH}_4)_2\text{A}''$ 2aq (Duisberg, *A* 213, 1(3)— BaA'' — CaA'' 5aq— $\text{Ca}(\text{HA}'')_2$ 5aq— PbA'' — $\text{Ag}_2\text{A}''$ green yellow pp

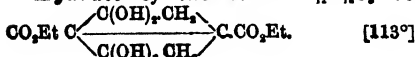
Mono ethyl ether EtHA'' [184°] Made by the action of dilute KOH on the di ethyl ether in the cold. Pale yellow needles (from water), sl sol water, alcohol, and ether— $\text{Ba}(\text{EtA}'')_2$ 5aq— $\text{Ca}(\text{EtA}'')_2$ 5aq greenish yellow needles

Di ethyl ether Et_2A'' [133°] S (ether) 16 at 20°. Prepared by adding bromine to a solution of its dihydride in CS_2 , or to the dihydride suspended in water (Herrmann, *A* 211, 327, *B* 19, 2229). Formed also by the action of sodium on di bromo acetoacetic ether dissolved in ether (Wedel, *A* 219, 74)

Properties—Greenish yellow trimetric tables (from benzene) or needles (from ether). May be sublimed. Its alcoholic solution shows blue fluorescence. FeCl_3 gives a bluish green colour. Does not react with phenyl cyanate (Goldschmidt *a* Meissler, *B* 23, 259). Does not react with phenyl hydrazine or hydroxylamine. Sol alkalis, forming a yellow liquid, from which it is ppd by CO_2 , and, as a scarlet pp, $\text{Na}_2C_2H_2O_4$, by conc NaOHaq . Ac_2O has no action at 510°

Reactions—1 Reduced to its dihydride by *zinc* and HClAq (Baeyer, *B* 19, 428)—2 *Bromine vapour* forms $C_8H_4Br_2O_4(\text{CO}_2\text{Et})_2$ [157°] and afterwards $C_8Br_2O_4(\text{CO}_2\text{Et})_2$ (Herrmann, *B* 19, 2234, Böninger, *B* 21, 1758)—3 *Dry nitrous acid gas* passed into its solution in absolute ether oxidises it to di oxy quinone dicarboxylic ether (Hantzsch *a* Loewy, *B* 19, 26), an intermediate crystalline powder $C_8H_4NO_6$ or $C_8H_4NO_6$ [148°] giving a violet colour with alkalis being first formed (Loewy, *B* 19, 2393)

Hydrate of the ether $C_8H_4O_4$ is



Formed, together with the ether, by the action of bromine on the dihydride (succinyl-succinic ether) (Hantzsch *a* Zeckendorf, *B* 20, 2800). Yellow needles. On boiling with alcohol it changes to the ether $C_8H_4O_4$ [133°]. Hydroxylamine reduces it to $C_8H_4H_2O_4$ [128°]

Di acetyl derivative of the ether $C_8H_4(\text{OAc})_2(\text{CO}_2\text{Et})_2$ [154°] Got by the action of AcCl on the ether or on $C_8H_4(\text{ONa})_2(\text{CO}_2\text{Et})_2$ (Wedel, *A* 219, 81, Nef, *A* 258, 306). Monoclinic crystals. Not attacked by Br . Its alcoholic solution is not fluorescent

Di benzoyl derivative of the ether $C_8H_4(\text{OBz})_2(\text{CO}_2\text{Et})_2$ [174°] Made from the ether, NaOHAq , and BzCl . Colourless needles (from alcohol). Not attacked by Br in CHCl_3 . Conc H_2SO_4 sets free $C_8H_4(\text{OH})_2(\text{CO}_2\text{Et})_2$. Zinc dust and conc HClAq form three isomeric dihydrides $C_8H_4(\text{OBz})_2(\text{CO}_2\text{Et})_2$, melting at 165°, 85°-95°, and 100°-110°

Di methyl derivative $C_8H_4(\text{OMe})_2(\text{CO}_2\text{H})_2$ [265°] Formed by saponification of $C_8H_4(\text{OMe})_2(\text{CO}_2\text{Et})_2$ (*v infra*) with alcoholic potash (Nef, *A* 258, 298). Colourless needles. Its aqueous solution shows bluish-violet fluorescence— $(\text{NH}_4)_2\text{A}''$ sol water. Not fluorescent

Di methyl derivative of the ethyl ether $C_8H_4(\text{OMe})_2(\text{CO}_2\text{Et})_2$ [101 5°] Made from $C_8H_4(\text{ONa})_2(\text{CO}_2\text{Et})_2$ and MeI at 100° (Nef, *A* 258, 297). Colourless tables. Its solutions show bluish violet fluorescence. Not reduced by zinc dust and HOAc

Di benzyl derivative of the ethyl ether $C_8H_4(\text{OCH}_2\text{Ph})_2(\text{CO}_2\text{Et})_2$ [96 5°] Monoclinic needles (from alcohol), insol KOHaq

Oxy-amide $C_8H_4(\text{OH})_2(\text{CO}_2\text{NH}_2)_2$ Formed by the action of hydroxylamine on the ether (Jeaurenand, *B* 22, 1278). Prisms (containing 2aq), sol water and alcohol

Dioxyterephthalic acid dihydride $C_8H_6O_4$, Succinyl succinic acid S 015 at 19 5° Got from its ether and the calculated quantity of cold NaOHAq . Minute needles, decomposed by heat, yielding $C_8H_6O_4$ [78°] and CO_2 . FeCl_3 colours its solution violet (Baeyer *a* Noyes, *B* 22, 2168)— BaA' 2 1/2 aq— $(\text{NH}_4)_2\text{A}'$ 2aq

Methyl ether $\text{Me}_2\text{A}''$ [132°] Made by the action of sodium on methyl succinate (Ebert, *A* 229, 50). Got also from bromo acetoacetic ether and ammonia (Geuther, *A* 244, 204). Sol. alcohol and ether

Mono ethyl ether EtHA'' [98°] Got from the di ethyl ether and cold NaOHAq (Herrmann). Yellowish prisms (from ether). Its alcoholic solution shows blue fluorescence. FeCl_3 gives a violet colour. Decomposed by boiling water into CO_2 and $\text{C}_8\text{H}_4\text{EtO}_4$

Di-ethyl ether $\text{Et}_2\text{A}''$ is

$$\text{CO}_2\text{Et} \text{ CH} \begin{array}{c} \diagup \text{CH} \text{CO} \\ \diagdown \text{CO} \text{CH}_2 \end{array} \text{ CH CO}_2\text{Et} \text{ or } \text{CO}_2\text{Et} \text{ CH} \begin{array}{c} \diagup \text{CH}=\text{C}(\text{OH}) \\ \diagdown \text{C}(\text{OH}) \text{CH} \end{array} \text{ CH CO}_2\text{Et} \quad [127^\circ]$$

 S G Ψ 141 S (ether) 16 at 17°

Formation—1 By the action of K or Na on ethyl succinate (Fehling, *A* 49, 186, Herrmann, *A* 211, 306, Duisberg, *B* 16, 133)—2. By the action of NaOEt (free from alcohol) on succinic ether (Volhard, *B* 16, 134)—3 From bromo-acetoacetic ether by the action of alcoholic NH_3 (Duisberg, *A* 218, 183) or of NaOEt

(Wedel, *A* 219, 92) —4. By reducing di oxy-terephthalic ether with zinc and HClAq (Baeyer, *B* 19, 428)

Properties—Green triclinic crystals with blue fluorescence (from ether), v sl sol. hot water Insol NH₄Aq, but forms a yellow solution in NaOHAq Its alcoholic solution shows blue fluorescence FeCl₃ gives a red colour Decomposed by excess of KOHAq, yielding black products containing syrupy 'succinyl-propionic' acid C₆H₄O₄ and a crystalline acid C₆H₄O₄ [139°], which forms BaA'' 2aq The alkaline solution is turned brown by oxygen Does not react with phenyl cyanate (Goldschmidt & Meissler, *B* 23, 258)

Reactions—1 Converted by bromine into di-oxy terephthalic ether Bromine and potash yield brominated quinones —2 Nitrous acid forms a di nitroso-derivative C₆H₂(N₂O)₂ [114°], a white powder, insol water and alcohol, decomposed by boiling alcohol with formation of di-oxy-terephthalic ether, and by boiling water with formation of the oxim of pyruvic ether —3 Hydroxylamine in presence of NaOHAq form C₆H₄(NOH)₂CO₂Et, crystallising from alcohol in plates which begin to decompose at 160° (Jeaurenaud, *B* 22, 1282) An alcoholic solution of phenyl-hydrazine forms the white hydrazo-compound C₆H₄(NH NHPH₂)(CO₂Et)₂ [165°] whence bromine produces the diazo compound C₆H₄(N₂Ph)₂(CO₂Et)₂ [126°], which on saponification yields the acid C₆H₄(N₂Ph)₂(CO₂H)₂ [over 250°] The white hydrazo-compound is accompanied by C₆H₄(NH NHPH₂)(CO₂Et)₂ [208°], a yellow isomeride identical with Knorr's body [206°] (Baeyer, *B* 24, 2690) —4 Phenyl hydrazine (2 mols) forms in presence of toluene and some glacial acetic acid, the four following compounds C₆H₄(N,NHPh)(CO₂Et)₂ [160°], C₆H₄(N,NHPh)₂(CO₂Et)₂ [206°], C₆H₄N₂O₂ [212°], and C₆H₄N₂O₂, decomposing above 300° (Knorr & Bulow, *B* 17, 2654) The last body yields a di-methyl derivative C₆H₄(Me₂N)₂O₂ and is converted by nitrous acid into a blue substance C₆H₄N₂O₂ —5 PCl₅ forms the chlorides of dichloro dihydroterephthalic and dioxyterephthalic acids (Levy & Curchoff, *B* 22, 2106)

Salts—Na₂Et₂C₆H₄O₄2EtOH Got from the ether by ppg with alcoholic soda (Remsen, *B* 8, 1409) Red, on drying it changes to colourless Na₂Et₂C₆H₄O₄ (Hantzsch & Herrmann, *B* 21, 1756) —K₂Et₂C₆H₄O₄ Colourless —K₂Et₂C₆H₄O₄ orange —MgEt₂C₆H₄O₄ 2aq dark red amorphous mass —BaCl₂H₄O₄ aq red pp —CaCl₂H₄O₄ aq

Di-acetyl derivative of the ether C₆H₄(OAc)₂(CO₂Et)₂ [171°] Made from the ether and AcCl Needles (by sublimation) Its solutions are not fluorescent

Di-benzoyl derivative of the ether C₆H₄(OBz)₂(CO₂Et)₂ [165°] Formed from C₆H₄(ONa)₂(CO₂Et)₂ and BzCl (Nef, *A* 258, 810) Formed also, together with two isomerides [188°] and [102 5°], by reducing the di-benzoyl derivative of di-oxy-terephthalic ether with zinc and HClAq The three isomerides may be separated by crystallisation from ether-ligroin The three compounds are converted into C₆H₄(OH)₂(CO₂Et)₂ by H₂SO₄, and into C₆H₄(OBz)₂(CO₂Et)₂ by Br in CS₂.

(a)-**Di-benzyl derivative of the ether** C₆H₄(OCH₂Ph)₂(CO₂Et)₂ [169°] Made in small quantity (5 pc) from C₆H₄(OCH₂Ph)₂(CO₂Et)₂ by reducing with Zn and HCl (Nef, *A* 258, 801) Colourless needles May be sublimed

(β)-**Di-benzyl derivative of the ether** C₆H₄(OCH₂Ph)₂(CO₂Et)₂ [148 5°] Made, together with a (γ)-isomeride [140 5°], by the action of benzyl chloride on C₆H₄(ONa)₂(CO₂Et)₂ at 100° Less sol alcohol than the (γ)-isomeride H₂SO₄ converts it into a crystalline polymeride [272°] Hydroxylamine and phenyl hydrazine have no action

Di-ethyl derivative C₆H₄(OEt)₂(CO₂H)₂ [126 5°] Got, with EtBr, by heating bromo-ethyl acetate ether at 100° (Wedel, *A* 219, 104) Trimetric octahedra, *a* *b* *c* = 245.1 641 Sol water and alcohol, forming acid solutions, coloured red by FeCl₃ —(NH₄)₂A'' —Na₂A'' 2aq —K₂A'' aq —BaA'' 2aq —CaA'' aq —MgA'' 2aq —ZnA'' 3aq —MnA'' 4aq —PbA'' —Et.A''

Di-oxy-terephthalic acid tetrahydride C₆H₄O₄ & C₆H₄(OH)₂(CO₂H)₂ [191°] Made by the action of hydroxylamine on dioxyterephthalic ether (Jeaurenaud, *B* 22, 1279) Prisms, sl sol cold water, v e sol alcohol and ether FeCl₃ gives a brown colour —(NH₄)₂A'' needles —BaA'' white powder —Ag₂A'' 2aq needles

Ethyl ether Et.A'' [128°] Made by the action of hydroxylamine on the hydrate of di-oxyterephthalic ether [113°] (Hantzsch & Zeckendorf, *B* 20, 2801) Yelow crystalline body

Di-oxy-terephthalic acid hexahydride C₆H₄(OH)₂(CO₂H)₂ **Di oxy hexamethylene dicarboxylic acid** The salt BaA'' 3aq is got by saponifying the nitrile with baryta water (Baeyer & Noyes, *B* 22, 2177) Thick prisms or needles, v sl sol water

Nitrile C₆H₄O₄(CN)₂ [180°] Made from quinone tetrahydride (diketohexamethylene) and HCl Colourless, v sol hot water and alcohol

c-Di-oxy-terephthalic acid C₆H₄(OH)₂(CO₂H)₂ [1 2 3 4] [290°] Formed from sodium pyrocatechin and CO₂ (Schmitt & Hahle, *J pr* [2] 44, 1) Thin needles (containing aq) forming solutions with deep blue fluorescence FeCl₃ gives a blue colour —Na₂A'' 2aq fluorescent prisms —Pb₂C₆H₄O₄ —Ag₂A''

Methyl ether Me.A'' [145°] Needles

Ethyl ether Et.A'' [90°] Tables

Tetra-oxy terephthalic acid C₆H₄O₄ & C₆(OH)₂(CO₂H)₂ **Ethyl ether Et.A''** [178°] Formed by passing SO₂ into a feebly alkaline solution of di-oxy-quinone dicarboxylic ether (Loewy, *B* 19, 2388) Golden plates (from chloroform), v sl sol water, alcohol, and ether In presence of NaOHAq it is oxidised by air to di-oxy-quinone dicarboxylic ether (Böniger, *B* 22, 1284) Phenyl cyanate (4 mols) and some chloroform at 170° yield C₆(O CO NHPh)₂(CO₂Et)₂ [264°], an orange powder, insol most solvents (Goldschmidt & Meissler, *B* 23, 266) Ac₂O yields the tetra acetyl derivative C₆(OAc)₄(CO₂Et)₂ [202°], a colourless crystalline powder (Hantzsch & Zeckendorf, *B* 20, 2798) Hydroxylamine forms C₆(O NH₂OH)₂(CO₂Et)₂ [170°] Alcoholic phenyl-hydrazine forms, on boiling, the corresponding salt C₆O₂(O N₂H₂Ph)₂(CO₂Et)₂ [184°] (Böniger)

Reference—DI CHLORO-DI-OXY-TEREPHTHALIC ACID

DI OXY-TETRADECANE *v* BUTYRO PINACONE
and Di isobutyl-pinacone

DI OXY-TETRADECANOIC ACID $C_{14}H_{26}(OH)_2O_2$ [60°] A product of oxidation of oil of millet by $KMnO_4$ (Kassner *Ar Ph* [3] 25, 1081) Insol water, *v* sol alcohol and ether

OXYTETRIC ACID, formed by heating methyl acetoacetic ether with bromine and a little water, is identical with Mesaconic acid (Cloeze, *Bl* [3] 3, 598, *C R* 110, 583, Walden, *B* 24, 2033, Gorbhoff, *J R* 1887, 605)

o-OXY-THIENYL-ACETIC ACID

$C_6H_5S \cdot CH(OH)CO_2H$ [115°] Formed by reducing thienyl-glyoxylic acid with sodium amalgam (Ernst, *B* 19, 3280) Needles (from benzene), *v* sol water, alcohol, and ether Yields thiophenic aldehyde on boiling with MnO_2 Reduced by HI and P to thienyl acetic acid— $BA'A'$, *zaq*— $CA'A'$, *zaq* *v* sol water— Ag_2A' white pp

o-OXY THIOBENZOIC ACID $C_6H_5SO_2$ *ie*

$C_6H_4(OH)CO_2SH$ Made from *o*-oxy benzoyl chloride and KSH (Carius, *A* 129, 11) Brownish-yellow amorphous mass, insol water— $BA'A'$

Methyl derivative of the ethyl ether

$C_6H_4(OMe)CO_2SEt$ (198° at 80 mm) Made from $C_6H_4(OMe)CO_2OPh$ and $NaSEt$ in ether (Seifert, *J pr* [2] 31, 475) Oil, smelling like mercaptan

o Oxy-thiobenzoic acid $C_6H_4(OH)CSOH$

Amide $C_6H_4(OH)CSNH_2$ [118°] Made by fusing salicylamide with P_2S_5 (Spilker, *B* 22, 2767) Colourless needles, *v* sol alcohol $FeCl_3$ colours its aqueous solution violet Slowly converted into $C_6H_4(OH)CONH_2$ by boiling water

Di oxy dithio benzoic acid $C_6H_4(OH)_2CS_2H_2$ [4 2 1] Made by heating resorcin with potassium xanthate at 100° (Lippmann, *M* 10, 618) Yellow needles (containing aq) Melts at 131° when hydrated, but decomposes at 124° when anhydrous Potash fusion yields (4,2,1)-di-oxy-benzoic acid

Tri oxy di-thio-benzoic acid

$C_6H_4(OH)_3CS_2H_2$ [4 3 2 1] [154°] Made by heating pyrogallol with potassium xanthate and alcohol (L) Yellow crystals (containing aq), *v* sol alcohol and ether Yields *c*-tri-oxy benzoic acid when fused with potash

OXY THIONAPHTHENE $C_{10}H_8SO$ *ie*

$CH \cdot C(=O)CH \cdot CH$ [72°] Made by heating thiophenic aldehyde with sodium succinate and Ac_2O at 135° (Biedermann, *B* 19, 1618) Needles (by sublimation), *sl* sol water, *v* sol $NaOHAq$ Gives the indophenine reaction Chloroform and $KOHAq$ give a bluish-green colour on warming

OXY THYMOQUINONE $C_{11}H_{10}O_2$ *ie*

$C_6HMe(C_6H_4)(OH)O_2$ [1 4 6 2 5] [165°]

Formation—1 By the oxidising action of $FeCl_3$ on di-amido-thymol (Carstanjen, *J pr* [2] 15, 399, Ladenburg a Engelbrecht, *B* 10, 1218) 2 By dissolving bromo thymoquinone in $KOHAq$ (Carstanjen, *J pr* [2] 8, 57)—3 By the action of H_2SO_4 or $HClAq$ on methylamidothymoquinone (Zincke, *B* 14, 97) or dimethylamidothymoquinone (Schulz, *B* 16, 898)

Properties—Yellow needles, *sol*. alcohol, ether, and hot water Forms a violet red solution in alkalis May be sublimed

Reactions—1 EtI at 100° forms $C_{11}H_{11}EtO_2$

which sublimes in golden plates—2 *Aniline* in alcohol forms $C_6HMe(C_6H_4)(OH)O_2(NHPh)$ [135°], crystallising in violet-black needles, and forming a bluish violet solution in NH_4Aq —3 *p-Tolu idine* yields the corresponding *p*-toluide [165°]

(8) **Oxy-thymoquinone**

$C_6HMe(C_6H_4)(OH)O_2$ [1 4 3 2 5] [183°] Made from di nitro carvacrol by reduction and subsequent oxidation with $FeCl_3$ (Mazzara, *B* 23, 1390) Orange prismatic tables, volatile with steam Na_2CO_3Aq forms a violet solution

Di-oxy thymoquinone $C_6Me(C_6H_4)(OH)_2O_2$

[213°] Formed by boiling chloro-oxy thymoquinone with $KOHAq$ (E a L) and by boiling dimethylamido thymoquinone with alcohol and H_2SO_4 or KOH (Zincke, *B* 14, 95) Red needles or prisms, forming a violet solution in alkalis Phenylene phenyl *o* diamine, in presence of alcohol and $HOAc$, forms oxythymophenindulone

$C_6H_4 \cdot \langle \overset{N}{NPh} \rangle \cdot C_6Me(C_6H_4)(OH)O_2$ as ruby-red crystals [175°] (Kehrmann a Messinger, *B* 24, 590) — $BA'A'$ *aq*— PbA' green pp

Di-acetyl derivative [81°] Needles

Di-benzoyl derivative [163°]

Reference—CHLORO OXY THYMOQUINONE

DI OXY-DITHYMYL ETHANE $C_{22}H_{20}O_2$ *ie*

$CH_3CH(C_6H_4)_2(OH)_2$ [185°] Formed by reducing $CCl_2CH(C_6H_4)_2OH$, with zinc-dust and alcohol (Jager, *C J* 31, 262) and also by adding a mixture of chloroform and $SnCl_4$ to a cooled mixture of paraldehyde and thymol (Steiner, *B* 11, 287) Efflorescent plates (from alcohol)

Di acetyl derivative [100°] Needles

Di benzoyl derivative [191°] Needles

Di ethyl ether $C_{22}H_{20}Et_2O_2$ [72°]

DI OXY-DI THYMYL-ETHYLENE

$CH_3C(C_6H_4)_2(OH)_2$ [171°] Formed, together with the preceding body, by boiling the compound $CCl_2CH(C_6H_4)_2OH$, with zinc dust and alcohol (Jager, *C J* 31, 263) Needles (from $HOAc$) K_2FeCy_4 oxidises it, in alcoholic solution, to $C_6H_4O_2$ [215°], crystallising in green needles Alkaline K_2FeCy_4 forms $C_{22}H_{20}O_2$ [215°], which forms dark-red crystals

DI-OXY DI-THYMYL SULPHIDE

$(C_6H_4)_2(OH)_2S$ [152°] Formed from thymol and $SOCl_2$ (Tassinari, *G* 17, 92)

OXY-TOLUAMIDOXIM

[3 6 1] $C_6H_4Me(OH)C(NO_2)NH_2$ [124°] Formed from the thioamide in alcohol by treatment with hydroxylamine hydrochloride and $NaOHAq$ (Goldbeck, *B* 24, 3662) Yellowish plates (from benzene), *v* sol hot water Coloured cherry red by $FeCl_3$ — $B'HCl$ [215°]

Acetyl derivative Plates (from benzene) gives $C_6H_4(OH)C \cdot \langle \overset{N}{N}O \rangle \cdot CMe$ [45°] on heating with water in a sealed tube

Benzoyl derivative

$C_6H_4(OH)C(NO_2)NH_2$ [182°] Plates, *sl* sol alcohol Coloured green by $FeCl_3$ in acetone

Di-benzoyl derivative [143°]

c-Oxy-toluamidoxim

[3 2 1] $C_6H_4Me(OH)C(NO_2)NH_2$ [126 5°] Formed from the nitrile and hydroxylamine (Paschen, *B* 24, 3670) Tables, *v* sol alcohol and hot water Yields a dibenzoyl derivative $C_6H_4Me(OBz)C(NO_2)NH_2$ [164°], whence KOH forms $C_6H_4Me(OH)C \cdot \langle \overset{N}{N}O \rangle \cdot CPh$ [150°]

OXY-TOLUENE v. CRESOL and BENZYL

ALCOHOL

Di-oxy-toluene $C_8H_8O_2$, $\pm e$

$C_8H_7Me(OH)_2$ [124] *Cresorein Lutorcin* Mol w 124 [104°] (c 269°) Formed from amido *p*-cresol and from amido *o*-cresol [161°] by the diazo reaction (Knecht, *B* 15, 92, *B* 15, 298, Wallach, *B* 15, 2835) Formed also by potash-fusion from the corresponding toluenedisulphonic acid (Nolting, *B* 19, 136) and from bromo *p*-cresol (Vogt & Henninger, *C R* 94, 650) Spherical crystalline groups, v sol water, alcohol, and ether, sl sol benzene and ligroin $FeCl_3$ colours its solutions blue NH_3 with damp air turns it brown, green, and finally blue Bleaching powder gives a yellow colour Its solution in NaOH aq is turned brown by air On heating with chloroform and NaOHAq it gives a rose red colour On heating with phthalic anhydride it forms the phthalein $C_{22}H_{14}O_6$, which is a brick red powder, exhibits green fluorescence in alkaline solution, and yields $C_{22}H_{14}Ac_2O_6$ [260°] Unlike resorcin, cresorein gives no colour when the product got by heating with H_2SO_4 and nitro-benzene is diluted and made alkaline

Di-acetyl derivative $C_8H_8(OAc)_2$ [c 160°]

Di-oxy-toluene $C_8H_7Me(OH)_2$ [126] [63°-66°] Made by the diazo reaction from amido cresol [124°-128°] (Ullmann, *B* 17, 1960) Needles, v sol water and alcohol Coloured red by bleaching powder With phthalic anhydride it gives a compound analogous to fluorescein Reduces cold ammoniacal $AgNO_3$

Isocrescin $C_8H_8(OH)_2$ [87°] (260°) Made by potash fusion from toluene (γ) disulphonic acid (Senhofer, *A* 164, 181) Needles (containing aq) Sol water, alcohol, and ether Tastes sweet $FeCl_3$ gives a brownish green colour Reduces cold ammoniacal $AgNO_3$

Other di-oxy-toluenes are described as Hydrotoluquinone, Isohydrotoluquinone, Orcin, and Methyl pyrocatechin

***o*-OXY-TOLUENE PHOSPHINIC ACID**

$C_8H_7CH(OH)P(OH)_2$ [90°] Formed, together with $(C_8H_7CH(OH))_2PO_3H$ [165°] by heating benzoic aldehyde with hypophosphorous acid (Ville, *C R* 107, 659, 110, 348) Plates, decomposing at 140° with formation of benzoic aldehyde Reduces $AgNO_3$ — $Ba(HA'')_2$ aq small plates

Acetyl derivative $CHPh(OAc)P(OH)_2$ ***o*-OXY-TOLUENE PHOSPHONIC ACID**

$C_8H_7CH(OH)PO(OH)_2$ [173°] Made from benzoic acid by successive treatment with PCl_5 and water (Fossek, *M* 7, 34) Hard crusts (from benzene and HOAc)— BaA'' — $Ba(HA'')_2$

OXY-TOLUENE SULPHONIC ACID v**CRESOL SULPHONIC ACID****OXY *o*-TOLUIC ACID** $C_8H_8O_3$, $\pm e$

$C_8H_7Me(OH)CO_2H$ [231] *Oxy methyl-benzoic acid* Mol w 152 [183°] Formed by the diazo-reaction from amido *o*-toluic acid [191°], and by potash-fusion from sulpho *o*-toluic acid (Jacobsen, *B* 16, 1963, 17, 163) Needles, sol cold water, alcohol, and ether, sl sol chloroform Volatile with steam $FeCl_3$ gives a brown pp

Methyl derivative $C_8H_7Me(OMe)CO_2H$ [146°] Needles, sl sol cold water— CaA' , 2aq

Oxy-*o*-toluic acid $C_8H_7Me(OH)CO_2H$ [241] [179°] *Formation*—1 By potash fusion from the corresponding aldehyde (Tiemann & Schotten, *B* 11, 778), and from the corresponding

$C_8H_7Me(SO_2NH_2)CO_2H$ (Jacobsen, *B* 14, 40)—2 By heating *m*-cresol with CCl_4 and alcoholic NaOH (Schall, *B* 12, 819)—3 From amido-*o*-toluic acid [c 165°] by the diazo reaction (Jacobsen, *B* 17, 164)

Properties—Needles (containing $\frac{1}{2}$ aq), v sol hot water, alcohol, and ether, insol cold chloroform Not coloured by $FeCl_3$ Yields *m*-cresol on distillation— CaA' , 2aq crystals, v sol water

Methyl derivative $C_8H_7Me(OMe)CO_2H$ [176°] Needles (from water)

Oxy-*o*-toluic acid $C_8H_7Me(OH)CO_2H$ [251] [172°] Formed by potash fusion from the corresponding $C_8H_7Me(SO_2NH_2)CO_2H$, and also from amido *o*-toluic acid [196°] by the diazo reaction (Jacobsen, *B* 14, 41, 17, 163) Prisms, sol alcohol, ether, and hot water, v sl sol chloroform Volatile with steam $FeCl_3$ gives a brown pp The Cu salt forms bluish green plates

Oxy-*o*-toluic acid $C_8H_7Me(OH)CO_2H$ [261] [168°] S 143 at 25° Formed by potash-fusion from bromo *o*-toluic acid (Jacobsen, *B* 16, 1962) Long needles, v sol hot water, alcohol, and ether Volatile with steam At 200° it yields *m*-cresol $FeCl_3$ gives a bluish violet colour

Oxy-*m*-toluic acid $C_8H_7Me(OH)CO_2H$ [221] *Homosalicylic acid* (β) *Cresotic acid* [164°]

Formation—1 By the action of CO_2 on sodium *o*-cresol (Engelhardt & Latschneff, *Z* 1869, 623, Kekulé, *B* 7, 1006, Ihle, *J pr* [2] 14, 456)—2 By heating *o*-cresol with CCl_4 and NaOH at 100° (Schall, *B* 12, 86)—3 By means of potash fusion from the corresponding $C_8H_7Me(SO_2NH_2)CO_2H$ (Jacobsen, *B* 11, 902)—4 From *c*-amido toluic acid (Jacobsen, *B* 14, 2354)

Properties—Long needles, sl sol cold water, v sol chloroform Volatile with steam $FeCl_3$ gives a violet colour Poisonous (Dunstan & Black, *Ph* [3] 21, 429)

Methyl derivative $C_8H_7(OMe)CO_2H$ [81°] Feathery needles

Ethyl ether MeA' (235°) SG $\frac{33}{3}$ 1 1444 (Pinner, *B* 23, 2938)

Ethyl ether EtA' (248°) SG $\frac{33}{3}$ 1 1020 With benzamide it forms $C_{22}H_{11}N_2O$ crystallising in yellowish needles, insol acids and alkalis (Pinner, *B* 23, 2939)

Nitrile $C_8H_7(OH)CN$ [885°] Got from its acetyl derivative, which is formed by the action of Ac_2O on the oxim of *c*-toluic aldehyde (Paschen, *B* 24, 3669) Tables, v sol alcohol

Oxy-*m*-toluic acid $C_8H_7Me(OH)CO_2H$ [341]

Formation—1 By potash fusion from the corresponding $C_8H_7Me(SO_2NH_2)CO_2H$ (Jacobsen, *B* 11, 897, Remsen & Iles, *Am* 1, 87, *B* 11, 462, Mahon, *Am* 4, 186), from bromo *m*-toluic acid and from chloro *m*-toluic acid [210°] (Jacobsen, *B* 14, 2351, 18, 1761)—2 By heating *o*-cresol with CCl_4 and NaOH, the yield being 40 p c (Schall, *B* 12, 816)—3 By the diazo reaction from amido *m*-toluic acid (Remsen & Kuhara, *Am* 8, 428)—4 By oxidation of the corresponding aldehyde (Tiemann & Schotten, *B* 11, 777)

Properties—Long needles (containing $\frac{1}{2}$ aq), sol hot water, alcohol, and ether, insol cold chloroform Gives no colour with $FeCl_3$ Yields *o*-cresol when heated with HCl aq at 180° Gives a nitro-compound [85°].

Salts — CaA' , 3aq minute needles — BaA' , 3aq — CuA' , 1½aq blue flocculent pp

Methyl derivative $\text{C}_6\text{H}_4\text{Me}(\text{OMe})\text{CO}_2\text{H}$ [193°] Minute needles Got by saponifying $\text{C}_6\text{H}_4\text{Me}(\text{OMe})\text{CO}_2\text{Me}$ [67°] (Schall) The amide $\text{C}_6\text{H}_4\text{Me}(\text{OMe})\text{CONH}_2$ [144°] is prepared from $\text{C}_6\text{H}_4\text{Me}(\text{OMe})$ [1 q], ClCONH_2 , and AlCl_3 (Gattermann, *A* 244, 64)

Ethyl derivative $\text{C}_6\text{H}_4\text{Me}(\text{OEt})\text{CO}_2\text{H}$ [199°] Obtained by saponification of the amide $\text{C}_6\text{H}_4\text{Me}(\text{OEt})\text{CONH}_2$ [167°] which is got from $\text{C}_6\text{H}_4\text{Me}(\text{OEt})$ and ClCONH_2 (G) — CaA' , 2aq (Brown, *Am* 4, 375)

s Oxy m toluic acid

$\text{C}_6\text{H}_3\text{Me}(\text{OH})\text{CO}_2\text{H}$ [3 5 1] [208°] Formed from m toluic acid by sulphonation followed by potash fusion, the resulting acid being freed from its isomeride by steam distillation (Jacobsen, *B* 14, 2357) Formed also by the action of baryta water on $\text{C}_{12}\text{H}_{10}\text{O}_8$ [90°], which is obtained from acetyl pyruvic ether and NaOHaq (Claisen, *B* 22, 3271) Needles or prisms, v sol hot water, not volatile with steam — CaA' , 2aq prisms — SrA'_2

Methyl ether MeA' [93°] Needles.

Oxy m toluic acid

$\text{C}_6\text{H}_3\text{Me}(\text{OH})\text{CO}_2\text{H}$ [3 6 1] *p*-Homosalicylic acid

(a) *Cresotic acid* [151°]

Formation — 1 By passing CO_2 over *p* cresol in which sodium has been dissolved (Engelhardt a La'schmoff, *Z* 1869, 622, 712, Kolbe a Lautemann, *A* 115, 203, *Ihle, J pr* [2] 14, 455) 2 By heating *p* cresol with CCl_4 and NaOH (Schall, *B* 12, 816) — 3 By potash fusion from chloro m xylene sulphonic acid (Voigt, *Z* [2] 5, 577, *B* 2, 284), from m xylene sulphonic acid (E a L), from (1,3,4) xylenol or its (8) sulphonic acid, from bromo m toluic acid, and from sulpho-m toluic acid (Jacobsen, *B* 11, 374, 14, 2352, Staedel a Holz, *B* 13, 2919) — 4 By oxidation of the corresponding aldehyde (Tiemann a Schotten, *B* 11, 778) — 5 From amido toluic acid [172°] by the diazo reaction (Panaotovic, *J pr* [2] 33, 64)

Properties — Colourless trimetric prisms (from water), sl sol water, v sol alcohol, ether, and chloroform Volatile with steam Its aqueous solution is turned violet by FeCl_3 , Conc HClaq at 180° yields *p* cresol POCl_3 , forming various dehydration products (Schiff, *A* 245, 43) Distillation with As_2O_3 yields $\text{C}_{12}\text{H}_{10}\text{O}_8$ [148°] (Kostanecki, *B* 18, 1988)

Salt — BaA' , 2aq leaflets

Methyl ether MeA' (242°) SG $\frac{2}{3}$ 1 1438 Liquid (Pinner, *B* 23, 2938)

Ethyl ether EtA' (251°) SG $\frac{2}{3}$ 1 1037 Reacts with benzamidine, forming $\text{C}_{12}\text{H}_{11}\text{N}_3\text{O}$ crystallising in yellowish needles (P)

Amide $\text{C}_6\text{H}_3\text{Me}(\text{OH})\text{CONH}_2$ [178°] Got from the acid by treatment with POCl_3 at 60° followed by alcoholic NH_3 at 160° (Schiff, *G* 17, 559, *A* 245, 43), and from the ether and NH_3 (Goldbeck, *B* 24, 3659) Needles (from alcohol) P_2O_5 forms $\text{C}_6\text{H}_3(\text{OH})\text{CSNH}_2$ [127°]

Anilide $\text{C}_6\text{H}_3(\text{OH})\text{CONHPh}$ [53°] Plates

Nitrile $\text{C}_6\text{H}_3(\text{OH})\text{CN}$ v sol alcohol Its acetyl derivative [57°] is got by the action of Ac_2O on the oxim of toluic aldehyde

Methyl derivative $\text{C}_6\text{H}_3\text{Me}(\text{OMe})\text{CO}_2\text{H}$ [70°] (L), [67°] (Schall, *B* 22, 749) Long needles — AgA'

Amide of the methyl derivative $\text{C}_6\text{H}_3\text{Me}(\text{OMe})\text{CONH}_2$ [163°] Made from $\text{C}_6\text{H}_3\text{Me}(\text{OMe})$ and ClCONH_2 in presence of AlCl_3 (Gattermann, *A* 244, 66) Needles (from hot Aq)

Nitrile of the methyl derivative $\text{C}_6\text{H}_3\text{Me}(\text{OMe})\text{CN}$ (270° uncor) Made by warming diazotised amido *p* cresol with CuCy , (Limpach, *B* 22, 351) Oil

Amide of the ethyl derivative $\text{C}_6\text{H}_3\text{Me}(\text{OEt})\text{CONH}_2$ [152°] Made from $\text{C}_6\text{H}_3\text{Me}(\text{OEt})$, ClCONH_2 , and AlCl_3 (G) Silky needles (from dilute alcohol)

Oxy p toluic acid $\text{C}_6\text{H}_3\text{Me}(\text{OH})\text{CO}_2\text{H}$ [4 2 1].

(7) *Cresotic acid m-Homosalicylic acid* [177° cor] **Formation** — 1 By passing CO_2 into *m* cresol in which sodium has been dissolved (E a L, Biedermann a Pike, *B* 6, 323) — 2 By oxidation of the corresponding aldehyde (Tiemann a Schotten, *B* 11, 777) — 3 By heating *m* cresol with CCl_4 and NaOHaq (Schall, *B* 12, 816) — 3 By potash fusion from *p* xylenol (Jacobsen, *B* 11, 570)

Properties — Needles (from water) or monoclinic prisms (from alcohol), m sol hot water, v sol alcohol and chloroform Volatile with steam FeCl_3 gives a red colour HClaq at 170° yields *m* cresol

Salts — CaA' , 3aq — BaA' , 3aq prisms

Methyl ether MeA' (243°) SG $\frac{2}{3}$ 1 1395 (Pinner, *B* 23, 2938)

Ethyl ether EtA' (254°) SG $\frac{2}{3}$ 1 0973 Converted by benzamidine into $\text{C}_{12}\text{H}_{11}\text{N}_3\text{O}$ crystallising in needles [235°]

Methyl derivative $\text{C}_6\text{H}_3\text{Me}(\text{OMe})\text{CO}_2\text{H}$ [104°] Leaflets, sol hot water (Schall)

Ethyl derivative $\text{C}_6\text{H}_3\text{Me}(\text{OEt})\text{CO}_2\text{H}$ [109°] Got by oxidising the ethyl derivative of thymol (Paterno, *J* 1879, 519)

Oxy p toluic acid $\text{C}_6\text{H}_3\text{Me}(\text{OH})\text{CO}_2\text{H}$ [4 3 1] [207° cor]

Formation — 1 By potash fusion from sulpho *p* toluic acid or its mono amide (Flesh, *B* 6, 481, Remsen a Hall, *Am* 2, 54, Weinreich, *B* 20, 981), and from chloro or bromo *p* toluic acid (Gerichten, *B* 11, 368) — 2 From nitro-toluic acid [190°] by reduction, followed by the diazo reaction (Fittica, *B* 7, 927)

Properties — Needles, sol hot water, alcohol, and ether, insol chloroform Volatile with steam Not coloured by FeCl_3 Gives *o* cresol on distillation with lime

Salts — CaA' , 4aq — PbA' , 2aq needles

Ethyl ether EtA' [75°] Prisms

Methyl derivative $\text{C}_6\text{H}_3\text{Me}(\text{OMe})\text{CO}_2\text{H}$ [156°] Needles (by sublimation) — BaA' , 4aq

o Oxy o toluic acid $\text{CH}_3(\text{OH})\text{C}_6\text{H}_3\text{CO}_2\text{H}$ [118°] Made by dissolving phthalide in boiling NaOHaq (Hessert, *B* 10, 1446, 11, 237) Powder, v sl sol cold water, v sol alcohol and ether Changes on fusion, and on boiling with water, into its anhydride — BaA' , — AgA' octahedra

Anhydride v PHTHALIDE

o Oxy p toluic acid $\text{CH}_3(\text{OH})\text{C}_6\text{H}_3\text{CO}_2\text{H}$ *Oxymethyl-benzoic acid* [181°] Formed, together with terephthalic acid and $\text{C}_6\text{H}_3(\text{CH}_3\text{OH})$, by boiling terephthalic aldehyde with NaOHaq (Löw, *A* 231, 373) Formed also by boiling CH_3Br , $\text{C}_6\text{H}_3\text{CO}_2\text{H}$ with baryta-water (Kekulé a Dittmar, *A* 162, 342) Plates or needles, v sol water and ether Not reddened or resublimed by H_2SO_4 — AgA'

Di-oxy-o-toluic acid $C_6H_4Me(OH)_2CO_2H$ [3 5 3 1] *Cresorsellic acid* Mol. w 168 [145°] S 86 at 0° Formed by fusing di sulpho o-toluic acid with potash (Jacobsen, *B.* 16, 1960) Long needles, v sol alcohol and boiling water $FeCl_3$ gives a brownish-black colour Reduces alkaline Ag and Cu solutions Conc H_2SO_4 gives a bright red colour on heating $-NH_4A'$ 2aq prisms

Di oxy-toluic acid *Methyl derivative* $C_6H_4Me(OMe)(OH)CO_2H$ [1 3 4 5] *Creosol carboxylic acid* [182°] Obtained by the action of CO_2 upon sodium creosol (Wende, *B.* 19, 2324) Concentric needles Sublimable V sol alcohol, ether, and chloroform, sl sol water, nearly insol benzene Gives a blue colouration with $FeCl_3$

Salts — NH_4A' needles — KA' small readily soluble needles — BaA'_2 small needles — PbA'_2 white pp — CuA'_2 yellow powder

Methyl ether MeA' [92°], small colourless trimetric crystals, $abc = 5285 \ 1 \ 0 \ 7384$ Gives a bluish green colour with $FeCl_3$

Ethyl ether EtA' [77°], small needles

Di oxy-m-toluic acid $C_6H_4Me(OH)_2CO_2H$ [1 2 4 5] *Cresorcin carboxylic acid* [208°] Made by boiling cresorein (di oxy toluene) (1 pt) with $NaHCO_3$ (4 pts) and water (8 pts) (Kostanecki, *B.* 18, 3203) Prisms (containing aq), v e sol water — KA'_2 2aq prisms, v sol water

Di - oxy p-toluic acid $C_6H_4Me(OH)_2CO_2H$ [4 5 3 1] [176°] Formed by heating disulpho p-toluic acid with KOH (Weinreich, *B.* 20, 981) Needles, v sol water, alcohol, and ether

Di - oxy - toluic acid $C_6H_4Me(OH)CO_2H$ *Hydrotoluquinone carboxylic acid* [206°-210°] S 07 at 8° Formed by heating hydrotoluquinone with $KHCO_3$ and conc K_2SO_4 aq at 160° (Brunner, *M.* 2, 458) Trimetric plates (from alcohol) or indistinct crystals (containing $\frac{1}{2}$ aq) (from water), m sol hot water and alcohol $FeCl_3$ gives a blue colour Reduces ammoniacal $AgNO_3$ in the cold Yields hydrotoluquinone on fusing Conc H_2SO_4 at 160° yields tetra oxy di methyl anthraquinone, a dark red powder — CaA'_2 2aq — BaA'_2 2aq — PbA'_2 2aq crystalline *Ethyl ether EtA'* [98°] Needles

Di oxy p-toluic acid $C_6H_4Me(OH)_2CO_2H$ *Orselllic acid* *Orsellinic acid* *Orsellitic acid* [176°] S (ether) 22 at 20° Formed by boiling lecanoric acid or erythrin with lime water, baryta water, or $NaOHAq$ (Stenhouse, *P.M.* [3] 32, 300, *Pr.* 12, 263, *A.* 68, 61, Hesse, *A.* 117, 312, 139, 35) Prisms (containing aq), v e sol alcohol, sol water Splits up on fusion, or on boiling with water, into CO_2 and orcin $FeCl_3$ gives a purple colour Bromine gives tri bromo orcin $POCl_3$ acting on the anhydrous acid at 90° forms a blue liquid which, when poured into water, ppts $C_{10}H_8P_2O_{11}$, a blue substance with coppery lustre resembling indigo The solutions of this 'phosphorsellic acid' in water and alcohol are intensely blue, and it is pptd from aqueous solution by HCl , by H_2SO_4 , and by $NaCl$ Phosphorsellic acid forms $C_{10}H_8Pb_2P_2O_{11}$ and $C_{10}H_8Pb_2P_2O_{11}$, an anilide $C_{10}H_8(NHPh)_2P_2O_{11}$ which gives $C_{10}H_8Pb_2(NHPh)_2P_2O_{11}$, and an acetyl derivative $C_{10}H_8Ac_2P_2O_{11}$ as a dark violet mass which gives $C_{10}H_8(PbOH)_2Ac_2P_2O_{11}$ (Schiff, *A.* 228, 56)

Salt — $Ba(C_6H_3O)_2$ 2aq prisms

Methyl ether MeA' Made by boiling

lecanoric acid or erythrin with $MeOH$ Silky needles (from boiling water)

Ethyl ether EtA' [182°] Got in like manner (Heeren, *Schw.* *J.* 59, 841, Liebig, *P.* 21, 32, Kane, *Tr.* 1840, 237, 279, Schunck, *A.* 41, 160, 61, 72, Rochleder & Heldt, *A.* 48, 5, Stenhouse, *C.* *J.* 20, 224, Strecker, *A.* 68; 111, Hesse, *A.* 117, 297) May be prepared by repeatedly exhausting *Roccella tinctoria* with boiling alcohol and evaporating the extract Thin needles, nearly insol cold water, v sol alcohol and alkalis Yields $C_6Cl_2Me(OH)_2CO_2Et$ [162°] on chlorination

Isoamyl ether C_5H_{11}A' [76°] Prisms

Di oxy - o - toluic acid $C_6H_4Me(OH)_2CO_2H$ [6 4 2 1] *Paraorsellic acid* [172°] S 17 in the cold Formed by heating orcin with a solution of ammonium carbonate, $KHCO_3$, or $NaHCO_3$ (Senhofer a Brunner, *M.* 1, 236, Bistrzycki a Kostanecki, *B.* 18, 1986), or by heating potassium orcin in a current of CO_2 at 250° (Schwarz, *B.* 13, 1643) Needles (containing aq), sl sol water, v sol alcohol and ether $FeCl_3$ gives a blue colour Boiling water decomposes it into CO_2 and orcin $POCl_3$ followed by water gives $P_2(C_6H_3O)_6$, a chrome-green powder (Schiff, *A.* 228, 56) — KA' — BaA'_2 6aq — $Ba_2(C_6H_3O)_6$ 8aq — CuA'_2 4aq — AgA'_2 nearly insol water

Di oxy-m-toluic acid

$CH_3(OH)C_6H_4(OH)CO_2H$ [3 2 1] [142°] Formed by reduction of aldehyde o-oxy-benzoic acid with sodium-amalgam (Reimer, *B.* 11, 792) Prisms, v sol water, alcohol, and ether $FeCl_3$ gives a violet colour H_2SO_4 gives a red colour Boiling HCl aq resinifies it

Di oxy-p-toluic acid

$CH_3(OH)C_6H_4(OH)CO_2H$ [5 2 1] [above 270°] Formed by reduction of aldehyde p oxy benzoic acid (R) White powder, sl sol water Not coloured by $FeCl_3$, or by H_2SO_4

Di oxy m-toluic acid

$CH_3(OH)C_6H_4(OH)CO_2H$ [3 6 1] Made by reducing aldehyde o oxy benzoic acid (R) Prisms (from ether), sl sol water, m sol alcohol and ether $FeCl_3$ gives a violet colour Oxidised by $KMnO_4$ to oxy isophthalic acid, and by aqueous $K_2Cr_2O_7$ to aldehyde oxy benzoic acid

References — **DI-BROMO OXY-TOLUIC ACID** and **DI-iodo-ORSELLIC ACID**

OXY-O-TOLUIC ALDEHYDE $C_6H_4O_2$ 1 e $C_6H_4Me(OH)CHO$ [2 4 1] [110°] Prepared, together with the (4,2,1) isomeric, by heating m cresol with chloroform and $NaOHAq$ (Tiemann a Schotten, *B.* 11, 773) Flat leaflets (from water), not volatile with steam $FeCl_3$ gives a red colour

Phenyl hydrasile

$C_6H_4Me(OH)CHN_2HPh$ [151°] (Paschen, *B.* 24, 3671)

Oxy m-toluic aldehyde $C_6H_4Me(OH)CHO$ [3 4 1] [115°] Formed, together with the (3,2,1) isomeric, by the action of chloroform and potash on o cresol (Tiemann a Schotten, *B.* 11, 772) Long prisms (from water), not volatile with steam $FeCl_3$ gives a bluish violet colour Yields a nitro-derivative [152°]

Acetyl derivatives $C_6H_4(OAc)CHO$ [40°] (275°) Needles (Staats, *B.* 13, 138, Barbier, *Bl.* [2] 33, 52, *C.R.* 90, 37).

Oxy-m-toluic aldehyde $C_6H_4Me(OH)CHO$ [3 2 1] [17°] (209°) Formed as above (T. a S.).

Crystals, volatile with steam. FeCl_3 gives a bluish colour. Forms a yellow solution in NH_4Aq .

Acetyl derivative $\text{C}_9\text{H}_9\text{O}_2$, [267°] (B).

Oxism. [98°] (Paschen, B 24, 3668)

Phenyl hydrazide [95°] Tables

Oxy-m-toluic aldehyde $\text{C}_8\text{H}_7\text{Me}(\text{OH})\text{CHO}$ [8 6 1] [56°] [218°] Formed by the action of CHCl_3 and KOH on *p*-cresol (T & S). White leaflets. FeCl_3 gives a green colour. Yields a nitro-derivative [141°]

Acetyl derivative $\text{C}_8\text{H}_7\text{Me}(\text{OAc})\text{CHO}$ [57°] Long needles. Made by the action of Ac_2O on the ethereal solution of the K salt. Combines with NaHSO_3 . With Ac_2O it unites in the cold, forming $\text{C}_8\text{H}_7\text{Me}(\text{OAc})\text{CH}(\text{OAc})_2$, which crystallises in prisms [94°] and does not combine with NaHSO_3 .

Methyl derivative $\text{C}_8\text{H}_7\text{Me}(\text{OMe})\text{CHO}$ (254°) Colourless liquid

Oxism. $\text{C}_8\text{H}_7\text{Me}(\text{OH})\text{CH NOH}$. [105°] Needles (Goldbeck, B 24, 3658)

Oxy-p-toluic aldehyde $\text{C}_8\text{H}_7\text{Me}(\text{OH})\text{CHO}$ [4 2 1] [54°] [223°] Made, together with the (2,4,1)-isomere, by the action of chloroform and potash on *m*-cresol (T & S). Crystals, volatile with steam. FeCl_3 gives a violet colour.

Di-oxy o-toluic aldehyde $\text{C}_8\text{H}_6\text{Me}(\text{OH})_2\text{CHO}$ [6 4 2 1] *Orcyl aldehyde* [178°] Formed by heating orcin with chloroform and potash (The mann & Helkenberg, B 12, 999), and also by potash-fusion from methyl umbelliferone (Pechmann & Welsh, B 17, 1646). Needles, sol alcohol and hot water. FeCl_3 gives a reddish-brown colour. Aniline forms the compound $\text{C}_8\text{H}_6\text{Me}(\text{OH})_2\text{CH NPh}$ [126°], crystallising in yellow prisms.

Di-oxy m-toluic aldehyde *Methyl derivative* $\text{C}_8\text{H}_6\text{Me}(\text{OH})(\text{OMe})\text{CHO}$ [5 2 3 1] [270°-275°] Formed by the action of chloroform and NaOH upon cresol (Tiemann & Koppe, B 14, 2026). Oil, sol alcohol. FeCl_3 colours its alcoholic solution green.

OXY-TOLUIDINE v AMIDO CRESOL

OXY-TOLUQUINOLINE v OXY-METHYL-QUINOLINE

DI-OXY-TOLUQUINONE $\text{C}_9\text{H}_6\text{Me}(\text{OH})_2\text{O}_2$ [177°] Made by digesting oxy phenylamido-toluquinone anilide with dilute KOH (Hagen & Zincke, B 16, 1562). Brownish yellow plates, v sol most solvents. May be sublimed.

Tri-oxy toluquinone $\text{C}_9\text{H}_5\text{Me}(\text{OH})_3\text{O}_3$ Formed from tri-amido-orcin by treatment with FeCl_3 , the resulting amido dimido-orcin being heated with HClAq at 150° (Merz & Zetter, B 12, 2044). Dark crystals, nearly insol cold alcohol. Yields a crystalline tri acetyl derivative.

Reference — *Di-bromo-* and *Di-chloro-* **OXY-TOLUQUINONE**

OXY-TOLUQUINOXALINE

[8] $\text{C}_8\text{H}_6\text{Me} \begin{smallmatrix} \text{N COH} \\ \text{N OH} \end{smallmatrix}$ [267°] Formed by oxidation of its dihydride [o 127°], which is got by condensing tolylene diamine with chloroacetic ether (Hinsberg, B 18, 2870, A 248, 76), or by reducing nitro-tolyl amido acetic acid (Plöchl, B 19, 6). Yields a methyl derivative [71°], and an ethyl derivative [67°].

Isomeric v **OXY METHYL-QUINOXALINE**

Di-oxy-toluquinoxaline $\text{C}_8\text{H}_5\text{Me} \begin{smallmatrix} \text{N COH} \\ \text{N COH} \end{smallmatrix}$ [above 300°] Formed by heating tolylene-c

diamine with oxalic acid at 150° (Hinsberg, B 18, 2690, A 287, 348) by reducing nitro-tolyl-oxamic acid, and by heating tolylene-diamine dicyanide with HClAq at 150° (Bladin, B 18, 670). Needles — NaHA — AgA — HA — HOAc . Plates.

DI-OXY-DITOLYL

[8 4 1] $\text{C}_9\text{H}_6\text{Me}(\text{OH})_2\text{C}_8\text{H}_6\text{Me}(\text{OH})$ [1 8 4] *o-Di-cresol* [157°] (H), [161°] (G). Made from di-amido-ditolyl by the diazo-reaction (Gerber, B 21, 749, Hobbs, B 21, 1067). Needles, sl sol hot water, v sol alcohol and ether. Yields a di-nitro derivative [273°]

Di-acetyl derivative $\text{C}_{12}\text{H}_{12}(\text{OAc})_2$ [131°]

Di-benzoyl derivative [185°] Needles

Di-ethyl ether $\text{C}_{11}\text{H}_{10}(\text{OEt})_2$ [156°] Made, together with ditolyl, by the action of alcohol on di-amido-ditolyl (Schultz, B 17, 468). White plates, sol hot alcohol.

Di-propyl ether [115°] Leaflets.

Di-isooamyl ether [69°]

Tetra-oxy-ditolyl $\text{C}_8\text{H}_4\text{O}_4$ *ie*

[1 2 5 6] $\text{C}_8\text{H}_4\text{Me}(\text{OH})_2\text{C}_8\text{H}_4\text{Me}(\text{OH})_2$ [6 1 2 5] [202° uncor] Formed by oxidising hydrotoluquinone, dissolved in HOAc with MnO_2 and H_2SO_4 in the cold (Brunner, M 10, 174). FeCl_3 oxidises it to the quinhydrone $\text{C}_{10}\text{H}_8\text{O}_2$, crystallising in violet scales [220° uncor], and finally to the quinone $\text{C}_6\text{H}_4\text{O}_2$ [163°], crystallising in yellow prisms.

Tetra acetyl derivative

$\text{C}_{12}\text{H}_{10}(\text{OAc})_4$ [135°] Made by heating with NaOAc and Ac_2O at 160°. Needles

Anhydride $\text{C}_{12}\text{H}_8\text{Me}_2(\text{OH})_2\text{O}$ [232°] Got by heating $\text{C}_{12}\text{H}_8\text{Me}_2(\text{OMe})_2\text{O}_2$ with HClAq at 180° (Nietzki, B 11, 1281). Colourless plates (containing aq) (from dilute alcohol).

Di-methyl ether $\text{C}_{12}\text{H}_{10}\text{Me}_2(\text{OMe})_2(\text{OH})_2$ [173°] Formed by partial methylation. Colourless needles, quickly oxidised in alcoholic solution by air. Yields a di acetyl derivative $\text{C}_{12}\text{H}_8(\text{Me}(\text{OMe})(\text{OAc}))_2$ [6 1 2 5], crystallising in needles [123°] (Noelting & Werner, B 23, 3248).

Dehydride of the di-methyl ether $\text{C}_{11}\text{H}_9\text{O}_2$ *ie* $\text{C}_{12}\text{H}_8\text{Me}_2(\text{OMe})_2\text{O}_2$ [153°] Formed by oxidation of the di-methyl ether of hydrotoluquinone by chromic acid mixture (Nietzki, A 215, 161). Dark-red needles (from alcohol).

Tetra-methyl ether $\text{C}_{12}\text{H}_{10}(\text{OMe})_4$ [129°]

Di-ethyl ether $\text{C}_{12}\text{H}_{10}(\text{OEt})_2(\text{OH})_2$ [138°] Formed by the action of alcoholic ammonium sulphide on the dehydride (or quinone) $\text{C}_{11}\text{H}_9(\text{OEt})_2\text{O}_2$, which crystallises in green-black needles [139°], and is made by oxidising $\text{C}_8\text{H}_6\text{Me}(\text{OEt})_2$ with chromic acid mixture (Noelting, B 23, 3247, B [3] 4, 806). White needles.

α-OXY m-TOLYL-ACETIC ACID $\text{C}_8\text{H}_7\text{O}_3$ *ie*

[1 3] $\text{C}_8\text{H}_6\text{Me} \begin{smallmatrix} \text{CH}(\text{OH}) \text{CO}_2\text{H} \\ \text{m-Methyl mandelic acid} \end{smallmatrix}$ [84°] Formed from *m*-toluic aldehyde, KCr , and HCl (Bornemann, B 17, 1469). Small plates (from benzene), v sol water.

α-OXY-p-tolyl-acetic acid

[1 4] $\text{C}_8\text{H}_6\text{Me} \begin{smallmatrix} \text{CH}(\text{OH}) \text{CO}_2\text{H} \end{smallmatrix}$ [146°] Made by reduction of *p*-tolyl-glyoxylic acid by zinc dust and ammonia (Claus & Krosenberg, B 20, 2050). Tables (from water) — NaA — KA — aq — CaA — BaA — aq

Ethyl ether EtA [77°] Needles.

α-OXYTOLYL-ALLYL-THIO-UREA

[1 2] $\text{C}_8\text{H}_6(\text{CH}_2\text{OH})\text{NH CS NH C}_6\text{H}_5$ Formed from *o*-amido benzyl alcohol and allyl

thiocarbamide in benzene Oil, changed by warming with HClAq to $\text{C}_6\text{H}_4\text{CH}_2\text{NC}_6\text{H}_5$. [91°] Converted by HgO in alcohol to oxy allyl-quinoxaline dihydride $\text{C}_6\text{H}_4\text{CH}_2\text{NC}_6\text{H}_5$, which crystallises in four sided prisms [78°], and yields $\text{B}^1\text{H}_2\text{PtCl}_6$ [171°] (Söderbaum a. Widmann, *B* 22, 1870, 2937)

OXY-TOLYL-BENZYLIDENE-TOLYLENE-DIAMINE $\text{C}_{12}\text{H}_{16}\text{N}_2\text{O}$ *te* $\text{C}_6\text{H}_4\text{NHCH}_2\text{NCH}_2\text{C}_6\text{H}_4\text{OH}$ [160°] Made from *o*-amido di *p* tolyl amine and salicylic aldehyde (O Fischer a Sieder, *B* 23, 3801) Yellow crystals, sl sol water, v sol benzene

OXY-*p*-TOLYL BENZYL-METHYL-PYRIMIDINE $\text{C}_8\text{H}_8\text{MeC} \begin{smallmatrix} \text{N CMe} \\ \text{N C(OH)} \end{smallmatrix} \text{C CH}_3\text{Ph}$ [240°] Made from *p* tolenyl amidine hydrochloride and benzyl aceto acetic ether (Pinner, *B* 23, 3826) Slender needles (from pyridine), insol water, v sl sol hot alcohol

OXY-TOLYL CARBINOL $\text{C}_6\text{H}_4\text{Me(OH)CH}_2\text{OH}$ [5 2 1] [105°] *S* 67 at 15° Made by reduction of (2,5,1) oxytoluol aldehyde with sodium amalgam (Schotten, *B* 11, 784) Colourless leaflets

TRI-OXY TRI TOLYL CARBINOL *Anhydride* $\text{C}_{18}\text{H}_{18}\text{O}_3$, *te* $(\text{C}_6\text{H}_4\text{Me(OH)})_3\text{C} \begin{smallmatrix} \text{O} \\ \text{O} \end{smallmatrix} \text{C} \begin{smallmatrix} \text{O} \\ \text{O} \end{smallmatrix} \text{C}_6\text{H}_4\text{Me}$

Cresaurin Formed by heating *p* cresol with formic acid and ZnCl_2 at 110° (Nencki, *J pr* [2] 25, 275) Red amorphous powder, insol water, v sol HOAc Forms a crimson solution in alkalis
DI-OXY-DITOLYL DICARBOXYLIC ACID $\text{C}_{16}\text{H}_{14}\text{O}_6$ Made by heating sodium di oxy ditolyl with CO_2 in a closed vessel at 160° (Deninger, *B* 21, 1639) Crystals, insol water, sl sol alcohol and ether Not melted at 290° FeCl_3 gives a blue colour Yields a di acetyl derivative $\text{C}_{16}\text{H}_{12}\text{O}_6$ [163°]

OXY-TOLYL-CROTONIC ACID *Anhydride* *v* **DI METHYL COUMARIN**

Di oxy-tolyl crotonic acid *Anhydride* $\text{C}_6\text{H}_4\text{Me(OH)} \begin{smallmatrix} \text{CMe CH} \\ \text{O} \text{---} \text{CO} \end{smallmatrix}$ *Di methyl umbelliferone* [250°] Made by the action of H_2SO_4 on a mixture of acetoacetic ether and orcin (Pechmann a. Cohen, *B* 17, 2188) Needles (from alcohol), nearly insol water On boiling with NaHSO_4 it forms a solution coloured red by FeCl_3
Acetyl derivative $\text{C}_{11}\text{H}_{14}\text{AcO}_2$ [195°] Needles, v sol alcohol

TRI OXY-TRI-TOLYL ETHANE $\text{C}_6\text{H}_4\text{Me(OH)CH}_2\text{CH(C}_6\text{H}_4\text{Me OH)}$ Three isomerides with this formula are got by warming di-chloro-ethyl ether with *o*, *m*, and *p*-cresol respectively (Brückner, *A* 257, 322) They are amorphous powders, v sol alcohol and ether, insol water, and yield tri acetyl derivatives which are oxidised by FeCl_3 to anhydrides $(\text{C}_6\text{H}_4\text{Me(OH)CH}_2\text{C(OHMe(OH))})_3\text{O}$

OXY-TOLYL-HYDRAZINE *Methyl derivative* [1 4 3] $\text{C}_6\text{H}_4\text{Me(OMe)NH NH}_2$ [45°] Got from the methyl ether of amido *p* cresol by treatment with nitrous acid and subsequent reduction (Lumpach, *B* 22, 351) Crystalline

Oxy-*om*-di-tolyl-hydrazine *Ethyl derivative* [2 1] $\text{C}_6\text{H}_4\text{Me NH NH C}_6\text{H}_4\text{Me(OEt)} [1 3 4]$ [78°] Made by reducing the azo-compound

with alcoholic ammonium sulphide (Noelting a. Werner, *B* 23, 3260, *BI* [2] 4, 796) White crystals, insol water, sol alcohol

The isomerides

[2 1] $\text{C}_6\text{H}_4\text{Me NH NH C}_6\text{H}_4\text{Me(OEt)} [1 5 2]$ [138°], [4 1] $\text{C}_6\text{H}_4\text{Me NH NH C}_6\text{H}_4\text{Me(OEt)} [1 3 4]$ [77°], and [4 1] $\text{C}_6\text{H}_4\text{Me NH NH C}_6\text{H}_4\text{Me(OEt)} [1 5 2]$ [153°] are formed in like manner

DI OXY-DI TOLYL KETONE $\text{C}_{12}\text{H}_{14}\text{O}_2$, *te* $\text{C}_6\text{H}_4\text{Me(OH)CO C}_6\text{H}_4\text{Me(OH)}$ [138°] Got by potash-fusion from *o* cresol benzoin, which is a product of the action of benzotrichloride on *o* cresol (Schroeter, *A* 257, 74) Colourless needles, v sol alcohol and ether

Di oxy di tolyl ketone $\text{CO(C}_6\text{H}_4\text{Me OH)}_2$ [104°] Made from *p* cresol phthalen $\text{C}_6\text{H}_4\text{O}$, by potash fusion (Drewson, *A* 212, 344) Yellow needles (from alcohol), insol water, v sol alkalis

OXY *p* TOLYL METHYL ETHYL PYRIMIDINE $\text{C}_8\text{H}_8\text{MeC} \begin{smallmatrix} \text{N CMe} \\ \text{N C(OH)} \end{smallmatrix} \text{C Et}$ [218°] Made from *p* tolenyl amidine and ethyl acetoacetic ether (Pinner, *B* 23, 3826) Hair like needles, m sol hot alcohol

OXY-TOLYL METHYL KETONE $\text{C}_6\text{H}_4\text{O}_2$, *te* [1 2 5] $\text{C}_6\text{H}_4\text{Me(OH)CO CH}_3$ [104°] Formed from amido *m* tolyl methyl ketone by the diazo reaction (Klingel, *B* 18, 2699) Flat white prisms, v sol alcohol and hot water FeCl_3 gives a yellowish brown colour

Di oxy tolyl methyl ketone $\text{C}_6\text{H}_4\text{Me(OH)CO CH}_3$ [146°] Made from orcin, HOAc , and POCl_3 (Rasinsky, *J pr* [2] 26, 59) Needles, v sol alcohol, ether, and HOAc FeCl_3 colours its aqueous solution black

OXY *o* TOLYL METHYL PYRAZOLE

$\text{C}_{11}\text{H}_{11}\text{N}_2\text{O}$, *te* $\text{C}_6\text{H}_4\text{MeN} \begin{smallmatrix} \text{CO CH} \\ \text{N} \end{smallmatrix} \text{CMe}$ [183°] Formed by heating *o* tolyl hydrazine with acetoacetic ether at 140° (Knorr, *B* 17, 549) Colourless crystals With KOH and MeI it yields $\text{C}_{11}\text{H}_{11}\text{MeN}_2\text{O}$ [97°]

Oxy *p* tolyl methyl pyrazole $\text{C}_{11}\text{H}_{11}\text{N}_2\text{O}$ [140°] From *p* tolyl hydrazine and acetoacetic ether (K) Yields a methyl derivative [137°] The compound $\text{C}_6\text{H}_4\text{N} \begin{smallmatrix} \text{CO C NHC}_6\text{H}_4 \\ \text{N} \end{smallmatrix} \text{CMe}$ [217°]

is formed, together with $\text{C}_{12}\text{H}_{12}\text{N}_2\text{O}_2$ by heating *p* tolyl hydrazine with thio acetoacetic ether (Buchka a Sprague, *B* 22, 2555, *C J* 59, 340)

OXY TOLYL METHYL PYRIMIDINE

$\text{C}_8\text{H}_8\text{MeC} \begin{smallmatrix} \text{N CMe} \\ \text{N C(OH)} \end{smallmatrix} \text{CH}$ [216°] Made from tolenyl amidine and acetoacetic ether (Glock, *B* 21, 2658) Long white needles, m sol hot alcohol — Salts $\text{B}^1\text{H}_2\text{PtCl}_6 \cdot 2\text{aq}$ [241°] — $\text{B}^1\text{H}_2\text{CrO}_7 \cdot 7\text{aq}$ [c 170°] Yellow plates — $\text{B}^1\text{C}_6\text{H}_4\text{N}_2\text{O}$ [196°] Yellow needles

***o* OXY TOLYL-DIPHENYL-ETHYLIDENE-AMINE** $\text{C}_{12}\text{H}_9\text{N CPh CHPh OH}$ [141°] Made by heating benzoïn with *o* toluidine at 150° (Bandrowski, *M* 9, 693) Canary yellow needles

OXY-TOLYL-PROPIONIC ACID $\text{C}_{10}\text{H}_{11}\text{O}_3$, *te* $\text{CH}_2\text{C(C}_6\text{H}_4\text{OH)CO}_2\text{H}$ Got from di bromo oxy tolyl-propionic acid $\text{CHBr}_2\text{C(C}_6\text{H}_4\text{OH)CO}_2\text{H}$ by reducing with sodium amalgam (Böttlinger, *B* 14, 1598) Long four sided tables, v sol water

OXY-DI *p*-TOLYL PYRAZINE *Tetra-*

hydride $C_6H_4Me \cdot N < \begin{smallmatrix} CH_2 & CH_2 \\ | & | \\ CH_2 & CO \end{smallmatrix} > N \cdot C_6H_4Me$

'*Di-tolylmonooxypyrazine*' [168 5°] Made from *p*-toluidine by successive treatment with ethylene bromide and a mixture of chloro-acetic acid and NaOAc (Bischoff, *B* 22, 1785, 23, 2035) Crystals, v sol water and ether

Di-oxy-di-o-tolyl-pyrazine Dihydride
 $C_6H_4Me \cdot N < \begin{smallmatrix} CH_2 & CH_2 \\ | & | \\ CO & CO \end{smallmatrix} > N \cdot C_6H_4Me$ *Di-tolyl-di-*

acopyrazine [184°] Formed from di-o tolyl-ethylene-diamine and oxalic acid at 200° (Bischoff, *B* 22, 1805, 23, 2034)

Di oxy di p tolyl pyrazine dihydride [268°] Made by dissolving oxy di-*p* tolyl pyrazine tetrahydride in HOAc and oxidising with nitrous or chromic acid (B) Colourless plates, v sol. aniline and alcohol, sl sol ether and hot water. Alcoholic potash yields di tolyl ethylene-diamine [97°] and oxalic acid

Di oxy s di o tolyl pyrazine Dihydride
 $C_6H_4Me \cdot N < \begin{smallmatrix} CO & CH_2 \\ | & | \\ CH_2 & CO \end{smallmatrix} > N \cdot C_6H_4Me$ [160°]

Formed by heating *o* tolyl-amido acetic acid at 220° (Bischoff a Nastvogel, *B* 22, 1787) and by the action of potash on the product of the action of chloro acetyl chloride on *o* toluidine (Widman, *J pr* [2] 58, 293, 305) Rectangular plates (from alcohol), insol water, sol conc HClAq

Reactions —1 PCl_5 forms the compound $C_6H_4N < \begin{smallmatrix} CO & CCl \\ | & | \\ CCl & CO \end{smallmatrix} > N \cdot C_6H_4$, crystallising from alcohol in white needles [201°] —2 *Alcoholic potash* forms $C_6H_4N(CH_2CO_2H)COCH_2NHC_6H_4$, a white crystalline solid [129°] decomposed by conc HClAq at 160° into MeCl, *o* toluidine, and $C_6H_4NHCH_2CO_2H$

Salt —($C_6H_4N_2O_2$) $_2H_2PtCl_4 \cdot 4aq$ [176°]

Di oxy di p tolyl-pyrazine Dihydride [253°] Formed from oxalic acid and *p* tolyl amido-acetic *p* toluide (B & N) Made also by boiling the bromo acetyl derivative of *p* toluidine with alcoholic potash, or by heating *p* tolyl amido acetic acid at 200° (Abenius, *J pr* [2] 40, 433) Long white needles, insol water, m sol alcohol

Di oxy op di-tolyl pyrazine Dihydride

[1 2] $C_6H_4Me \cdot N < \begin{smallmatrix} C(OH) & OH \\ | & | \\ OH & C(OH) \end{smallmatrix} > N \cdot C_6H_4Me$ [1 4]

[180°] Made by heating *p*-toluidine with the chloro acetyl derivative of *o* tolyl amido acetic acid at 140° (Abenius, *J pr* [2] 40, 448) White matted needles, v sol hot alcohol, insol ether

Tetra-oxy di phenyl-pyrazine. Dihydride of the di-ethyl derivative

[1 4] $C_6H_4(OEt) \cdot N < \begin{smallmatrix} CH_2 & CO \\ | & | \\ CO & CH_2 \end{smallmatrix} > N \cdot C_6H_4(OEt)$ [4 1]

[265°] Slender needles

*OXY *p*-TOLYL QUINAZOLINE* $C_{13}H_{11}N_3O_2$
 $C_6H_4 \cdot N < \begin{smallmatrix} CH \\ | \\ CO \end{smallmatrix} N \cdot C_6H_4Me$ [146°] Made by oxidising *p*-tolyl quinazoline dihydride with $KMnO_4$ (Laal a Busch, *B* 22, 2698) Plates (from dilute alcohol) or needles (from ether), sl sol. hot water Does not react with hydroxylamine or phenyl-hydrazine — $BHCl$ [214°] Needles — $B' \cdot H_2PtCl_6$ [above 800°] Golden plates, sol alcoholic HCl

*OXY-(*Py* 8) TOLYL-QUINOLINE* $C_{13}H_{11}NO$
 $C_6H_4 \cdot N < \begin{smallmatrix} CH & CH \\ | & | \\ N & O \end{smallmatrix} C_6H_4Me(OH) \cdot$ ψ - *Flavemol*.

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[196°] Got by the diazo reaction from (4,3,1)-amido tolyl-quinoline (Weidel a Bamberger, *M* 9, 104) Small plates (from alcohol) Yields (*Py* 8) tolyl-quinoline on distillation with zinc-dust. Reduction followed by potash-fusion yields oxy-isophthalic acid — $B'HOl$ 2aq — $B' \cdot H_2PtCl_6$ (dried at 105°) Yellow needles

Acetyl derivative $C_{13}H_{11}NO_2$ [106°]

Di-oxy-tolyl-quinoline $C_{13}H_{11}NO_2$ [89°]

Formed, as well as oxy-tolyl quinoline, by the above method of preparation Plates (from alcohol), sol. benzene

DI-OXY-DI-o TOLYL SULPHIDE $C_{11}H_9SO_2$
 $2e$ $8(C_6H_4Me(OH)(214))$. *Dioxytoluene*

[135°] Formed from di-amido di-tolyl sulphide by the diazo-reaction (Truhlar, *B* 20, 676) Amorphous, sl sol water, alcohol, and ether

Di oxy-di-tolyl sulphide [210°-215°]. Formed from *m* cresol by treatment with SO_2 in CS_2 (Tassinari, *G* 17, 92, *C C* 1888, 1354) Yields an acetyl derivative [44°], which forms, on oxidation, the corresponding sulphone [268°], which on saponification gives $SO_2(C_6H_4OH)_2$ [133°]

Di oxy-di-tolyl sulphide [118°] Formed from *p* cresol and SO_2 . Yields an acetyl derivative [84°] which is oxidised by $KMnO_4$ to a sulphone [209°], which on saponification gives $SO_2(C_6H_4OH)_2$ [209°] (Tassinari, *Rend Accad Linc* [4] 4, 47)

OXY-o TOLYL-THIO-UREA $C_8H_9N_3SO_2$
 $C_6H_4Me \cdot NH \cdot CS \cdot NH \cdot OH$ [92°] Formed from *o* tolyl thiocarbimide and hydroxylamine in chloroform solution (Tiemann, *B* 22, 1939, Voltmer, *B* 24, 381) Slender needles, almost insol chloroform and water, sol alcohol, ether, and KOHAq $FeCl_3$ colours its alcoholic solution violet Yields *o* tolyl-cyanamide on long standing

Benzyl ether $C_8H_9Me \cdot NH \cdot CS \cdot NH \cdot OCH_2Ph$ [125°] Made from benzyl hydroxylamine and *o* tolyl-thiocarbimide

OXY-o TOLYL-UREA $C_8H_9N_2O_2$
 $CH_2(OH)C_6H_4 \cdot NH \cdot CO \cdot NH_2$ [180°] Formed from amido benzyl alcohol, potassium cyanate, and HCl (Söderbaum a Widman, *B* 22, 1668) Tables or prisms, in sol boiling water, v sl sol alcohol Decomposes on fusion, yielding di-oxy-di-tolyl-urea $CO(NH \cdot C_6H_4 \cdot CH_2OH)_2$ [108°], which crystallises in needles

OXYTROPINE C_8H_9NO [242°] A base occurring in crude belladonnine (Merling, *B* 17, 384, Ladenburg, *B* 17, 153) Crystalline — $B' \cdot H_2PtCl_6$ 2aq red prisms

OXY TRUXILLIC ACID

$(C_6H_4(OH)CH \cdot CH \cdot CO_2H)_2$ [273°]. Formed by potash fusion from (α)-sulpho truxillic acid (Liebermann a Bergami, *B* 22, 783) Prisms, m sol cold water Probably a polymeride of *p* coumaric acid An isomeric acid, formed from the amido-acid by the diazo-reaction, does not melt at 360° It yields an acetyl derivative [244°] (Homans, *B* 24, 2591)

OXY-URAMIDO-BENZOIC ACID Methyl derivative $NH \cdot CO \cdot NH \cdot C_6H_4(OMe) \cdot CO_2H$
Amsuramic acid S 05 at 100° Formed from potassium cyanate and a solution of the hydrochloride of amido-amic acid (Menschutkin, *A* 158, 99) Needles (from water) — CaA , 7aq

OXY-UREA $NH_2 \cdot CO \cdot NH \cdot OH$ [180°] Formed from aqueous hydroxylamine nitrate and potas-

s E

sium cyanate in the cold (Dressler & Stein, *Z.* [2] 5, 202) Needles (from alcohol) v. s. sol Ag. Decomposed by heating, yielding urea Reduces AgNO_3 , FeCl_3 gives a blue-violet colour — $\text{KH}(\text{CH}_2\text{N}_3\text{O})_2$ crystalline pp, got by adding alcoholic potash to the alcoholic solution (Hodges, *A* 192, 214) — $(\text{Ph}_3\text{H}(\text{OAc})_2)(\text{CH}_2\text{N}_3\text{O}_2)_2$ crystals — $(\text{CuCH}_2\text{N}_3\text{O}_2)_2 \cdot \text{H}_2\text{OAc}$ green mass

Benzyl ether $\text{NH}_2\text{CO NH OCH}_2\text{Ph}$ [188°] Formed from (a) benzyl hydroxylamine hydrochloride and potassium cyanate (Behrend & Leuchs, *A* 257, 203) Needles

Oxy-biuret $\text{C}_2\text{H}_4\text{N}_4\text{O}_2$ [134°] Formed from conc hydroxylamine sulphate and potassium cyanate, the mixture being evaporated with alcohol Minute four sided prisms, sol water and alcohol Gives a white pp with AgNO_3 Reduces warm ammoniacal AgNO_3 , FeCl_3 gives no colour — $\text{KOC}_2\text{H}_4\text{N}_4\text{O}_2$ minute needles

(a) **OXY-UVITIC ACID** $\text{C}_8\text{H}_8\text{O}_4$ 18 $\text{C}_8\text{H}_7\text{Me}(\text{OH})(\text{CO}_2\text{H})_2$ [1235] *Cresol dicarboxylic acid* [285°] (B) [295°] (H & R) [278°] (J) Formed from (a)-amido-uvitic acid by the diazo reaction (Böttinger, *B* 9, 804, 13, 1934, *A* 189, 177) and by potash-fusion from sulpho-uvitic acid (Jacobsen, *A* 206, 187, Hall & Remsen, *Am* 2, 137) Needles (from alcohol), v sol alcohol, insol chloroform Decomposes on fusion Conc HClAq at 200° yields o cresol FeCl_3 gives a purple colour — $\text{Ca}(\text{HA}')_2 \cdot 2\text{aq}$ — $\text{CaA}'_2 \cdot 2\text{aq}$ — $\text{CaA}'_4 \cdot 4\text{aq}$ — $\text{Ca}_2(\text{C}_8\text{H}_7\text{O}_4)_2 \cdot \text{Ag}_2\text{A}''$

Methyl ether $\text{Me}_2\text{A}''$ [126°] Needles

Mono-ethyl ether $\text{EtHA}''\text{aq}$ Needles

(b) **Oxy-uvitic acid**

$\text{C}_8\text{H}_7\text{Me}(\text{OH})(\text{CO}_2\text{H})_2$ [1435] [225°–235°] (J) [220°] (B) 8 13 at 12°, 52 at 100° (J) Formed by the action of nitrous acid on (b)-amido uvitic acid (Böttinger) Formed also by potash-fusion from s mesitol $\text{C}_8\text{H}_7\text{Me}_2\text{OH}$ and from oxy mesitylenic acid (Jacobsen, *A* 195, 285) Needles (from water), v sol alcohol and ether FeCl_3 gives a red colour HClAq at 200° forms p cresol (J) — $\text{Ag}_2\text{A}''$ prisms

Methyl ether $\text{Me}_2\text{A}''$ [79°] Needles

Oxy-uvitic acid $\text{C}_8\text{H}_7\text{Me}(\text{OH})(\text{CO}_2\text{H})_2$

[1346] Got by saponification of its ether which is formed from sodium acetoacetic ether by the action, in presence of NaOEt , of chloroform, chloral, trichloro acetic ether, or CCl_4 (Oppenheim & Pfaff, *B* 7, 929, 8, 884, 9, 321, Conrad & Guthzeit, *A* 222, 249) Needles, sl sol cold water, v sol alcohol and ether FeCl_3 gives a reddish violet colour Softens at 290°, decomposing at the same time Yields m cresol on distillation with lime PCl_5 forms a mixture of chlorides, whence water forms oxyuvitic acid and $\text{C}_8\text{H}_7\text{O}_4$ crystallising in needles — $\text{K}_2\text{A}''\text{aq}$ — $\text{BaA}''_2 \cdot 1\frac{1}{2}\text{aq}$ — $\text{CaA}''_2 \cdot 1\frac{1}{2}\text{aq}$ — CuA''_2 — $\text{Ag}_2\text{A}''$

Methyl ether $\text{Me}_2\text{A}''$ [108°] Plates

a OXY-VALERIC ACID $\text{C}_5\text{H}_{10}\text{O}_4$ 18

$\text{CH}_3\text{CH}_2\text{CH}_2\text{CH}(\text{OH})\text{CO}_2\text{H}$ [31°] Formed by the action of HClAq on its nitrile, which is the cyanhydrin of n-butyric aldehyde (Menozzi, *G* 14, 46) Got also by boiling a-bromo-butyric ether with aqueous Na_2CO_3 (Jushin, *B* 17, 2504) Silky hygroscopic laminae; v sol water, alcohol and ether — $\text{BaA}''_2 \cdot 1\frac{1}{2}\text{aq}$ plates — CaA''_2 8 36 at 100° — $\text{ZnA}''_2 \cdot 2\text{aq}$ 8 1 01 at 100° — CdA''_2 — CuA''_2 — $\text{Ag}_2\text{A}''$ small scales, sl sol cold water.

Ethyl ether EtA'' (190°). Oil.

a-Oxy-isovaleric acid

$(\text{CH}_3)_2\text{CH CH}(\text{OH})\text{CO}_2\text{H}$ [38°].

Formation — 1 By heating bromo isovaleric acid with Ag_2O and water (Clark & Fittig, *A* 189, 199) or with KOHAg (Ley & Popoff, *A* 174, 61, Schmidt & Schutleben, *A* 193, 87) — 2 By digesting chloro-isovaleric acid with baryta, water (Schlebusch, *A* 141, 322) — 3 From its ether, which is a product of the action of zinc and isopropyl iodide on oxalic ether (Markownikoff, *Z* 1870, 517) — 4 From its nitrile, which is made by combination of HCy with isobutyric aldehyde (Lipp, *A* 205, 24)

Properties — Rectangular tables, v sol water, alcohol, and ether Not deliquescent Volatile with steam Dilute H_2SO_4 at 140° splits it up into isobutyric aldehyde and formic acid

Salts — NaA' — BaA'_2 — ZnA'_2 — $\text{CaA}'_2 \cdot \text{aq}$ — $\text{CaA}'_2 \cdot 1\frac{1}{2}\text{aq}$ — $\text{CaA}'_2 \cdot 4\text{aq}$ — $\text{MgA}'_2 \cdot 2\text{aq}$ — $\text{CuA}'_2 \cdot \text{aq}$ — AgA' feathery crystals (from hot water)

Ethyl ether EtA' (175°) Oil

Anhydride $\text{C}_5\text{H}_8\text{O}_4$ **Valerolactide** [126°] (220°–240°) Made by heating the acid in sealed tubes at 200° Needles, v sol alcohol and ether Not attacked by dilute alkaline solutions

Amide $\text{Pr CH}(\text{OH})\text{CONH}_2$ [104°] Got from the nitrile and HClAq Crystals

Nitrile $\text{Pr CH}(\text{OH})\text{CN}$ SG § 96 Oil, decomposed at 186° into isobutyric aldehyde and HCy

a-Oxy-valeric acid $\text{CMeEt}(\text{OH})\text{CO}_2\text{H}$ [68°] (M) [66°] (B)

Formation — 1 By saponification of its ether, which is made by the action of zinc on a mixture of MeI , EtI and oxalic ether (Frankland & Duppa, *A* 135, 37) — 2 By boiling $\text{OEtMeBr CO}_2\text{H}$ with baryta water (Böcking, *A* 204, 14) — 3 From methyl ethyl ketone by combination with HCy and saponification of the resulting nitrile (B) — 4 By oxidation of $\text{CMeEtH CO}_2\text{H}$ with dilute KMnO_4 (Miller, *A* 200, 282)

Properties — Needles (by sublimation), v e sol water, alcohol, and ether Oxidised to methyl ethyl ketone by chromic acid mixture Reduced by HI to $\text{CMeEtH CO}_2\text{H}$ Rotates on water

Salts — ZnA'_2 — AgA' Nodules

Ethyl ether EtA' (165.5°) SG § 977

V D 498 (calc 5.04) Liquid, sol water

B-Oxy-isovaleric acid

$(\text{CH}_3)_2\text{C}(\text{OH})\text{CH}_2\text{CO}_2\text{H}$

Formation — 1 By oxidation of the alcohol $\text{CMe}_2(\text{C}_2\text{H}_5)\text{OH}$ with cold chromic acid mixture (Saytzeff, *A* 185, 168, 197, 78), or with KMnO_4 at 0° (Schirokoff, *J pr* [2] 23, 206) — 2 By the action of zinc on a mixture of acetone and chloro acetic ether (Reformatsky, *B* 20, 1210) — 3 By oxidation of tri oxy hexane by KMnO_4 (Reformatsky, *J pr* [2] 40, 404)

Properties — Syrup, v sol water, alcohol, and ether Not volatile with steam Reduced by HI to isovaleric acid

Salts — $\text{CaA}'_2 \cdot 12\text{aq}$ — CuA'_2 — $\text{CuA}'_2 \cdot 2\text{aq}$ — AgA' monoclinic crystals, sl sol water.

Ethyl ether EtA' (180°)

B-Oxy-valeric acid

$\text{CH}_3\text{CH}(\text{OH})\text{CHMe CO}_2\text{H}$ Formed by reduction of methyl-acetoacetic ether with sodium amalgam (Rohrbeck, *A* 193, 229) Syrup, resolved by distillation into water and methyl-crotonic acid [62°] — NaA' (dried at 100°). [210°] (Miller, *A*

900, 269) Crystalline powder—BaA', aq—AgA laminae, sl sol hot water

(γ)-Oxy-valeric acid

$\text{CH}_3\text{CH}(\text{OH})\text{CH}_2\text{CH}_2\text{CO}_2\text{H}$ The salts of this acid are made by dissolving the lactone in bases. The free acid quickly changes back to the lactone

Salts—The Ba and Ca salts are deliquescent amorphous masses—AgA' trichioneneedles

Ethyl ether EtA' Oil

Lactone $\text{CH}_3\text{CH}(\text{OH})\text{CH}_2\text{CH}_2\text{CO}$ (208° i v)

SG 2 1072 Occurs in crude wood vinegar (Grodski, F 17, 1369) Prepared by boiling γ-bromo valeric acid with water (Messerschmidt, A 208, 96), and by reduction of acetyl propionic (levulic) acid with sodium amalgam (Wolff, A 208, 104) Formed also by heating γ-oxy propylmalonic acid (Hjelt, A 216, 56), and by the action of NaOHAq on nitroso oxy methyl pyrrole dihydride (Tafel, B 22, 1884) Liquid, miscible with water Neutral to litmus Separated from solution by K_2CO_3 Reduced by HI and P at 250° to n-valeric acid Oxidised by HNO_3 to succinic acid Boiling alcoholic NaOEt forms $\text{C}_6\text{H}_5\text{O}$, [c 32°] (Fittig, A 256, 126) Phenyl hydrazine forms $\text{C}_{11}\text{H}_{11}\text{N}_2\text{O}$, [76°-79°], crystallising in needles, v sol water (W Wislizenus, B, 20, 402)

Amide $\text{CH}_3\text{CH}(\text{OH})\text{CH}_2\text{CH}_2\text{CONH}_2$, [56°] Formed by heating the lactone or the ether with NH_3 Aq Thin plates, v e sol water and alcohol, sl sol ether At 170° it is split up into NH_3 and the lactone (Neugebauer, A 227, 97)

Di-oxy-valeric acid

$\text{CH}_3\text{CH}(\text{OH})\text{CMe}(\text{OH})\text{CO}_2\text{H}$ Di-methyl glyceric acid [107°] Formed by the action of water at 99° on di methyl glycidic acid

$\text{O} < \text{CMeCO}_2\text{H}$ [62°], which is formed from

tiglic acid $\text{CH}_3\text{CH}(\text{OH})\text{CMeCO}_2\text{H}$ by successive treatment with HCl and boiling KOHAq (Meli koff, A 234, 228, B [2] 47, 166)—KA'—AgA'

Tetra-oxy-valeric acid $\text{C}_6\text{H}_5\text{O}_4$

$\text{CH}_2(\text{OH})\text{CH}(\text{OH})\text{CH}(\text{OH})\text{CH}(\text{OH})\text{CO}_2\text{H}$ Arabinic acid $[\alpha]_D = -67^\circ$ Formed by allowing arabinose (10 g), water (75 g), and Br (20 g) to stand for 36 hours (Bauer, J pr [2] 30, 379, 34, 46, Kihani, B 19, 3081, 20, 344) Hygroscopic crystalline mass—CaA', 5aq—SrA', 5aq prisms

References—Bromo and Chloro-oxy-valeric acid

TETRA-OXY-VALERIC ALDEHYDE

$\text{CH}_2(\text{OH})\text{CH}(\text{OH})\text{CH}(\text{OH})\text{CH}(\text{OH})\text{CHO}$ Arabanose [160°] Mol w 150, by Raoult's method (Brown a Morris, C J 53, 619) H C p 557, 100 H.F 259, 400 (Berthelot, C R 111, 12) A product of the hydrolysis of ARABIC ACID (q v) (Scheibler, B 1, 58, 108, 6, 612, 17, 1731, Kihani, B 13, 2804, 15, 87, 19, 3031, 20, 344, Claesson, B 14, 1271, O'Sullivan, C J 45, 41) Trimetric prisms, v sol hot water, nearly insol alcohol and ether Dextrorotatory (v vol 1 p 297) Tastes sweet Does not undergo alcoholic fermentation Yields a phenylhydrazide [158°] Arabum $\text{C}_6\text{H}_{11}\text{O}_5$ is an anhydride of this aldehyde (O'Sullivan, C J 57, 59).

OXY-VALERO-CYAMINE v α-GUANIDO-VALERIC ACID

OXY-VINYL-BENZOIC ACID Anhydride v. METHYLENE-PHTHALIDE

OXY-XANTHONE v OXY-DIPHENYLENE KETONE OXIDE

OXY-XYLENE v. XYLENOL and TOLYL-CARBINOL

Di-oxy-xylene The (6,3,2,1)-, (5,2,3,1)-, and (5,2,4,1) di oxy-xylenes are described as HYDROXYLQUINONES OXY-TOLYL-CARBINOL is an α-endo-di-oxy xylene (5,3,4,1)-Di-oxy-xylene is described as BEXOROLIN

Di-oxy-m-xylene $\text{C}_8\text{H}_6\text{Me}_2(\text{OH})_2$, [184-6], Xylorcin Mol w 188 [125°] (278°) Formed from amido m xynolol [161°] by the diazo-reaction (Kostanecki, B 19, 2324) White monoclinic plates (from chloroform), v sol water, alcohol, and ether Not affected by air containing NH_3

Di-acetyl derivative $\text{C}_8\text{H}_4(\text{OAc})_2$, [45°] (286°) Prisms, insol cold water

Di oxy-xylene $\text{C}_8\text{H}_6\text{Me}_2(\text{OH})_2$, [120°] Got by potash fusion from chloro m xylene sulphonic acid (Gundelach, B [2] 28, 345) Gives a red colour with bleaching powder solution

Di-oxy-m-xylene $\text{C}_8\text{H}_6\text{Me}_2(\text{OH})_2$, [182-4] [146°] Formed from m-xylene by heating with H_2SO_4 at 150°, converting the resulting disulphonic acid into chloride and fusing the $\text{C}_8\text{H}_6\text{Me}_2(\text{SO}_2\text{Cl})_2$ with potash (Wischni, B 23, 3113) White needles (by sublimation), v sol water, alcohol, and ether FeCl_3 colours its solution deep violet Fusion with phthalic anhydride forms a fluoresein

Di-α-oxy-o-xylene $\text{C}_8\text{H}_4(\text{CH}_2\text{OH})_2$, Phthalalcohol o Tolylene alcohol Xylylene alcohol. Di methyl benzene glycol [64°] 8 (ether) 25 at 18° Formed by the action of sodium amalgam upon a boiling solution of phthalyl chloride in HOAc (Hessert, B 12, 646) Formed also by boiling di α-bromo-o-xylene with Na_2CO_3 Aq (Baeyer a Perkin, jun, B 17, 124, C J 53, 6; Colson, C R 98, 1643, B [2] 43, 6, 45, 6; A Ch [6] 6, 106) Tables (from ether), v e sol water and alcohol HBr forms $\text{C}_8\text{H}_4(\text{CH}_2\text{Br})_2$ and HCl acts in like manner KMnO_4 oxidises it to phthalic acid Resinified by cold H_2SO_4 Hot H_2SO_4 forms amorphous insoluble $(\text{C}_8\text{H}_4\text{O})_n$ and syrupy $\text{C}_8\text{H}_4\text{O}_2$ (Hjelt, B 19, 1588) HNO_3 forms phthalide

Di acetyl derivative $\text{C}_8\text{H}_4\text{O}_4$, [37°] Di ethyl ether $\text{C}_8\text{H}_4(\text{OEt})_2$, (249°) at 720 mm Liquid (Leser, B 17, 1325)

Di-α-oxy m-xylene $\text{C}_8\text{H}_6(\text{CH}_2\text{OH})_2$, [47°] SG (liquid) 11 161, 22 1385 Formed by boiling $\text{C}_8\text{H}_4(\text{CH}_2\text{Br})_2$, [77°] (1 mol) with water containing K_2CO_3 (1 mol) (Colson, A Ch [6] 6, 112, C R 99, 40) Got in like manner from $\text{C}_8\text{H}_4(\text{CH}_2\text{Cl})_2$ (Colson a Gautier, B [2] 45, 6) Crystalline solid, with bitter taste, v e sol water and alcohol, m sol ether HBr regenerates di-α-bromo-m-xylene Gives isophthalic acid on oxidation

Ethyl ether $\text{C}_8\text{H}_6(\text{CH}_2\text{OEt})_2$, (248°) at 712 mm Got by boiling $\text{C}_8\text{H}_4(\text{CH}_2\text{Br})_2$ with alcoholic potash (Kipping, B 21, 46, C J 53, 46) Oil Yields isophthalic acid on oxidation

Di-α-oxy-p-xylene $\text{C}_8\text{H}_6(\text{CH}_2\text{OH})_2$, p-Xylylenol alcohol, [113°] Formed from $\text{C}_8\text{H}_4(\text{CH}_2\text{Cl})_2$ (1 pt.) by heating with water (80 pts) at 175° (Grimaux, A 155, 342, Colson a Gautier, B [2] 45, 7) It is one of the products of the action of boiling NaOHAq upon terephthalic aldehyde (Löw, A 231, 374) Needles, v. sol. water, al-

cohol, and ether. Yields terephthalic acid on oxidation.

Acetyl derivative $C_8H_7(OH, OAc)$, [47°] Made from $C_8H_7(OH, Cl)$, and $KOAc$ in alcohol.

Benzoyl derivative $C_8H_7(OH, OBz)$. Needles, v sol alcohol and ether (Grimaux).

Mono-ethyl ether $C_8H_7(OH, OEt)$ (251°). Oil. Formed from di- α -chloro-*p*-xylene and conc alcoholic potash at 100° (G). Successive treatment with $POCl_3$ and water converts it into terephthalic aldehyde (Colson, *C R* 99, 975).

Tri-oxy-xylene $C_8H_6(OH)_3$, [122°]. Formed by reducing oxy-isoxylquinone with aqueous SO_2 (Fittig a Siepermann, *A* 180, 87). Crystallises from water in tables (containing aq). Melts at 90° when hydrated. Colours the skin brown. On spontaneous evaporation of the aqueous solution in air it forms a quinoxaline as dark lustrous needles [143°]. Yields *m*-xylene on distillation with zinc dust.

Tri-acetyl derivative [99°]. Prisms. Reference—TETRA CHLORO DI OXY-XYLENE.

DI-OXY-XYLENE CARBOXYLIC ACID $C_8H_6(OH)_2CO_2H$ [1 3 4 6 5]. *Xylorcan carb oxylie acid* [196°]. Got by heating *m*-xylorcan with $NaHCO_3$ and some water at 180° (Kostanecki, *B* 19, 2328). Prisms from dilute alcohol, sl sol water. Gives off CO_2 on fusion. $FeCl_3$ gives a deep blue colour.

Oxy-*m*-xyloquinone $C_8H_6(OH)O_2$, [108°]. Formed by distilling di-amido mesitylene with chromic acid mixture and water, Me being displaced by OH (Fittig, *B* 8, 16, *A* 180, 27). Orange needles, smelling like quinone, m sol hot water, v sol alcohol and ether. Volatile with steam. Its alkaline solution is reddish violet. Reduced by SO_2 to tri-oxy-xylene. Acetyl chloride at 100° forms a crystalline body [124°], insol water— $C_8H_6O_2(OK)$. Small black needles, v s. sol. Aq, m sol alcohol, insol ether— $(C_8H_6O_2)_2Ba$. Brownish-red pp.

α -OXY-XYLYL-ACETIC ACID [1 3,4] $C_8H_6Me_2CH(OH)CO_2H$ [119°]. Got by reducing (1,3,4)-xylyl glyoxylic acid (Claus, *J pr* [2] 48, 148). Rhombohedra (by sublimation), v. sl. sol. cold water, v sol alcohol and ether.

α -Oxy-xylyl-acetic acid [1 4 2] $C_8H_6Me_2CH(OH)CO_2H$ [114°]. Got in like manner from [1 4 2] $C_8H_6Me_2COCO_2H$ (Claus). Needles or prisms, v sol hot water.

DI-OXY-XYLENE-DI-METHYL-DI-PYRIMIDINE $C_8H_4(CH_3)_2C \begin{smallmatrix} N(OH) \\ NMe \end{smallmatrix} > CH_3$, [above 250°]. Made from acetoacetic ether and *p*-phenylene-diact imido- ether (Glock, *B* 21, 2861). Crystalline mass, insol ordinary solvents.

OXY-XYLYL-METHYL-PYRAZOLE $C_8H_6Me_2N \begin{smallmatrix} C(OH)CH \\ N=CH \end{smallmatrix}$ [159°]. Got from the product $C_8H_6N_2O_4$ of the action of (1,3,4)-xylyl-hydrazine on acetoacetic ether by heating with conc HCl aq at 150° (Klauber, *M* 12, 215). Small white needles— B^*HCl [185°]— $B^*H_2FeCy_2$, white crystals.

Oxy-xylyl-di-methyl-pyrazole $C_8H_6Me_2N \begin{smallmatrix} CO-CH \\ NMe \end{smallmatrix}$ [118°]. Made by heating the compound $C_8H_6N_2O_4$ (*v supra*) with MeI and $MeOH$ at 180° (K). Small white needles, v sol alcohol and ether, sol cold water.

Reduces Fehling's solution. $FeCl_3$ gives a violet red colour— B^*HCl [35°]. Small crystals.

DI-OXY-DI-XYLYL-PYRAZINE DIHYDRIDE $C_8H_6Me_2N \begin{smallmatrix} CO \ CH_2 \\ CH_2 \ CO \end{smallmatrix} N C_8H_6Me_2$, [203°]. Formed by boiling bromo acetyl (1,1',2) xylidine with alcoholic potash (Abenius, *J pr* [2] 40, 486). Flat needles, insol water and ether.

OZOKERIT. A fossil resin, consisting chiefly of a hydrocarbon called lekene (*q v*). On chlorination in presence of $BrCl_3$ at 360° it yields CCl_4 , C_2Cl_6 , C_3Cl_8 , and C_4Cl_{10} (Hartmann, *B* 24, 1019) (*V* also PARAFFIN).

OZONE O_3 , Mol w 47.91. A blue gas (Hautefeuille a Chappuis, *C R* 91, 522), it usually occurs mixed with oxygen, and possesses a characteristic odour (−106°) (Olszewski, *M* 8, 69, *W* 87, 337). *VD* 24 *S* at 760 mm. 366 at 18° (Schöne, *B* 6, 1224), 834 at 1° (Carius, *A* 174, 80), and 2745 at 14° (McLeod, *C J* 49, 607). Andrews states that it is insoluble in water.

Occurrence.—Ozone is believed to be a normal constituent of pure air. Hartley (*C J* 39, 111), Chappuis (*C R* 91, 985, 94, 858), and E Schöne (*J R* 1884 2, 250), who have examined the absorption spectrum of ozone, have attributed the blueness of the sky to its presence. But the recent observations of Laveag a Dewar (*P M* [5] 26, 286) show that the absorption spectrum of compressed oxygen exhibits certain bands identical with those of the solar spectrum, which Angstrom found to be equally strong whether the atmosphere was wet or dry, and that daylight when observed through a column of oxygen 18 m in length and at 90 atmos possesses a blue tint (*v* also Olszewski, *W* 42, 663). The proportion of ozone in the air varies very considerably, and is supposed by many observers to be greater at high than at low altitudes (*v* also Thorpe, *C J Proc* 72). Houzeau judges the maximum proportion at ordinary levels to be $\frac{1}{10000}$ by volume (*C R* 74, 712). Andrews found that a temperature of 250° destroys the constituent of the air which exhibits the reactions of ozone, whereas air containing traces of chlorine, or of the higher oxides of nitrogen, is not so affected by heat (*Pr* 16, 68). On the other hand, Hovsøy (*Bl* [3] 2, 877) and Schöne (*B* 13, 1503) conclude that the presence of ozone in the air is still unproved.

Formation.—1. Ozone is formed in the electrolysis of dilute sulphuric acid (Schönbein, *P* 50, 616, Margnac, *C R* 20, 808, Meidinger, *A* 88, 57, *C J* 7, 251, Baumeit, *P* 89, 38, Andrews, *T* 146, 1, Soret, *Arch des Sciences*, 16, 218, *C R* 56 390, Berthelot, *C R* 86, 71, *A Ch* [5] 14, 345, Schöne, *B* 6, 1224, Carius, *A* 174, 1, Brodie, *C J* 17, 293, McLeod, *C J* 49, 591). Berthelot and Schönbein also obtained ozone by the electrolysis of other acid solutions. It has been supposed that the ozone formed in electrolysis is accompanied by H_2O_2 , but Brodie (*C J* 17, 281), and, more recently, McLeod (*loc cit*), have shown that the oxidising body which remains in solution is probably S_2O_8 . The proportion of ozone present in electrolytic oxygen appears to depend to a great extent on current-density. By using a positive electrode of very small area McLeod obtained O containing 17.4 per cent of ozone, when electrodes of large

area are used the yield of ozone is sometimes very small.—3 When air or oxygen is exposed to the electric discharge, especially if it be the silent discharge, the O is partly converted into ozone. If air be used, oxides of N may be formed and mistaken for ozone. According to Berthelot (*C R* 92, 82) and Hautefeuille a Chappuis (*C R* 92, 80, and 184), oxides of N may be formed to a slight extent even by the silent discharge, and H and O have obtained a new and unstable oxide of nitrogen in this way. Giannetti a Volta (*G* 5, 439) found that with the discharge from a Holtz machine the yield of ozone is increased by using a wire brush as negative electrode. Bichat a Guntz (*C R* 107, 534, *A Ch.* [6] 19, 131), who used an ozone generator consisting of a wire stretched along the axis of a metallic tube, found that the negative effluve produced by far the greatest yield of ozone. Thus they attribute to its higher temperature. It has been suggested that the production of ozone by the electric discharge is an effect of a condition of electro static stress. But Thomson a Threlfall (*Pr* 40, 329) find that oxygen is only converted into ozone when there is an actual luminous discharge. This has been confirmed by Bichat a Guntz, and some experiments by the writer seem to show that even when oxygen is illuminated by the ultra violet rays ozone is only formed by actual luminous discharge. Dewar has obtained a body giving the reactions of ozone from air by passing a current of water through a glass tube, surrounded by a larger tube of platinum which was heated by the oxyhydrogen flame, the air from the annular space between the hot and cold tubes being sucked into the inner tube by the stream of water through a minute hole in the glass tube, and collected and examined. If the substance thus obtained was really ozone, this result seems to confirm the idea that the action of the electric discharge on oxygen is due to temperature (v. also Elster a Geitel, *W* 39, 321, and Hoesvay, *Bl* [3] 2, 734).

The formation of ozone from O by electric discharge is greatest at low temperatures and under high pressure (von Babo, *A* 1863 *Suppl.* 11, Hautefeuille a Chappuis, *C R* 91, 228). But the exact influence of temperature and pressure have probably not yet been made out. Hautefeuille a Chappuis have noticed that at a pressure of about 50 mm. ozone is alternately produced and destroyed by the silent discharge (*C R* 94, 646). Von Babo a Claus, and Hautefeuille a Chappuis, consider that prolonged action of the discharge is favourable to ozonification. But Brodie, with the apparatus described below, found that the maximum effect was quickly reached. The writer's experience agrees with that of Brodie, and tends to show that observations to the contrary effect have been due to the irregular working of the machine employed.

Bichat a Guntz, using the apparatus described above, have failed to find any simple quantitative relation between the potential difference of the discharging surfaces and the yield of ozone, though they, and also Giannetti a Volta (*G* 5, 439) and Berthelot, find that an increase of potential produces an increased yield of ozone. (For details on various points v. Marignac a. De la Rive, *Arch. of Elect.* 5, 5; Fremy a Bec-

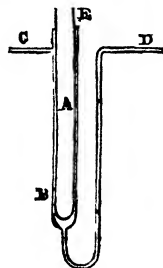
querel, *A Ch.* [8] 85, 62, Andrews; *T.* 146, 1; 150, 113, Brodie, *T* 162, 435, Berthelot, *C R* 88, 50, *A Ch.* [5] 12, 448, Hautefeuille a. Chappuis, *C R* 92, 80, 134, 94, 646, Shenstone a. Cundall, *C J* 51, 610.) Hautefeuille a. Chappuis (*C R* 91, 762) find that the production of ozone by the action of the electric effluve on oxygen is prevented by the presence of Cl, but that N, H, and SiF₄ are favourable to its production.

3 Brodie (*T* 164, 88), by the action of the silent discharge on carbon dioxide, under the most favourable conditions, obtained as much as 85 p.c. of the liberated oxygen in the form of ozone.

4 Ozone has long been regarded as one of the products of various cases of oxidation such as the slow oxidation of P, Et₃O, and turpentine, the decay of organic matter, and the combustion of compounds containing H in the air, Thorpe a Tutton observe that it is not formed in the oxidation of P₂O₃ (*C J* 57, 569). It is possible, however, that ozone is less frequently formed in such changes than has been supposed, and that the reactions attributed to ozone may often have been due to H₂O₂. The experiments of Kingzett (*C J* 37, 792) and McLeod (*C J* 37, 118), however, seem to make it certain that ozone is produced in the slow oxidation of P (v. also Hoesvay, *Bl* [8] 2, 860, 4, 707, Leeds, *C. N* 39, 157, 40, 70, 41, 164, 42, 17, Cundall, *C J* *Proc* 78, 26, Loew, *B* 22, 8325).

Heat of formation— $3O_2 = 2O_3 = -59,200$ (Berthelot, *C R* 82, 1281), —66,720 (Mulder a van der Meulen, *B* 15, 511).

Preparation.—1 From oxygen. The following method, which was introduced by Siemens and Brodie, is perhaps the most convenient. A tube, A, is sealed into a slightly larger tube B,

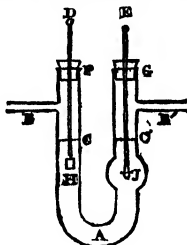


at E, before the blowpipe, or by means of solid paraffin. A and B should be of thin glass, and two narrow tubes, O and D, should be attached to B. A is filled with dilute H₂SO₄, and the apparatus is then immersed in dilute acid to the level OD, the electrodes of a Ruhmkorff coil, or of an electric machine, are respectively connected with the acid in A, and the contents of the cylinder, and a slow current of O is led through the apparatus from D to C while the discharge passes. The liquid in the cylinder should be cooled by ice, or it may be replaced by a freezing mixture, in which case a platinum wire should be wrapped round the outside of B. The gas which escapes at C is well charged with ozone. The ozonised oxygen may be collected over oil of vitriol. It must not be brought into contact with india-rubber; joints that will bear contact with ozone may be made by slipping a wider tube over the

ends of those that are to be connected, warming them, and running a little melted paraffin between the inner and outer tubes

The character of the discharge in the ozone generator has been studied by Hautefeuille & Chappuis (*C. R.* 91, 281), Thomson & Threlfall (*Pr.* 40, 829), Shenstone & Cundall (*C. J.* 51, 622), and by Bichât & Guntz (*A. Ch.* [6] 19, 181)

2 By electrolysis When dilute H_2SO_4 is electrolysed with electrodes of considerable area the proportion of ozone in the O is usually small McLeod (*C. J.* 49, 591) recommends the following method Place dilute H_2SO_4 (*S. G.* 11) in a U tube, A, attach delivery tubes to the



arms B B' Let the negative electrode H be a sheet of Pt foil suspended by the glass D from a cork closing the mouth at F, and let the positive electrode J consist of two fine wires, sealed into a glass tube E, which is afterwards filled with mercury On connecting D and E with a galvanic battery, O highly charged with ozone will escape at B' It is best to immerse the U tube in ice cold water McLeod has obtained O containing as much as 17.3 p.c. of ozone by means of this apparatus

3 To prepare ozone by the oxidation of phosphorus Place clean sticks of P half submerged in water in a series of flasks, and conduct a rather slow stream of air through the flasks, and then through a little water A temperature of 24° – 25° gives the best results H_2O_2 is also formed, but it remains dissolved in the water (Kingzett, *C. J.* 37, 792)

Properties and Reactions—Ozone is more strongly magnetic than common oxygen (Bequerel, *C. R.* 92, 348) Its rate of diffusion is near to that required for the density 24 (Sorêt, *A. Ch.* [4] 13, 257) It is entirely destroyed at 270° (Andrews, *T.* 150, 113) by contact with platinum black at ordinary temperatures (Mulder & van der Meulen, *B.* 1, 167), and on contact with pure Hg (Hg is not visibly oxidised if both be perfectly dry, Shenstone & Cundall, *C. J.* 51, 619) Ozone is also decomposed by dry Ag, the silver being but slightly oxidised, and by contact with MnO_2 (Andrews) Volta (*G.* 9, 521) states that Au, Pt, Pd, and dry Ag are without action on ozone Brodie came to a similar conclusion in regard to Au, Al, and Cu Berthelot (*C. R.* 86, 76) and others have considered that dry ozone has no fixed pressure of dissociation If suddenly compressed, or compressed without cooling, it explodes with a yellowish flame (Hautefeuille & Chappuis, *C. R.* 91, 522). The action of ozone on salts has been studied by Maquenne (*C. R.* 94, 795), and by Mailfert (*C. R.* 94, 860). The latter has also

studied its action on S, Se, Te, and several sulphides, and on CH_4 , C_2H_4 , C_2H_6 , C_3H_8 , C_4H_{10} , C_2H_5OH , and C_2H_5I (*C. R.* 94, 1186) Ozone oxidises alcohols of small molecular weight more readily than the polyhydric alcohols (A Renard, *A. Ch.* [5] 16, 289) Liquid ozone explodes violently on corking into contact with C_2H_4 (Olszewski, *A. Ch.* [2] 37, 887) It acts as a strong bleaching agent on vegetable colours, and quickly destroys india rubber, its action on cork is much less rapid It does not oxidise CO at 300° (Remsen, *Am.* 4, 50) It oxidises NH_4Ag , forming ammonium salts of nitrous and nitric acid (Carius, *A.* 174, 31), unless the solution is dilute (Hartley, *C. J.* 39, 123) It is believed to be completely absorbed by turpentine and oil of cinnamon (Sorêt, *A. Ch.* [4] 7, 113), peroxidised compounds being formed which react with water (Kingzett, *C. J.* 37, 800) When ozonised O acts on $KIAg$, I is set free and the ozone is destroyed, but the volume of the gas remains unaltered With neutral solutions the initial action may correspond to the equation $2KIAg + O_3 = K_2OAq + O_2 + I_2$, but usually KIO , is a final product of the change P glows freely in ozonised air (Thorpe & Tutton, *C. J.* 57, 571) Ozone oxidises $TiOH$ and As_2O_3 , the latter action has been used in determining its heat of formation It is destroyed by solutions of KHO , BaO , and CaO to a certain extent (Andrews) Hartley has observed the formation of peroxide of potassium by its action on solid KOH (*C. J.* 39, 124) But it is unaffected by solution of Na_2CO_3 (Brodie) It readily attacks I, forming periodic acid and lower oxides of I (Ozier, *C. R.* 86, 722) It converts ether into an ozonised substance which acts with water to form H_2O_2 (Kingzett, *C. N.* 34, 127, Berthelot, *C. R.* 86, 71, Dunstan & Dymond, *C. J.* 57, 584) Several observers have concluded that it arrests putrefaction of animal matter, and have proposed its use as a preservative for meat When present in the air in large quantities it frequently produces irritation of the mucous membrane

Although its general action is that of a strong oxidiser, in certain cases ozone acts as a reducing agent Thus when ozone acts with Na_2O_2 , an expansion occurs which is due to the simultaneous decomposition of the two bodies in equivalent proportions (Brodie, *T.* 162, 454), probably according to the following equation— $Na_2O_2 + O_3 = Na_2O + 2O_2$ Probably other unstable oxides, such as H_2O_2 , react with it similarly under favourable conditions Its action on blood has been said to resemble that of a reducing agent (Dogiel, *C. C.* 1875, Binz, *C. C.* 1882) It changes the red colour of the compound formed when sulphamic acid is mixed with naphthyl amine to orange yellow (Ilosvay, *Bl.* [1] 2, 351)

Detection—Paper impregnated with $TiOH$ is turned brown by ozone even when dry (Schöne, *B.* 13, 1508) The similar change of colour produced by nitrous acid is destroyed by excess The colour is also permanent when ozone is mixed with a relatively small proportion of nitrous acid (Ilosvay) The action of ozone on the red compound of naphthylamine and sulphamic acid (*see Properties*) also affords a test for ozone, even in the presence of traces of nitrous acid (Ilosvay, *Bl.* [3] 2, 860) Ozone may be distinguished from H_2O_2 by not yielding water when strongly heated, and by the absence

of any action with titanio acid or chromic acid, also by resisting the action of Na_2CO_3 , which destroys H_2O_2 . Papers saturated with solutions of KI and starch are often used for the detection of ozone, but are only trustworthy when the absence of Cl and of oxides of nitrogen can be assured. Houshau employed test papers steeped in faintly acid solution of litmus and then treated with KI. These he found to be insensitive to O_3 , to traces of oxides of nitrogen, and to H_2O_2 . When exposed to ozone, however, they are turned blue by the alkali that is set free.

Estimation.—1 Rough estimations of ozone are frequently made by comparing the tints produced by exposing some of the test papers mentioned above with a scale of tints. Such methods are not very satisfactory.—2 Thénard has estimated the proportion of ozone in the air volumetrically by means of a standard solution of As_2O_3 in HClAq of such strength that 1 c.c. is equivalent to 1 mgrm. of O . 10 or 20 c.c. of the solution are thoroughly agitated with the sample of gas, 30 c.c. of a 1 p.c. solution of H_2SO_4 are added, and it is then titrated in the usual manner with permanganate. For 8 grms. of O absorbed 24 grms. of O_3 are destroyed (Berthelot, *C R* 82, 1251). As oxides of nitrogen and H_2O_2 , if present, tend to reduce the amount of ozone found, this method gives the minimum, and not the maximum, amount of ozone present.—3 If a solution of potassium arsenite containing 73 grm. per litre, with an excess of pure KI, be agitated with air containing ozone, part of the arsenite is converted into arsenate by the ozone, and the amount of unaltered arsenite can be found by titrating with very dilute I solution after adding some ammonium carbonate and starch. This method is strongly recommended by Hartley (*C J* 39, 120). The solution of the arsenite should be acidified for keeping, and neutralised before use with KHCO_3 .—4 The action of ozone on KIAq may also be employed to estimate ozone. The iodide must be perfectly free from iodate, and must be thoroughly brought into contact with the gas. When the action is complete, and *not before*, the product must be acidified with dilute HClAq or $\text{H}_2\text{SO}_4\text{Aq}$. The liberated I may then be titrated in the usual manner. The O equivalent to the I liberated, multiplied by three, gives the amount of ozone.—5 The amount of ozone produced in the electrification of oxygen may also be calculated from the contraction that occurs under the influence of the discharge (*v. Babo, A Suppl.* 2, Andrews, *T* 150). Shennstone, a Cundall (*C J* 51, 610) have devised an apparatus for this purpose. S & O have shown that if the ozonised gas comes into contact with oil of vitriol in such a process, the acid should be previously thoroughly treated with ozone, otherwise the results are likely to be high.

History.—The earliest known record concerning ozone relates to an observation by van Marum, 1785, that air or O when submitted to the electric spark acquires a characteristic odour. About fifty years later, 1840, Schönbein, who published about sixty papers on this subject, published his first memoir on ozone in Poggen-dorff's *Annalen*. As the result of his numerous observations, Schönbein recognised ozone as a distinct form of matter, ascertained that it could be obtained by the electrolysis of dilute acid, by

means of the electric discharge on O , and in the slow oxidation of P. And it is interesting to note that these still constitute the chief methods for the production of ozone. He also observed many of its chief properties. For several years after Schönbein's original discovery, comparatively little progress was made in ascertaining the nature of ozone till the experiments of Marignac and De la Rive, 1845 (*Arch. of Elect.*), and of Fremy and Becquerel, 1853 (*A Ch.* [3] 35, 62), showed that the purest O then obtainable could be entirely converted into ozone, provided that the action of the spark took place in the presence of excess of KI, or of moist silver, which appeared to be capable of completely absorbing ozone. But it was not till a much later period that the researches of Andrews (*T* 146, 1) and Soret (*C R* 58, 390) finally showed that ozone from all sources is identical. Although the researches of Marignac & De la Rive, and of Fremy & Becquerel, thus established the character of ozone and its production from oxygen, the exact relation of the two bodies was still imperfectly understood, and ozone seems to have been regarded as differing from oxygen either in regard to its electrification, or, by some, in being a compound of oxygen and water, until in 1860 Andrews and Tait (*loc. cit.*) examined the volumetric relations of ozone and oxygen, and by a masterly research showed that the ozonising of oxygen is accompanied by a contraction in the volume of the gas, and that, on the other hand, the reproduction of O from the ozone by heat causes the gas to recover its original volume. In this research A and T also showed that the iodine titre of a sample of ozonised oxygen corresponds in every case to the contraction that has occurred on ozonising it, a fact which, in the hands of Soret and Brodie, materially contributed to the further elucidation of the subject. Andrews and Tait also observed that when ozonised oxygen acts with KIAq , the gas remains unaltered in volume after the action is complete. The true bearing of these facts was not, however, perceived till Odling (*Manual of Chemistry*) pointed out, in 1861, that they were consistent with the adoption of O_3 as the simplest formula for ozone. This interpretation was afterwards supported by the experiments of Soret and Brodie. The former showed that when ozonised oxygen is allowed to act on turpentine, a contraction occurs that is equal to about twice the iodine titre of the gas, *i.e.* is about twice as great as the contraction which accompanies its formation, from which, if it be assumed that the ozone formed is wholly absorbed by the turpentine, it follows that two volumes of gaseous ozone contain three volumes of gaseous oxygen. Soret's experimental numbers did not agree very closely with this hypothesis. But the hypothesis was confirmed by subsequent experiments, in which he compared the rates of diffusion of ozone, O_2 , and CO_2 , and found that the rate of diffusion of ozone approximates to that of a gas having the SG 24 (*A Ch.* [4] 18, 257). Finally, in 1872, Brodie (*T* 162, 435), as one of the results of a beautiful series of experiments, found that, while in some cases, as in that of neutral KIAq , the oxidation caused by ozone is unaccompanied by any contraction in the volume of the gas, in other cases various degrees of con-

traction occur. Thus the oxidation of HIAq is attended by a diminution in the volume of the gas equal to half the volume that would be occupied by the weight of gas that is absorbed, and the action of the gas on turpentine and on $\text{Na}_2\text{S}_2\text{O}_3\text{Aq}$ is accompanied by a diminution of volume equal to two-thirds of the volume that the O absorbed would occupy in the free state. Intermediate contractions were observed, but in no case did the contraction seem to be greater than in the action of turpentine, a result which afforded strong confirmation of the views of

Soret and Odling. An account of the early work on ozone exists in the *Handwörterbuch der Chemie*, 5, 835 (Braunschweig, 1853) (For later work v Odling, *R I* 1872).

Antozone This name was formerly given to a substance whose action with ozone resulted in their mutual decomposition. Before the relations of ozone and oxygen were established, these bodies were by some supposed to consist of oxygen in opposite conditions of electrification. Antozone was probably hydrogen peroxide. W A S

P

PACHYMOSE $\text{C}_{12}\text{H}_{22}\text{O}_{14}$. A substance, probably a glucoside, occurring in *Pachyma pinctorum*, a Chinese fungus (Champion, *J* 1872, 789). Insol water, sol alkalis, forming a solution ppd by Ca and Pb salts.

PACHYRHIZIDE A substance, not containing nitrogen, extracted from the seeds of *Pachyrhizus angulatus* (Greshoff, *B* 23, 3539). V sol alcohol, ether, and HCl , v sl. sol water and KOH . Tastes bitter, and is extremely poisonous, especially to fishes. Begins to melt at 61° , and decomposes at 161° . Yields salicylic and protocatechuic acids on fusion with potash. The alcoholic solution is acid in reaction. Probably identical with a similar body in the root-bark of *Derris* (*Pongamia elliptica*).

PÆONOL $\text{C}_{12}\text{H}_{16}\text{O}_4$, $\text{C}_6\text{H}_5\text{CO C}_6\text{H}_5(\text{OH})(\text{OMe})$ [124] [50°]. Got from the bark of *Pæonia montana* of Japan (Nagai, *B* 24, 2847). White needles (from alcohol). Yields an acetyl derivative [46.5°], a phenyl hydrazide [107°], and a crystalline oxim (Tiemann, *B* 24, 2855). Potash forms resacetophenone $\text{CH}_3\text{CO C}_6\text{H}_5(\text{OH})$, HIAq at 150° forms the same body.

PALISANDER RESIN $\text{C}_{12}\text{H}_{16}\text{O}_2$? [95°] SG 1.266. Extracted by alcohol from palisander wood, a red dye wood from Madagascar (Terreil a Wolff, *Bl* [2] 33, 435, cf Arnaudon, *Cemento*, 8, 278).

PALLADIUM Pd. At w 106.3. Mol w unknown. Melts between 1360° and 1380° (Bequerel, *C R* 57, 855), at c 1500° (Violle, *C R* 87, 981). SG 11.4 at 22.5° (Deville a Debray, *P M* [4] 50, 651). For other numbers v Clarke's *Table of Specific Gravities*, 2nd edit, 15. SH = 0.582 at 0° , = $0.582 + 0.0002t$ at t° (Violle, *l.c.*). Heat of fusion = 363 cal. CE (linear) = 0.0001176 at 40° (Fizeau, *C R* 63, 1125). EC. 12.64 at 17.2° (Ag at $0^\circ = 100$) (Matthiessen, *P* 103, 428). Volatilises in green vapour at c. 2000° . SVS c 9.3. For chief lines in emission-spectrum v *B A* 1884 434.

Occurrence—In small quantities, about 2 p.c., as metal alloyed with Pt, Ir, Os, Ru, and Rh. Certain kinds of South American gold contain from 5 to 10 p.c. Pd. Occurs also, with gold and lead selenide, in the Harz (Zinken, *P* 16, 491), and in small quantities in some specimens of silver (Röseler, *A* 180, 240). In 1803 Wollaston (*T* 1804, 428; 1805 216) separated two

new metals from Pt ore, to one of these metals he gave the name *palladium*, in allusion to the discovery, made about the same time, of the planet Pallas, and the other he called *rhodium*, because of the rose-coloured solutions of its salts (*rose* = a rose).

Formation—1 By heating PdCy_2 ,—2 By heating $\text{PdCl}_2 \cdot 2\text{KCl}$ and washing out the residual KCl —3 By ppn from solutions of its salts by means of Zn, Cu, or Fe—4 By reducing $\text{PdCl}_2 \cdot 2\text{KClAq}$ by oxalic or formic acid.

Preparation—When P. ore is heated in *aqua regia*, the Pd goes into solution, along with most of the Pt, as PdCl_2 . From this solution Pd is obtained by various processes. 1 The solution is made as nearly as possible exactly neutral by Na_2CO_3 , and HgCy_2Aq is added, PdCy_2 is ppd, along with Cu_2Cy_2 , if the ore contained Cu, the pp is washed, dried, and strongly heated, whereby Pd is obtained, mixed with Cu, the metallic residue is dissolved in HNO_3Aq , the solution is neutralised by Na_2CO_3 , and heated with HCO_2KAq and $\text{H}_2\text{C}_2\text{O}_4\text{Aq}$, whereby CO_2 is evolved plentifully, and Pd is deposited in lustrous plates, while Cu remains in solution (Dobereiner), or the metallic residue obtained by heating the cyanides may be dissolved in HNO_3Aq , the solution evaporated to dryness, and the residue strongly heated. By now treating with conc HClAq , CuO is dissolved away, while Pd remains.—2 NH_4Cl is added to the solution in *aqua regia*, and the liquid is filtered from $\text{PtCl}_2 \cdot 2\text{KCl}$, the filtrate may contain Ir, Rh, Ru, some Pt and Cu, besides Pd. These metals are ppd by addition of Zn or Fe. The pp is treated in various ways. Wilm (*B* 15, 241, v also *B* 13, 1198, 14, 629) recommends to dissolve the ppd metals in *aqua regia*, to boil the solution with excess of NaOHAq (von Schneider, *A Suppl* 5, 261), whereby the perchlorides of the metals, except that of Pt, are reduced to the lower chlorides, to acidify with HClAq , and add excess of NH_4Cl , in order to ppt $\text{PtCl}_2 \cdot 2\text{KCl}$. The filtrate from this pp. is boiled with excess of NH_4Aq , filtered, and excess of HClAq is added. After some time a yellow pp. forms, which is either almost pure $\text{PdCl}_2 \cdot 2\text{NH}_4\text{Cl}$, or if somewhat dirty-yellow in colour, it may contain $\text{Rh}_2\text{Cl}_6 \cdot 10\text{NH}_3$. This pp is treated with cold NH_4Aq , in which the Rh salt is insoluble,

and pure $\text{PdCl}_2 \cdot 2\text{NH}_4\text{Cl}$ is pptd by adding HClAq to the solution. The ppt is collected, washed with absolute alcohol, dried, and heated strongly: a grey spongy mass of Pd is obtained.

This process may be used for purifying commercial Pd.

For other methods of preparing Pd v Bunsen (A 146, 265), Philpp (D P J 220, 95), Guyard (O R 56, 1177), Rössler (Z 1866 175), Opifcius (D P J 224, 414).

Pd may be prepared from palladium containing gold by dissolving in *aqua regia*, pyg with HgCy_2Aq , and proceeding as described under 1 above. Or the ore may be fused with an equal weight of Ag and some KNO_3 , the regulus granulated, and treated with HNO_3Aq (Cook, P M [2] 23, 16), Ag pptd by addition of NaCl , and Pd pptd with other metals by Zn, the Pd may then be separated from this ppt as described under 2 above.

Properties—As prepared by heating $\text{PdCl}_2 \cdot 2\text{KCl}$ or PdCy_2 , or by pptn from solutions, Pd forms a grey metallic sponge which can be pressed together, more easily than Pt, to a compact mass. This compact form is also obtained by melting spongy Pd, Pd is a white metal, fairly malleable, ductile, and hard (somewhat softer than Pt). Compact Pd may be polished highly, it can be hammered into thin plates, and drawn into fine wire. In Brazilian gold ore Pd is found in regular octahedra, and in specimens of ore from the Harz it forms hexagonal plates. Joly (N 43, 541) obtained it in cubic octahedra by dusting Pd ribbon with powdered topaz and heating to bright redness for some time by an electric current. Heated in the O H flame to c. 2000° , Pd volatilises in greenish vapours, and condenses again to a brownish sublimate which is a mixture of metal and oxide. When the metal is melted in presence of O it absorbs O, which it gives up again on cooling (Deville a. Debray, A Ch [3] 56, 385). When strongly heated in air Pd is oxidised superficially, but the film of oxide is reduced at a higher temperature. Heated in an alcoholic flame, Pd black absorbs C and increases largely in volume. Pd absorbs H, it causes the combination of H and O when brought into electrolytic gas. Heated Pd foil brought into a mixture of NH_3 and O causes formation of NH_4NO_3 and NO_2 . Pd is more easily acted on by acids than any other of the Pt metals, it is dissolved by cold HNO_3Aq .

The at w has been determined by analysing $\text{PdCl}_2 \cdot 2\text{KCl}$ (Berzelius, P 13, 456), and by estimating Pd in PdNH_4Cl_2 by reducing in H (Keiser, Am 11, 398).

In its chemical relations Pd is closely allied to Ru and Rh, and less closely to Os, Ir, and Pt (v Noble metals, this vol p 628).

On account of its silver-like appearance, and its resistance to the action of H_2S , Pd is used for making scales and division-marks on scientific instruments, and also for coating and preserving silvered metallic ware. Pd wire is used in dentistry, an alloy with steel is used in making parts of physical instruments, and an alloy with steel, Cu, and small quantities of Au, Ni, Pt, Rh, and Ag is used in watch-making, as it is non-oxidisable, hard, and non-magnetic. Finely divided Pd is used in gas analysis for absorbing H from mixtures, and for effecting

the gradual combination of H or hydrocarbons with O (Hempel, B 23, 636, 1006).

Reactions and Combinations—1 When Pd is heated in air or oxygen Pd_2O is formed, but this oxide is reduced to Pd and O at a higher temperature.—2 Heated in hydrogen to c. 100° much H is absorbed, probably with formation of a hydride Pd_2H (v Hydrogen, vol 11 p 720).—3 Brought into a mixture of hydrogen and oxygen, in the ratio $2\text{H}_2\text{O}$, Pd black causes formation of H_2O without explosion (Coquillon, O R 83, 709).—4 Heated with sulphur Pd_2S is formed.—5 PdSe is formed by heating Pd with selenium.—6 Digested with bromine and water, PdBr_2 is produced.—7 With iodine tincture PdI_2 is produced.—8 Glowing Pd wire causes the decomposition of many hydrocarbons into C and H (Coquillon, C R 84, 1503, Wilm, B 14, 874).—9 In an alcoholic flame Pd becomes covered with C, spongy Pd increases largely in volume, probably with formation of a carbide (Wöhler, A. 184, 128).—10 Heated Pd foil brought into a mixture of ammonia and oxygen causes formation of NH_4NO_3 and NO_2 without explosion (Kraut, B 20, 1118).—11 Pd is oxidised to PdSO_4 by fusion with potassium-hydrogen sulphate.—12 Pd dissolves in acids more easily than any other of the Pt metals in nitric acid, even in the cold, it dissolves to $\text{Pd}(\text{NO}_3)_2$, in hydrochloric acid, especially when Cl is passed in, PdCl_2 is formed, in hydrotic acid, PdI_2 is produced, PdBr_2 is obtained by dissolving in hydrobromic acid, with a little HNO_3Aq , in sulphuric acid, with a little HNO_3Aq , PdSO_4 is produced, Pd dissolves in *aqua regia* to form PdCl_2 , PdCl_2 being perhaps produced at first.

Qualitative discrimination between palladium and platinum—If a drop of an alcoholic solution of I is dropped on to Pd, a black stain of PdI_2 is formed, and this stain disappears on heating, as Pt is not acted on by I tincture, this reaction serves to distinguish between the two metals.

Palladium, alloys of Alloys of Pd with several metals have been prepared. The alloy formed by heating equal weights of Pd and lead, and removing the excess of Pb, is a steel-grey powder, SG 11 255, agreeing in composition with the formula Pd_2Pb (Bauer, B 4, 451). Tin forms an alloy which seems to be a compound Pd_2Sn , (Deville a. Debray, A Ch [3] 56, 385). Alloys with Sb, As, Ba, Bi, Cu, Au, Hg, Ni, Pt, and Ag have been described (v Graham, C R 68, 1511, Mallet, C N 46, 216, Fischer, S 51, 197).

Palladium, ammonio-salts of, or Palladium-ammonium salts (*Palladamures* or *Ammoniacal palladium bases*) When HClAq is added to PdCl_2Aq containing an excess of NH_3 , the salt $\text{PdCl}_2 \cdot 2\text{NH}_3$ separates, by treatment with Ag_2O and H_2O , this salt gives the strongly alkaline base $\text{Pd}(\text{OH})_2 \cdot 2\text{NH}_3$, and by neutralising this base with acids various salts are obtained, e.g. $\text{PdSO}_4 \cdot 2\text{NH}_3$, $\text{PdCO}_3 \cdot 2\text{NH}_3$, &c. When a large excess of NH_3 is added to PdCl_2Aq and the solution is evaporated, or when a solution of $\text{PdCl}_2 \cdot 2\text{NH}_3$ in NH_4Aq is evaporated, the salt $\text{PdCl}_2 \cdot 4\text{NH}_3$ separates. Similarly, when PdSO_4 is dissolved in a large excess of NH_4Aq , the salt $\text{PdSO}_4 \cdot 4\text{NH}_3$ is obtained by evaporating this solution; decomposition of the sulphate with

BaOaq. and evaporation, gives the strongly alkaline base $\text{Pd}(\text{OH})_2 \cdot 4\text{NH}_3$, which yields salts by neutralisation with acids, e.g. $\text{PdCO}_3 \cdot 4\text{NH}_3$, $\text{PdSO}_4 \cdot 4\text{NH}_3$, &c. The compounds $\text{Pd}(\text{OH})_2 \cdot 2\text{NH}_3$ and $\text{Pd}(\text{OH})_2 \cdot 4\text{NH}_3$ represent the two series of palladium ammonio-salts. The reactions of these compounds and their derivatives lead to their representation as, in one case, compounds of the radicle PdN_2H_4 —derived from N_2H_4 by replacing 2H by Pd—and, in the other case, compounds of the radicle PdN_2H_3 —derived from PdN_2H_4 by replacing 2H by 2NH_3 . The two series of salts may be called *pallados diammonium compounds*, $\text{Pd}(\text{NH}_2\text{NH}_2)_2\text{Cl}_2$, &c., and *ammonium pallados diammonium compounds*, $\text{Pd}(\text{NH}_2\text{NH}_2\text{NH}_2\text{NH}_2)_2\text{Cl}_2$, &c. The members of the first series are also sometimes called *palladosamine compounds*, and those of the second series *pallado diamine compounds*. It is to be observed that both series are obtained from palladosochloride, PdCl_2 (cf. Iridium ammonio-salts, this vol p 47). The palladium ammonio salts have been examined chiefly by Kane (T 1842 275), Fehling (P 13, 466), Fischer (P 71, 431), and Hugo Müller (A 86, 341).

I PALLADOS DIAMMONIUM COMPOUNDS

$\text{N}_2\text{H}_4\text{PdX}'_2$, or $(\text{NH}_2\text{NH}_2)_2\text{PdX}'_2$, or perhaps $\text{NH}_2(\text{NH}_2)_2\text{PdX}'_2$

Pallados-diammonium chloride, $\text{N}_2\text{H}_4\text{PdCl}_2$ (simplest formula $\text{PdCl}_2 \cdot 2\text{NH}_3 = \text{di ammonio-palladium dichloride}$). This salt is known in two forms. 1 *Yellow crystals* are obtained by adding HClAq to PdCl_2 in excess of NH_3Aq , these crystals are scarcely sol water, sol with difficulty in cold acids, more sol hot acids, easily sol NH_3Aq , from which solution acids reppt the salt unchanged. The salt dissolves in KOH Aq , but no NH_3 is evolved even on heating, when the salt is suspended in water and Cl is passed in, solution is effected, addition of NH_3Aq now ppts the red form of the salt, and boiling with KOH Aq evolves NH_3 and leaves a liquid from which crystals of $\text{PdCl}_2 \cdot 2\text{NH}_3\text{Cl}$ separate. The continued passage of Cl produces $\text{PdCl}_2 \cdot 2\text{NH}_3\text{Cl}$, and finally PdCl_2Aq . 2 *A red salt* of the same composition is obtained by dissolving the yellow salt in conc HClAq and adding NH_3Aq , also by adding a slight excess of NH_3Aq to a rather dilute cold solution of PdCl_2Aq . The yellow salt is produced by dissolving the red variety in NH_3Aq and adding excess of HClAq , also by heating the red salt to 200° . Jørgensen (Gm K 3, 1242) regarded the red salt as polymeric with the yellow, and gave it the formula $\text{Pd}(\text{NH}_2\text{NH}_2)_2\text{Cl}_2 \cdot \text{Cl}_2\text{Pd}$. The salts may be isomeric, one being $\text{PdNH}_2\text{ClNH}_2\text{Cl}$ and the other $\text{PdNH}_2\text{Cl}_2\text{NH}_2$.

Pallados-diammonium hydroxide,

$\text{N}_2\text{H}_4\text{Pd}(\text{OH})_2$ (simplest formula $\text{Pd}(\text{OH})_2 \cdot 2\text{NH}_3 = \text{di ammonio palladium hydroxide}$). This base is obtained by decomposing the corresponding chloride with moist Ag_2O in presence of water, or the sulphate with BaO Aq , the solution thus obtained is yellowish, odourless, and has a strongly alkaline taste and reaction. The base is obtained as a curdy crystalline mass by evaporating at the ordinary temperature in absence of air. The solid rapidly absorbs CO_2 from the air, forming $\text{N}_2\text{H}_4\text{PdCO}_3$, it is decomposed by heating to somewhat above 100° . The base is sol. water, the solution decomposes NH_3 salts,

and ppts $\text{CuO} \cdot \text{H}_2\text{O}$ and Ag_2O from solutions of salts of Cu and Ag. The solution may be boiled with very slight change, on boiling with alcohol, Pd is pptd. The other salts of this series which have been described are as follows $\text{M} = \text{PdN}_2\text{H}_4$, —bromide MBr , carbonate MCO_3 , fluoride $\text{MF}_3(?)$, iodide MI , nitrate $\text{M}(\text{NO}_3)_3$, nitrite $\text{M}(\text{NO}_2)_3$, sulphate MSO_4 , and sulphite MSO_3 .

II AMMONIUM PALLADOS DIAMMONIUM COMPOUNDS $\text{N}_2\text{H}_4(\text{NH}_2)_2\text{PdX}'_2$, or perhaps $(\text{NH}_2\text{NH}_2)_2\text{PdX}'_2$

Ammonium pallados diammonium chloride $\text{Pd}(\text{NH}_2\text{NH}_2)_2\text{Cl}_2$, (or $\text{Pd}(\text{NH}_2\text{NH}_2)_2\text{Cl}_2 = \text{pallado diamine chloride}$, simplest formula $\text{PdCl}_2 \cdot 4\text{NH}_3 = \text{tetrammonio palladium dichloride}$). This salt is obtained in large, colourless, monoclinic prisms, with one molecule of water of crystallisation, by evaporating a solution of yellow $\text{N}_2\text{H}_4\text{PdCl}_2$ with excess of NH_3 , or a solution of PdCl_2 in considerable excess of NH_3Aq . Heating the salt to 120° , or adding acid to a solution of the salt, causes formation of yellow $\text{N}_2\text{H}_4\text{PdCl}_2$. Addition of PdCl_2 to a solution of the salt causes pptn of flesh red $\text{Pd}(\text{NH}_2\text{NH}_2)_2\text{Cl}_2 \cdot \text{PdCl}_2$ (Kane, Fehling). Treatment of $\text{Pd}(\text{NH}_2\text{NH}_2)_2\text{Cl}_2$ with NH_4Cl and *aqua regia* is said to produce $\text{Pd}(\text{NH}_2\text{NH}_2)_2\text{Cl}_2 \cdot \text{PdCl}_2$ (Croft, C N 16, 53).

Ammonium pallados diammonium hydroxide $\text{Pd}(\text{NH}_2\text{NH}_2)_2(\text{OH})_2$, (or $\text{Pd}(\text{NH}_2\text{NH}_2)_2(\text{OH})_2 = \text{pallado diamine hydroxide}$, simplest formula $\text{Pd}(\text{OH})_2 \cdot 4\text{NH}_3 = \text{tetrammonio palladium hydroxide}$). Obtained as a crystalline mass by decomposing a solution in water of the sulphate $\text{Pd}(\text{NH}_2\text{NH}_2)_2\text{SO}_4 \cdot \text{H}_2\text{O}$, produced by adding large excess of NH_3Aq to PdSO_4Aq and evaporating with BaO Aq , filtering and evaporating. The solution is odourless, it is strongly alkaline, and ppts hydroxides from solutions of salts of Al, Co, Cu, Fe, and Ni, but not from salts of Ag, it sets NH_3 free from NH_4ClAq . The solution absorbs CO_2 from the air, it is decomposed by boiling with organic matter. If this base is neutralised exactly by acids, salts of the base are obtained, e.g. $\text{Pd}(\text{NH}_2\text{NH}_2)_2\text{CO}_3$, $\text{Pd}(\text{NH}_2\text{NH}_2)_2\text{SO}_4$, &c., excess of acid, especially of a haloid acid, causes decomposition into a pallados ammonium salt and a salt of NH_3 , e.g. $\text{Pd}(\text{NH}_2\text{NH}_2)_2(\text{OH})_2 + 4\text{HClAq} = \text{PdN}_2\text{H}_4\text{Cl}_2 + 2\text{NH}_4\text{ClAq} + 2\text{H}_2\text{O}$. The other salts of this series which have been described are as follows $\text{M} = \text{Pd}(\text{NH}_2\text{NH}_2)_2$, —bromide MBr , and the double compound $\text{MBr} \cdot \text{PdBr}$, carbonate MCO_3 , fluoride $\text{MF}_3(?)$, iodide MI , nitrate $\text{M}(\text{NO}_3)_3$, double nitrate $\text{M}(\text{NO}_2)_3$, $\text{Pd}(\text{NO}_3)_3$, silicofluoride $\text{MSiF}_6(?)$, sulphate MSO_4 , sulphite MSO_3 .

The reactions of such substituted ammonias as NH_2Me , NH_2Et , and NH_2Ph on salts of Pd produce compounds analogous with the ammoniacal bases described above, e.g. *pallados diethylammonium chloride* $\text{N}_2\text{H}_5\text{Et}_2\text{PdCl}_2$, and *pallados diphenylammonium chloride* $\text{N}_2\text{H}_5\text{Ph}_2\text{PdCl}_2$, have been described (Müller, A 86, 341).

The substitution of *tri ethylarsine* and *tri-ethylphosphine* for ammonia, in the reactions with Pd salts, produces compounds similar to the N-containing bases, e.g. *pallados di-ethylphosphonium chloride* $\text{P}_2\text{Et}_2\text{PdCl}_2$.

(Cahours a Gal, *C R* 70, 897), and *pallados di-ethylarsonium chloride* $\text{As}_2\text{Et}_4\text{PdCl}_2$ (*C a G*, *C R* 71, 208)

Palladium, bromide of Only one bromide of Pd is known, and that has not been prepared free from impurities

PALLADIUM DIBROMIDE PdBr_2 (*Palladous* or *palladoso bromide*) The brownish solid obtained by dissolving Pd in a mixture of HBrAq and HNO_3Aq , or digesting finely divided Pd with BrAq , and evaporating, probably has the composition PdBr_2 , but it has not been obtained pure. **Double compounds of PdBr_2 with bromides of Ba, Mn, K, and Zn** have been prepared, but not fully examined, by von Bonsdorff (*P* 19, 347, 431), the K salt, K_2PdBr_4 , crystallises in thick rhombic forms (Joannis, *C R* 95, 295)

Palladium, chlorides of Two chlorides of Pd have been isolated, PdCl and PdCl_2 , a third, PdCl_3 , is known in combination. As none of these chlorides has been gasified, the formulæ are not necessarily molecular (*v* PALLADIUM, HALOID COMPOUNDS of, *post*) None of the chlorides has been formed by the direct union of Pd and Cl

PALLADIUM DICHLORIDE PdCl_2 (*Palladous* or *palladoso chloride*) By dissolving Pd in HClAq , Cl being passed into the liquid or a little HNO_3 being present, evaporating repeatedly with HClAq to remove HNO_3 , and finally concentrating, red brown prismatic needles of $\text{PdCl}_2 \cdot 2\text{H}_2\text{O}$ are obtained, at a moderate temperature H_2O is given off, and PdCl_2 remains as a black brown crystalline solid (Kane, *T* 1842 275, Bottger, *P* 106, 495) PdCl_2 is also obtained, partly as a sublimate and partly as a garnet red crystalline residue, by heating Pd_2S in a stream of Cl (Schneider, *P* 141, 519) PdCl_2 dissolves in water to form a dark red liquid, from which some oxychloride, $\text{Pd}_2\text{O}_2\text{Cl}_2$, separates on evaporation. In water containing HCl , PdCl_2 dissolves very readily. The solution is dark brown, and on treatment with bases it gives salts of the form M_2PdCl_4 —*chlorpalladites* (*v infra*), this solution may be supposed to contain *pallados chlorhydric acid* H_2PdCl_4

Double salts, chlorpalladites $\text{PdCl}_2 \cdot 2\text{MCl}$ or M_2PdCl_4 . These salts are obtained by the reaction of various oxides with PdCl_2 in dilute HClAq , or by adding various chlorides to conc PdCl_2Aq

Ammonium palladium dichloride or *Ammonium chlorpalladite* $\text{PdCl}_2 \cdot 2\text{NH}_4\text{Cl}$ or $(\text{NH}_4)_2\text{PdCl}_4$. Olive-green needles with a bronze coloured lustre, e sol water, insol conc alcohol. When strongly heated gives residue of finely divided Pd. Obtained by adding NH_4Cl to PdCl_2Aq containing HCl (Kane, *T* 1842 275, Wilm, *B* 18, 1202)

Potassium palladium dichloride, or *Potassium chlorpalladite* $\text{PdCl}_2 \cdot 2\text{KCl}$ or K_2PdCl_4 . Golden yellow needles formed by adding KClAq to conc PdCl_2Aq . The crystals are quadratic prisms (Joannis, *C R* 95, 295), they are fairly sol. in cold water, and much more sol. in hot water. Alcohol ppts the salt from a hot conc aqueous solution. An aqueous solution boiled with alcohol or SO_2 gives a pp of Pd, the dry salt is very slow, decomposed by heat to KCl and Pd, the decomposition is slow even in presence of oxalic acid (Rössler, *Z* 1866 175), but rapid in a stream

of H. The other chlorpalladites which have been described are $\text{PdCl}_2 \cdot \text{AlCl}_3 \cdot 10\text{H}_2\text{O}$ (Welkow, *B* 7, 803), BaPdCl_4 (von Bonsdorff, *P* 19, 347, 431), $\text{BePdCl}_4 \cdot 6\text{H}_2\text{O}$ (W, *lc*), CdPdCl_4 , CaPdCl_4 , MgPdCl_4 , MnPdCl_4 , NiPdCl_4 , ZnPdCl_4 , $\text{Co}_2\text{Cl}_2 \cdot 12\text{NH}_3$, 2PdCl_2 (Gibbs, *Am S* [2] 37, 58), $\text{Co}_2\text{Cl}_2 \cdot 10\text{NH}_3$, 2PdCl_2 (Carstanjen, *Gm K* 3, 1254), $\text{PdCl}_2 \cdot \text{Hg}_2\text{Cl}_2 \cdot 5\text{NH}_4\text{Cl}$ (Wilm, *B* 13, 1202)

PALLADIUM SUBCHLORIDE PdCl or Pd_2Cl_3 . Small quantities of this chloride are said to be formed, as a dark reddish brown crystalline solid, when PdCl_2 is heated to redness, the chloride deliquesces in air, and always contains PdCl_2 , and Pd (Kane, *T* 1842 275)

PALLADIUM TETRACHLORIDE PdCl_4 (*Palladi* or *pallados chloride*) This chloride probably exists in a solution of Pd in *aqua regia*, or of PdO_2 in conc HClAq , the compound has not been isolated. As this solution yields salts of the form M_2PdCl_6 when treated with various metallic chlorides, it perhaps contains *palladichlor hydric acid* H_2PdCl_6

Double salts, chlorpalladates $\text{PdCl}_2 \cdot 2\text{MCl}$ or M_2PdCl_4 . These salts are formed by adding metallic chlorides to a solution of Pd in *aqua regia*, or of PdO in conc HClAq , or to a solution of PdCl_2 in HClAq into which Cl has been passed, some of them are also produced by passing Cl into a solution of the corresponding chlorpalladite

Ammonium palladium tetrachloride or *Ammonium chlorpalladate* $\text{PdCl}_2 \cdot 2\text{NH}_4\text{Cl}$ or $(\text{NH}_4)_2\text{PdCl}_6$. A bright red pp obtained by adding NH_4Cl to conc PdCl_2Aq saturated with Cl or treated with conc HNO_3Aq *S G* 2 418 (Topsoë, *J* 1870 393). Reduced with difficulty, treated with NH_3Aq , N is evolved and $\text{PdCl}_2 \cdot 2\text{NH}_3$ is formed (H Müller, *A* 86, 341)

Potassium palladium tetrachloride or *Potassium chlorpalladate* $\text{PdCl}_2 \cdot 2\text{KCl}$ or K_2PdCl_6 . A cinnabar red powder, consisting of small regular octahedra, *S G* 2 738, obtained by mixing KClAq with PdCl_2Aq saturated with Cl, or with Pd in *aqua regia*, also formed by treating $\text{K}_2\text{PdCl}_4\text{Aq}$ with Cl (Topsoë, *J* 1870 393, Croft, *C N* 16, 53). When the aqueous solution is treated with NH_3Aq , N is evolved and K_2PdCl_6 is formed

The other chlorpalladates which have been described are $\text{BePdCl}_6 \cdot 8\text{H}_2\text{O}$ (Welkow, *B* 7, 38), $\text{MgPdCl}_6 \cdot 6\text{H}_2\text{O}$ (Topsoë, *J* 1870 393), $\text{NiPdCl}_6 \cdot 6\text{H}_2\text{O}$ (*T, lc*), $\text{ZnPdCl}_6 \cdot 6\text{H}_2\text{O}$ (*T, lc*)

Palladium, cyanides of, and **Double cyanides**, *v* vol II p 343

Palladium, fluoride of, PdF_2 . This compound is produced, according to Berzelius, by adding HFAq to conc $\text{Pd}(\text{NO}_3)_2\text{Aq}$, it is described as a brown pp scarcely sol. water or HFAq , and as forming double salts with alkali fluorides

Palladium, haloid compounds of The haloid compounds of Pd have not been at all fully investigated. The following table presents the compositions of those which have been isolated —

PdX	PdX_2	PdX_3
PdCl	PdF	PdCl_2
	PdBr	PdI
		in solution and combination

The compounds PdX_2 form double salts $\text{PdX}_2 \cdot 2\text{MX} = \text{M}_2\text{PdX}_4$, and PdCl_3 forms $\text{PdCl}_3 \cdot 2\text{MCl} = \text{M}_2\text{PdCl}_6$, the acids H_2PdCl_6 and

H_2PdCl_4 , probably exist in solution. The chloropalladates M_2PdCl_4 are not very easily reduced, the chloropalladites M_2PdCl_4 are readily oxidised to chloropalladates. None of the haloid compounds has been gasified, the formulae are the simplest that express the composition, but they are not necessarily molecular. $PdBr_2$ is formed by digesting together finely-powdered Pd and Br in presence of water. $PdCl_2$ and $PdCl$ are easily soluble in water, PdF_2 , $PdBr_2$, and PdI_2 are insoluble or but slightly soluble in water.

Palladium, hydride of Pd absorbs H very freely. A piece of Pd foil which has been strongly heated *in vacuo* absorbs 648 times its volume of H at 90° – 97° . When electrolytically *ppd.* Pd is used as the negative pole in the electrolysis of water, it absorbs 962 vols H. The physical properties of Pd charged with H make it very probable that a definite compound is formed, and that this compound has the composition Pd_2H (for details *v* HYDROGEN, vol. II. P 720).

Palladium, hydroxide of, *v.* Palladium, oxides and hydrated oxides of

Palladium, iodide of, PdI_2 . When $KIAq$ is added to $PdCl_2Aq$ or $Pd(NO_3)_2Aq$, an almost black *pp* of $PdI_2 \cdot H_2O$ is obtained, which loses H_2O *in vacuo* (Lassaigne, *J Chim méd* 11, 57). PdI_2 is scarcely sol water, alcohol, or ether, slightly sol $HIAq$, easily sol $KIAq$, from which solution dark-red deliquescent crystals of potassium iodopalladate, K_2PdI_4 , separate on concentration. As PdI_2 is scarcely sol water, the *ppn* of this salt may be used in the quantitative estimation of I in presence of Cl and Br. PdI_2 is decomposed to Pd and I at somewhat above 350° , the last traces of I are not removed by heat alone, but by heating in H (Berzelius).

Palladium, oxides and hydrated oxides of
Three oxides have been isolated, Pd_2O , PdO , and PdO_2 , another, Pd_2O_3 , intermediate between PdO and PdO_2 , seems to exist. PdO and PdO_2 seem to form hydrates, but the exact composition of these is not settled. These oxides all react with acids to form salts corresponding with PdO , Pd_2O at the same time forming Pd, and PdO_2 evolving O. It is, however, possible that some salts may be obtained corresponding with PdO_2 . All the oxides are reduced to Pd when strongly heated. The examination of the oxides of Pd is very incomplete.

PALLADIUM SUBOXIDE Pd_2O A black powder obtained by heating to low redness the *pp* formed by adding Na_2CO_3 to solution of a salt of PdO (Kane, *T* 1842 276), or by heating Pd black in a stream of air (Wilm, *B* 15, 2225). Decomposed to Pd and O by heating to full redness, reduced by H at ordinary temperatures, reacts with acids to form salts of PdO with excess of Pd.

PALLADIUM MONOXIDE PdO (Palladous oxide) A black powder, prepared by heating $Pd(NO_3)_2$ or by very gently heating a Pd salt with K_2CO_3 or Na_2CO_3 , and washing the residue with water. Reduced very easily by H (Wöhler, *A* 174, 160), gives Pd and O when heated to full redness. Soluble in acids with difficulty, forming salts PdX ($X = SO_3, 2NO_3, CO_3$, &c).

Hydrated palladium monoxide The dark-brown *pp* obtained by adding K_2CO_3 or Na_2CO_3 to solution of a salt of PdO was described

by Berzelius as $PdO \cdot H_2O$, but it may be a basic carbonate, heated to low redness this *pp* gives Pd_2O .

PALLADIUM DIOXIDE PdO_2 (Palladic oxide) A black powder, obtained by *ppg* $PdCl_2Aq$ or K_2PdCl_4Aq with excess of $KOHAc$, washing acid boiling with water, and drying at 100° , also by the action of ozone on compounds of Pd, and by the decomposition of water using Pd as the positive pole (Wöhler, *A* 146, 375, Maillet, *C R* 94, 860, 1186). At low red heat gives PdO and O, and at a higher temperature all O is given off. Treated with dilute $HClAq$, gives $PdCl_2Aq$ and Cl, with conc $HClAq$, probably forms $PdCl$. No corresponding salts have been isolated with certainty.

Hydrated palladium dioxide The *pp* obtained by adding excess of $KOHAc$ to $PdCl_2Aq$ or K_2PdCl_4Aq is probably $PdO_2 \cdot xH_2O$, but the exact composition of this *pp* is not known.

PALLADO PALLADIC OXIDE $Pd_2O_3 = 4PdO \cdot PdO_2$ According to Schneider (*P* 141, 519), an oxide of this composition is obtained by melting K_2PdS_2 or Na_2PdS_2 with KNO_3 and KOH , washing with water, and treating the residue with *aqua regia*. It is described as a dull brown powder, which gives off all its O when heated to redness in air, and is reduced by H at the ordinary temperature.

Palladium, salts of. Compounds obtained by replacing hydrogen of acids by Pd All the salts of Pd which have been isolated with certainty correspond with the oxide PdO , the salts of Pd are generally obtained by dissolving Pd in the various acids, with a little HNO_3Aq added, or in some cases by double decomposition from $PdCl_2Aq$ or $Pd(NO_3)_2Aq$. Only a very few salts have been examined, besides the salts of the haloid acids, the carbonate, nitrate, nitrite, sulphate, and sulphite have been isolated (*v* CARBONATES &c). $PdCl_2$, corresponding with PdO_2 , probably exists in solution, and several compounds of this salt—the chloropalladates—have been obtained (*v* Palladium tetrachloride, p 795).

Palladium, selenide of, $PdSe$ A grey in fusible solid, resembling osmium, with which it is perhaps isomorphous (Rössler, *A* 180, 244), formed by heating together Pd and Se (Berzelius).

Palladium, sulphides of Three sulphides are known, corresponding with the three oxides. The highest sulphide, PdS_2 , reacts with sulphides of more positive elements as a sulphanydride, forming *sulpho-palladates* M_2PdS_2 .

PALLADIUM SUBSULPHIDE Pd_2S A grey, hard solid, SG 7.803 at 15° . Prep red by fusing together, for 15–20 minutes over a blow-pipe, 100 parts yellow $N_2H_4PdCl_2$ (p 794), or 5 parts PdS , with 6 parts dry K_2CO_3 or Na_2CO_3 , 6 parts S, and 3 parts NH_4Cl , the fused mass is treated with water, and the lustrous blue-violet crystals of $K_2Pd_2S_3$, which are mixed with the Pd_2S , are removed by agitating with water. Pd_2S is very stable, it is not acted on by acids, scarcely by *aqua regia*, melts at a red heat without change, loses S slowly when very strongly heated (Schneider, *P* 141, 519).

PALLADIUM MONOSULPHIDE PdS (Palladous sulphide) Obtained by heating Pd, or certain

Pd salts, with S, also by ppg a salt of PdO by H₂S. Prepared in the dry way, PdS forms a blue white, lustrous, metal-like, very hard solid, prepared in the wet way, it is a black powder. Heated in air, slowly oxidises to a basic sulphate, heated in Cl forms PdCl₂ and S₂Cl₂ (Fellenberg, P 50, 66). A colloidal soluble form of PdS was obtained by Winkinger (Bl [2] 49, 462) by ppg from an extremely dilute solution and dialysing.

PALLADIUM DISULPHIDE PdS₂ (*Palladic sulphide*). A dark-brown powder, scarcely acted on by HNO₃Aq soluble *aqua regia* without separation of S (Schneider, P 141, 519), heated in a stream of CO, forms PdS and then Pd₂S. Obtained by decomposing solution of a sulphopalladate by dilute HClAq, and washing the pp *eg* Na₂PdS₂Aq + 2HClAq = 2NaClAq + PdS₂ + H₂S. This sulphide reacts as an acid anhydride with the sulphides of several of the more positive metals.

Sulphopalladates. These salts belong to two series, M₂PdS₂ and M₂Pd₂S₂ = M₂PdS, Pd₂S (Schneider, P 141, 519, 148, 625).

Sodium sulphopalladate Na₂PdS₂ (*Sodium-palladium sulphide*). This salt has not been obtained pure, as prepared by fusing 1 part N₂H₄PdCl₂ (p 794), or 5 part PdS, with 6 parts dry Na₂CO₃ and 6 parts S, to full redness, and washing the residue with water, it forms reddish-brown needles, which dissolve in water to form a brown liquid decomposed by HClAq with ppn of PdS₂.

Silver sulphopalladate Ag₂PdS₂. A black brown powder obtained by adding the Na salt to an alcoholic solution of AgNO₃.

The members of the other series of sulphopalladates, M₂Pd₂S₂, may be looked on as double compounds of M₂PdS₂ and Pd₂S, or as the salts of a hypothetical sulphopalladic acid H₂Pd₂S₂. These salts may perhaps be termed *meta sulphopalladates*.

Potassium meta-sulphopalladate, K₂Pd₂S₂, or K₂PdS₂Pd₂S. Formed by melting 2 parts N₂H₄PdCl₂ (p 794), or 1 part PdS, with 12 parts K₂CO₃ and 12 parts S, at a red heat, and washing the fused mass with water (Schneider, P 141, 519). Six sided, blue-violet, metal like crystals, insol water, treated with HClAq, K is removed but no H₂S is evolved, and the crystals become steel-grey, perhaps H₂Pd₂S₂ may be formed, but if so this soon decomposes, and a sulphide, said to be Pd₂S₂, remains (Schneider, P 141, 625). Another compound, which may perhaps be K₂PdS₂ (corresponding with the hypothetical acid Pd(SH)₂) is formed, along with Pd, by heating in H.

Silver meta-sulphopalladate Ag₂Pd₂S₂, or Ag₂PdS₂Pd₂S. A white-grey, lustrous, crystalline pp obtained by treating the K salt with an alcoholic solution of AgNO₃.

Palladium, sulphocyanides of, *vol ii*, p 350.

Palladium, thio. salts of, *cf. Sulphopalladates under Palladium, sulphides of, supra*.

M M P M.

PALM OIL. Extracted from the fruit of *Elais guineensis*. Soft orange mass containing palmitic and oleic acids and their glycerides (Pelouze a. Boudet, A 29, 42, Guibourt, J. Chem Med. 1, 177, Henry, J. Ph. 51, 241).

From palm kernels an oil is obtained containing the glycerides of oleic, stearic, palmitic, myristic, lauric, decolic, oleic, and hexoic acids (Oudemans, J pr [2] 2, 898).

PALMELLIN. A substance resembling hæmoglobin, which occurs in *Palmella cruenta*, a red fungus (Phipson, C R 89, 816, 1078, C N. 41, 216).

PALMITIC ACID C₁₆H₃₂O₂. Mol w 256. [80.75°] (Reissert, B 23, 2248) (271.5°; *v* at 100 mm) (Krafft, B 12, 1670, 16, 1721) (339°–356°) (Carnelley a. Williams, B 12, 1360) S G. (liquid) ²² 853 S (alcohol) 9.2 at 19.5° (C a. S) H C (solid) 2,371,788 (Longuine, A Ch [6] 11, 223). Occurs as glyceride in a very large number of animal and vegetable fats and fixed oils (Chevreul, *Recherches sur les corps gras*, Fremy, A 36, 44, Stenhouse, A 36, 50, Stahmer a. Meyer, A 43, 335, Schwarz, A 60, 69, Heintz, A 80, 299, 88, 298, 92, 291, von Böck, J pr. 49, 295, Berthelot, A Ch [3] 41, 216, 432, 47, 297, Maskelyne, C J 8, 1). Its myrcyl ether occurs in bees wax (Brodie, A 71, 150), and in human fat (Heintz), and its cetyl ether in spermaceti (L Smith, A 42, 241). Occasionally occurs in the free state, as in palm oil, and in *Lycopodium* spores (Langer, Ar Ph [3] 27, 625).

Formation—1 By saponification of palmitin, spermaceti, and melissin—2 By heating cetyl alcohol with potash lime (Dumas a. Stas, A Ch [2] 73, 113)—3 Together with acetic acid by fusing oleic or elaidic acid with potash (Varentz, A 36, 209)—4 By saponifying bird-lime with alcoholic potash (Divers a. Kawakita, C J 53, 271)—5 By saponifying the wax of *Myrica cerifera* (Chittenden a. Smith, Am 6, 217).

Preparation—1 Japanese wax (3 pts) is saponified with KOH (1 pt) and water (1 pt), the solution ppd by HClAq, and the acid rectified *in vacuo* (Krafft, B 21, 2265)—2 The fatty acids obtained by saponifying fats are dissolved in alcohol, and fractionally ppd with an alcoholic solution of lead acetate or a conc aqueous solution of barium or magnesium acetate (*cf. vol i* p 56).

Properties—Small hard crystals (from alcohol), insol water, *v* sol boiling alcohol and ether. Slightly decomposed on distillation. Chlorination, in presence of SbCl₃, yields CCl₄, C₂Cl₄, and C₃Cl₄ [224°] (Hartmann, B 24, 1018). Oxidation by alkaline KMnO₄ yields oxalic, succinic, adipic, acetic, butyric, hexoic, oxyvaleric, and dioxypalmitic acids (Gröger, M 8, 486). Oxidation by nitric acid (S G 136) yields succinic and glutamic acids (Carette, C R 102, 692). The Ba salt distilled with NaOMe yields pentadecane (Mai, B 22, 2133).

Salts—(NH₄)HA', —KA' pearly scales (from alcohol), insol ether. Dissolves in a small quantity of water, but a larger quantity ppt. KHA', which crystallises from alcohol in pearly scales [100°] (Schwarz)—NaA' laminae.—NaHA'. Got by adding hot water (1500 pts) to NaA'. Insol water, *v* sol hot alcohol.—BaA', S. (alcohol) 0035 at 20°. Pearly crystalline powder.—CaA', S (alcohol) 0103 at 20°.—MgA', [120°] (H)—PbA', [112°] Powder.—CuA', greenish blue powder.—AgA', amorphous, sl. sol water. Blackened by light.

Methyl ether MeA' [38°] (Berthelot)

Ethyl ether EtA' [24°] (Heintz)
Isoamyl ether $\text{C}_5\text{H}_{11}\text{A}'$ [9°] (Berthelot),
 [18-5°] (Duffy, *C J* 5, 814) Waxy

Octyl ether $\text{C}_8\text{H}_{17}\text{A}'$ [86°] Formed from the octyl alcohol of castor oil

Dodecyl ether $\text{C}_{12}\text{H}_{25}\text{A}'$ [41°] (K.).

Tetradecyl ether $\text{C}_{14}\text{H}_{29}\text{A}'$ [48°]

Hexadecyl ether $\text{C}_{16}\text{H}_{33}\text{A}'$ [54°] (Krafft, *B* 16, 3023) Probably identical with the *cetyl ether* $\text{C}_{16}\text{H}_{33}\text{A}'$ [49°], which is the chief component of spermaceti, from which it is prepared by crystallisation from boiling alcohol

Octadecyl ether $\text{C}_{18}\text{H}_{37}\text{A}'$ [59°]

Ceryl ether $\text{C}_{20}\text{H}_{41}\text{A}'$ [79°] The chief constituent of poppy wax (Hesse, *B* 3, 639)

Myrcyl ether $\text{C}_{20}\text{H}_{41}\text{A}'$ [72°] The constituent of bees wax that is insoluble in alcohol (Brodie, *A* 71, 159)

Glyceryl derivatives or Palmitins v vol ii p 621

Phenyl ether $\text{C}_6\text{H}_5\text{A}'$. [45°] (250° at 15 mm) (Krafft a Burger, *B* 17, 1379)

p-Tolyl ether $\text{C}_6\text{H}_4\text{A}'$ [47°] (258° at 15 mm)

Chloride $\text{C}_{16}\text{H}_{33}\text{OCl}$ [c 12°] (192.5° at 15 mm) (Krafft a Burger)

Amide $\text{C}_{16}\text{H}_{33}\text{CONH}_2$ [107°] Formed by the action of NH_3 on the chloride or ether, and also by heating tetradecyl malonic amide (Carlet, *Bl* 1859, 175, Krafft a Stauffer, *B* 15, 1730, Hell a Jordanoff, *B* 24, 990)

Anilide $\text{C}_{16}\text{H}_{33}\text{CONHPh}$ [90.5°] (283° at 17 mm) Made by boiling palmitic acid with excess of aniline (Hell a Jordanoff, *B* 24, 943) Silky needles (from alcohol), v e sol ether

Anhydride $(\text{C}_{16}\text{H}_{31}\text{O})_2\text{O}$ [64°] (Villier, *B* 9, 1932)

Nitride $\text{C}_{16}\text{H}_{31}\text{ON}$ [31°] (252° at 100 mm) $\text{SG } \frac{1}{2}$ 8186, $\frac{1}{2}$ 778 Formed by distilling the amide with P_2O_5 (K a S), and also by heating cyano palmitic acid (H a J) Six-sided tables Reduced by alcohol and sodium to hexadecylamine (Krafft, *B* 22, 811)

References — DI BROMO PALMITIC ACID AND HEXADECOIC ACID

PALMITIC ALDEHYDE $\text{C}_{16}\text{H}_{32}\text{O}$ [58°] (198° uncor at 22 mm) Prepared by distilling a mixture of calcium palmitate and calcium formate (Krafft, *B* 13, 1416, 16, 1714) Plates, sl sol ether Combines with NaHSO_3

Palmitic aldehyde $\text{C}_{16}\text{H}_{32}\text{O}$ [47°] S (alcohol) 64 at 16°, 12 at 78° S (ether) 16 at 16° Got by oxidation of cetyl alcohol with chromic acid mixture (Fridau, *A* 83, 23, Dollfus, *A* 181, 287) Crystalline

PALMITIN v GLYCERIN

PALMITIC ACID $\text{C}_{16}\text{H}_{32}\text{O}_2$ Mol w 252 [42°] Formed by the action of alcoholic potash at 180° on di bromo-palmitic acid (Schröder, *A* 143, 22) Silky needles (from alcohol), insol water, v sol ether Br forms $\text{C}_{16}\text{H}_{32}\text{Br}_2\text{O}_2$ and $\text{C}_{16}\text{H}_{32}\text{Br}_2\text{O}_2$ Fuming nitric acid oxidises it to suberic acid and suberic aldehyde — BaA' — AgA' amorphous powder blackened by light.

Reference — BROMO PALMITIC ACID

PALMITONE $(\text{C}_{16}\text{H}_{31})_2\text{CO}$ *Di-pentadecyl ketone* [88°] $\text{SG } \frac{1}{2}$ 799, $\frac{1}{2}$ 794 Formed by distilling calcium or barium palmitate with lime (Piria, *C R* 34, 140, Maskelyne, *C J* 8, 1, Krafft, *B* 15, 1714) Got also by heating palmitic acid with P_2O_5 at 210° (Kipping, *C J*

57, 986). Silvery plates, v sol alcohol and benzene Does not unite with NaHSO_3 (Lumprecht, *A* 94, 246) Bromine forms $\text{C}_{16}\text{H}_{32}\text{Br}_2\text{O}$ [55°] (Herez, *A* 186, 269)

Oxim $(\text{C}_{16}\text{H}_{31})_2\text{CNOH}$ [59°] Needles

PALMITOXYLIC ACID $\text{C}_{16}\text{H}_{32}\text{O}_3$ [67°]

Formed, together with suberic acid and suberic aldehyde by the action of fuming HNO_3 on palmitic acid (Schröder, *A* 143, 35) Laminæ (from alcohol), insol water, v sol ether — AgA' white pp, turned violet by light

PANAQUILONE $\text{C}_{16}\text{H}_{32}\text{O}_3$ Occurs in the root of *Panax quinquefolius* (Garrigues, *A* 90, 231) Amorphous powder, v sol water, and alcohol, insol ether Has a bitter sweet taste

Its solution is ppd by tannin H_2SO_4 forms a purple solution from which water ppts panacone $\text{C}_{16}\text{H}_{32}\text{O}_3$ (?) a crystalline powder HClAq also forms CO_2 and panacone on heating

PANICOLE $\text{C}_{16}\text{H}_{32}\text{O} + \text{C}_2\text{H}_5\text{OMe}$ [285°]

A crystalline substance in oil of millet (Kassner, *Ar Ph* [2] 25, 395, 26, 536) HClAq at 160° forms MeCl and $\text{C}_{12}\text{H}_{24}\text{O}$ [78°]

PAPAIN v PROTEIDS

PAPAVERINE $\text{C}_{20}\text{H}_{21}\text{NO}_4$ *fa.*

C(OMe)CHCCHCHN
 C(OMe)CHCCHCHN

$\text{C(OMe)CHCCHCHNCH}_2\text{C}_6\text{H}_4\text{(OMe)}$ [13.4] (Goldschmidt, *M* 9, 330, 349) [147°] S (ether) 4 at 10° Occurs in opium (Merck, *A* 66, 125, 73, 50, Anderson, *T E* 21, Pt 1, Hesse, *A* 153, 75, Suppl 8, 261, Z [2] 7, 641)

Preparation — 1 The aqueous extract of opium is ppd with Na_2CO_3 , the pp dissolved in ether and shaken with dilute HOAc The acetic acid solution is ppd by NaOH The pp digested with oxalic acid solution yields crystalline papaverine oxalate It is better, however, to dissolve the pp in acetic acid, remove thebaine by tartaric acid, ppt the mother liquor with ammonia, wash the pp with alcohol, and then treat with oxalic acid The oxalate is recrystallised from hot water, ppd by CaCl_2 , the filtrate ppd by NH_3 and the papaverine recrystallised from alcohol (Hesse) — 2 A solution of the hydrochlorides of the opium bases is ppd with conc NaOAc The pp, consisting of narcotine and papaverine, is dissolved in HClAq and diluted till it contains only 25 pc of narcotine K_2FeCy_6 is then added and, after 24 hours, the pp of papaverine ferrioyanide is collected and decomposed by NaOHAq (Plugge, *Ar Ph* [3] 25, 343)

Properties — Trimetric prisms (from ether alcohol), $a b c = 32.1 : 42$ Narcotic Nearly insol hot water, v sol hot alcohol and chloroform, m sol hot benzene Cannot be sublimed Inactive to light (Goldschmidt, *M* 9, 42) Conc H_2SO_4 forms a colourless solution, becoming dark violet on warming On adding dilute H_2SO_4 , HCl , or HNO_3 to a solution of papaverine in acetic acid, the sulphate, hydrochloride, or nitrate is ppd A solution of papaverine in conc H_2SO_4 gives a pp of sulphate on adding water Papaverine is a weak base, its solutions do not affect litmus With ammonium selenite dissolved in conc H_2SO_4 it gives a bluish colour changing to red (Ferreira da Silva, *Bl* [3] 6, 87, *C R* 112, 126)

Reactions — 1. **Potash-fusion** yields methylamine, $\text{C}_6\text{H}_5\text{Me(OMe)}$ [13.4], and protocatechuic acid (Goldschmidt, *M* 4, 704, 6, 969) — 2 Yields

MeCl on heating with HClAq—3 Aqueous KMnO₄ oxidises it to papaveric, veratric, m-hempic, pyridine (a) tri carboxylic, dimethoxy-sinchoic, dimethoxy phthalic, and oxalic acids, hemipic iso imide NH₂, papaveraldine and CO₂. Papaveric acid (*v infra*) is the main product (Goldschmidt, *M* 6, 372, 8, 510)

Salts—B'HCl [220°] S 27 at 18° Large monoclinic crystals, $abc = 83.1 \times 58$, $\beta = 92^\circ 20'$ (Foullon, *M* 6, 675)—B'₂H₂PtCl₂ 2aq [198°]—B'₂H₂ZnCl₂ white dimetric crystals, $ac = 1.84$ —B'₂H₂Cl₂ZnI₂ plates (from alcohol)—B'₂H₂CdCl₂ [176°]—B'₂H₂Cl₂CdBr₂ [185°] (Jahoda, *M* 7, 506)—B'₂H₂Cl₂CdI₂ [180°]—B'₂H₂HgCl₂ triclinic prisms—B'HBz [214°] Monoclinic crystals, $abc = 83.1 \times 58$, $\beta = 92^\circ 60'$ —B'HI [200°] Monoclinic, sometimes isomorphous with the hydrochloride, but when crystallised from alcohol $abc = 18.1 \times 12.2$, $\beta = 91^\circ$ —B'HI, Purple monoclinic prisms—B'HI, thin reddish needles—B'₂H₂HgI₂—B'HNO₃ monoclinic tables, $abc = 82.1 \times 55$, $\beta = 94^\circ 9'$ —B'H₂SO₄ monoclinic prisms, $a, b, c = 83.1 \times 1.38$, $\beta = 92^\circ 29'$ —B'H₂Cr₂O₇ Flat orange needles—B'H₂C₂O₄ S 26 at 10° Prisms—B'C₂H₃N₃O₇ [179°] Golden tables (from alcohol)—Succinate B'₂C₂H₂O₄ [171°]—Benzoate B'C₂H₂O₄ [145°]—Salicylate B'C₂H₂O₄ [130°]—Meconate B'C₂H₂O₄ aq prisms, al sol alcohol—Ferrocyanide B'₂H₂FeC₆ (Plügge, *Ar Ph* [3] 25, 793)

Methylo-iodide B'MeI 4aq Melts at 195° when anhydrous 55°-60° when hydrated Insol ether, sol water and alcohol Moist Ag₂O forms a caustic hydroxide, capable of giving a carbonate (Claus, *J pr* [2] 38, 496, Goldschmidt, *M* 10, 673) The following compounds have been prepared from the methylo iodide (Stransky, *M* 9, 751) B'MeOH 2aq Melting at 215° when anhydrous, B'MeCl [75°], B'MePtCl₂ 3aq, B'MeCr₂O₇ [85°], and the picrate B'MeC₂H₂N₃O₇ [205°]

Ethyl bromide B'EtBr 4aq [140°-145°] (Goldschmidt, *M* 6, 667) or B'EtBr 4aq [111°] (Claus a Huethlin, *B* 18, 1576) Trimetric needles, $abc = 70.1 \times 64$ Boiling KOHAq forms the alkaline oxide (B'Et)₂O, al sol cold water

Ethyl-chloride B'EtCl 4aq prisms [80°] Gives B'Et₂PtCl₂ 3aq [223°]

Ethyl-iodide B'EtI [216°]

Ethyl-nitrate B'EtNO₃ 3aq Prisms

Ethyl chromate B'Et₂Cr₂O₇ [78°]

Ethyl picrate B'EtC₂H₂N₃O₇ [175°]

Benzyl chloride B'CH₂Cl 7aq [165°] KMnO₄ oxidises it to veratric acid, benzyl-papaveraldine, papaveraldine, and hemipic iso benzyl imide Aqueous (20 p c) KOH forms (B'C₂H₂)₂O [165°]—B'₂(C₂H₂)₂PtCl₂ crystalline pp—B'C₂H₂Cr₂O₇ [185°]—B'₂(C₂H₂)₂Cr₂O₇ [85°]

o-Nitro-benzyl-chloride

B'C₂H₂(NO₂)Cl Crystallises with 4aq, 6aq, and 9aq (Seutter, *M* 9, 859)—(B'C₂H₂(NO₂)Cl)₂PtCl₂ crystalline pp—B'C₂H₂(NO₂)NO₃ 1½aq prisms.—(B'C₂H₂(NO₂)₂)Cr₂O₇ yellow prisms—B'C₂H₂(NO₂)₂H₂N₃O₇ yellow prisms (from alcohol)

Phenacylo-bromide B'CH₂BzBr 2½aq Efflorescent pyramids (Seutter, *M* 9, 1035) Forms the derivatives: B'CH₂BzCl 6aq yellow

needles—(B'CH₂Bz)₂PtCl₂—B'CH₂BzNO₂ 2aq—(B'CH₂Bz)₂Cr₂O₇ and B'CH₂BzC₂H₂N₃O₇ [182°] An aqueous solution of the phenacylo-bromide gives with dilute NaOHAq a pp. B'CH₂BzOH, whence boiling alcohol produces (B'CH₂Bz)₂O [186°], crystallising in colourless needles

Tetrahydride C₂₀H₂₂NO₄ [201°] Made by reducing papaverine with tin and HCl (Goldschmidt, *M* 7, 497) Small prisms (from dilute alcohol), m sol hot water, al sol ether—B'HCl 3aq [290°] Monoclinic needles, with very bitter taste, causes albuminuria when injected internally—Salts B'₂H₂PtCl₂ 3aq—B'H₂SO₄ 7aq—B'H₂C₂O₄ 6aq—B'₂H₂Cr₂O₇—B'C₂H₂N₃O₇ [270°] Yellow needles, v al sol alcohol

Bromo-papaverine C₂₀H₂₂BrNO₄ [145°] Got by adding bromine-water to a solution of papaverine hydrochloride Monoclinic crystals, insol water, v al alcohol and ether—B'HBz

Nitro papaverine C₂₀H₂₂N₃O₇ aq [163°] S (ether) 03 at 12° Made by boiling papaverine with dilute nitric acid (S G 1 06) (Hesse, *A. Suppl* 8, 292) Pale yellow prisms (from dilute alcohol) Violently decomposes when quickly heated—Salts B'HCl 1½aq S 35 at 16°—B'₂H₂PtCl₂—B'HI v al sol hot water—B'HNO₃ aq—B'H₂SO₄ 8aq—B'H₂C₂O₄ 2aq yellow prisms, v al sol water

Papaveraldine C₂₀H₂₂NO₄ s.e. [43 1] C₂H₂(OMe)₂CO C₂H₂(OMe)₂N [210°] Made by oxidation of papaverine with KMnO₄ and dilute H₂SO₄ (Goldschmidt, *M* 6, 954, 7, 486) Yellowish crystalline powder (from alcohol), insol water and alkalis, v sol hot HOAc H₂SO₄ gives a red colour, changing to dark violet on warming Potash fusion splits it up into veratric acid and dimethoxyisouquinoline Tin and HCl reduce it to papaverine tetrahydride—B'HNO₃ 2aq lemon yellow needles—B'HCl 2aq yellow crystals, decomposed by water—B'₂H₂PtCl₂ aq orange prisms—B'H₂SO₄—B'C₂H₂N₃O₇ [209°]

Methylo-iodide B'MeI 3aq [135°]

Ethyl-bromide B'EtBr 3aq [above 270°]

Benzyl-hydroxide B'C₂H₂OH or its anhydride (B'C₂H₂)₂O is a product of the oxidation of papaverine benzyl chloride by dilute (2 p c) KMnO₄ at 45° (Goldschmidt, *M* 9, 327) It crystallises from boiling water or alcohol in slender colourless needles [154°], converted by boiling HClAq into papaveraldine.

Ozim C₂₀H₂₂NO₄(NOH) [245°]. White needles, sol hot benzene

Phenyl-hydrazide C₂₀H₂₂NO₄(N₂HPH) [81°] Reddish yellow nodules (from alcohol)

Papaveroline C₂₀H₂₂NO₄ s.e. [43 1] C₂H₂(OH)₂CH₂C₂H₂(OH)₂N Formed by boiling papaverine (3 pts) with HIAq (25 pts of S G 1 96) and red P for 10 hours (G, Krauss, *M* 11, 851) White crystalline powder (containing 2 aq), insol water, m sol alcohol, al sol ether, v sol acids. Yields (a)-methyl-isouquinoline on distillation with zinc-dust—Salts B'HCl white needles—B'H₂SO₄ 8½aq—B'HI 2aq—B'₂H₂C₂O₄ 3aq needles, m sol hot water

Papaveric acid C₂₀H₂₂NO₄ s.e. C₂H₂(OMe)₂CO C₂H₂N(CO₂H)₂ [233°] Formed by oxidising papaverine with KMnO₄ (Goldschmidt, *M* 6, 890, 10, 158, 691). Minute

right-angled tables (containing aq), al sol water, alcohol, and ether, m, sol dilute alcohol. Its solutions are acid in reaction. Decomposes on fusion into CO_2 and pyropapaveric acid $\text{C}_{17}\text{H}_{11}\text{NO}_4$ [280°]. Potash-fusion forms protocatechuic acid.

Salts— KHA' 2½ aq white leaflets, v e sol. water— KHA' 2 aq needles— CaA'' 1½ aq— BaA'' — $\text{Cu}_2\text{A}''$ (OH) 6 aq— $\text{Ag}_2\text{A}''$ 2½ aq— $\text{AgH}_2\text{A}''$ aq— $\text{H}_2\text{A}'$ HCl 2½ aq yellow needles.

Anhydride. [170°]. Got by boiling papaveric acid with Ac_2O (Goldschmidt, M 10, 159).

Mono-ethyl ether EtHA'' [188°]. Got by boiling the anhydride with alcohol. Converted by NH_4Aq into the amic acid.

Oxym $\text{C}_{17}\text{H}_{11}\text{NO}_4(\text{NOH})$ [e 156°]. Small needles (from alcohol) (Goldschmidt, M 10, 692).

Phenyl-hydrazide $\text{C}_{17}\text{H}_{11}\text{NO}_4(\text{N}_2\text{HPh})$ [190°]. Yellow needles (from dilute alcohol).

Nitro-papaveric acid $\text{C}_{17}\text{H}_{11}(\text{NO}_2)\text{NO}$ [215°]. Formed by dissolving papaveric acid in conc HNO_3 . Golden needles (containing aq)— $\text{Ag}_2\text{A}''$.

Pyropapaveric acid $\text{C}_{17}\text{H}_{11}\text{NO}_4$ [230°]. Formed by fusing papaveric acid. Minute white leaflets— CaA'' 4 aq groups of needles— BaA'' 4 aq— HA' HCl aq orange red needles— AgA' crystalline pp.

Oxym $\text{C}_{17}\text{H}_{11}\text{N}_2\text{O}_5$ [226°]. Needles (from alcohol)— B' HCl aq yellow needles.

Phenyl-hydrazide $\text{C}_{17}\text{H}_{11}\text{NO}_4(\text{N}_2\text{HPh})$ [228°]. Yellow prisms— B' HCl.

PAPAVEROSINE An alkaloid in dried poppy heads of *Papaver somniferum* (Deschamps, A Ch [4] 1, 453). Nacreous needles (from alcohol). Coloured red by H_2SO_4 .

PAPAYOTIN An amorphous hygroscopic powder ppd by adding alcohol to an aqueous extract of the juice of *Carica papaya* (Peckholt, Ph [3] 10, 343). It has an astringent, slightly sweet taste.

PARABANIC ACID $\text{C}_6\text{H}_7\text{N}_2\text{O}_4$ + e

$\text{CO} < \text{NH CO}$ Oxalyl urea Mol w 114 S
 $\text{CO} < \text{NH CO}$ 47 at 8° H C 212,700 H F 2,200 (Matignon, C R 113, 198).

Formation—1 By dissolving uric acid (1 pt) in nitric acid (7 pts of S.G. 1.3) at 70° and evaporating to a syrup, when parabanic acid crystallises out on cooling (Liebig & Wöhler, A 26, 285, Menshutkin, A 172, 74)—2 By oxidation of alloxan—3 A product of the action of HOCl on guanine (Strecker, A 118, 156)—4 From uric acid by heating with MnO_2 and dilute H_2SO_4 (Wheeler, Bl [2] 7, 521) or with KNO_3 and H_2SO_4 (Gibbs, B 1, 341)—5 A product of the action of HCl and KClO_4 on uric acid (Laurent & Gerhardt, A Ch [3] 24, 175)—6 By the action of POCl_3 on oxaluric acid $\text{NH}_2\text{CO NH CO CO}_2\text{H}$ at 200° (Grimaux, C R 77, 1548)—7 By the action of PCl_5 on a mixture of oxalic acid and urea (Ponomareff, Bl [2] 18, 97)—8 By heating nitro-pyruvic ureide with bromine and water (Grimaux, C R 79, 1478)—9 By the action of Br and water on uric acid (Hardy, Bl [2] 1, 445, Magner, Bl [2] 22, 56).

Properties.—Monoclinic laminae (from water or alcohol); $a b c = 1.665, 478, \beta = 81^\circ 89'$. Decomposed above 200°, forming a white sublimate. Completely oxidised by KMnO_4 and H_2SO_4 to CO_2 and NH_3 . Not attacked by HNO_3 in the cold. Its solution is not ppd by Ca salts,

but on boiling with alkalis it is split up into oxalic acid and ammonia. In the same way lead salts and a little NH_3 gave a pp of lead oxalate (Maly, M 2, 264).

Reactions—1 In aqueous solution the salts rapidly change to oxalates—2 Zinc and HCl reduce it to oxalantin—3 Alcoholic NH_3 at 100° forms oxaluramide—4 Alcoholic EtI at 100° forms $\text{C}_6\text{H}_7\text{NO}_4\text{I}$, crystallising from alcohol in green prisms, nearly insol cold water (Hasiwetz, A 103, 200)—5 Urea (1 pt) at 180° forms $\text{C}_6\text{H}_7\text{N}_3\text{O}_4$, a sparingly-soluble powder (Grimaux, Bl [2] 32, 120).

Salts— $\text{NH}_4\text{A}'$ Crystalline powder, got by adding alcoholic ammonia to a solution of parabanic acid in absolute alcohol. Gives off its NH_3 at 100°. Warm water converts it into ammonium oxalurate— NaA' — KA' Got by using alcoholic KOEt— AgA' Crystalline pp, got by adding AgNO_3 to a solution of KA' — $\text{Ag}_2\text{O}_2\text{N}_2\text{O}_4$ aq Got by ppg a solution of parabanic acid with AgNO_3 . Insol water, sol HNO_3 —Urea salt $\text{CON}_2\text{H}_4\text{HA}'$ Four sided tables, al sol cold water—Phenyl-hydrazine salt $(\text{N}_2\text{H}_5\text{Ph})_2\text{A}'$ aq [170°]. Insol alcohol and ether. Boiling water converts it into $\text{NH}_2\text{CO NH CO CO N}_2\text{H}_5\text{Ph}$ [215°] (Skinner & Ruhemann, O J 53, 550)—Aniline salt [250°] (S A R).

Hydrate $\text{C}_6\text{H}_7\text{N}_2\text{O}_4$ aq S 135 at 8° Formed from uric acid (1 pt) and nitric acid (3 pts of S.G. 1.3) at 60°. The mixture is kept at 35°–55°, and finally heated to 70°, when the hydrate separates in large crystals (Tollens & Wagner, A 166, 321, 175, 227). Formed also by the action of POCl_3 on a mixture of urea and oxalic acid, heat not being applied (P). At 150°–160° it loses water and changes to ordinary parabanic acid.

Methyl parabanic acid $\text{CO} < \text{NMe CO}$
 NH CO

Methyl-oxalyl urea Oxalyl-methyl urea [149°]

Formation—1 By heating nitroso creatinin with HCl at 100° (Dessaignes, A 97, 342, Marcker, A 133, 315)—2 From methyl uric acid and HNO_3 (Hill, B 9, 1093, 13, 739)—3 By the action of AgNO_3 on methyl thioparabanic acid (oxalyl methyl thio urea) (Andreasch, B 14, 1449, M 2, 279)—4 By oxidation of theobromine or caffeine with chromic acid mixture (E Fischer, A 215, 297, Maly & Hinteregger, B 14, 727, M 2, 94).

Properties—Colourless trimetric prisms, sol hot water. Acid in reaction. Not volatile with steam. Decomposed by alkalis, even in the cold, into methyl-urea and oxalic acid.

Salt— AgA' Needles, sol hot water.

Di methyl parabanic acid $\text{CO} < \text{NMe CO}$
 NMe CO

Cholestrophane [146°]. (276°) S 187 at 20°

Formation—1 The final product of the action of chlorine on caffeine in water (Rochleder, A 73, 123)—2 By boiling caffeine with nitric acid (Stenhouse, A 45, 371, 46, 229)—3 By heating dry silver parabanate with MeI at 100° (Strecker, A 118, 174)—4 By oxidation of caffeine with chromic acid mixture (Maly & Hinteregger, B 14, 723)—5 By the action of AgNO_3 on $\text{CS} < \text{NMe CO}$ (Andreasch, M 2, 288).

6. By the action of dilute HClAq on murexolin

(Brunn, *B*, 21, 515) —7. By passing ozonised oxygen through water in which caffeine is suspended (Leipen, *M* 10, 184)

Properties —Trimetric prisms (from alcohol). May be sublimed. Completely decomposed by alkalis. Gives no pp with lead acetate until ammonia is added, when lead oxalate is ppt. Pure HNO_3 has no action (Franchumont, *R. T. C.* 6, 217)

Reactions —1 Alcoholic NH_3 forms dimethyl oxaluramide [225°] on heating —2 HClAq at 200° splits it up into oxalic acid, CO_2 , and methylamine (Calm, *B* 12, 624) —3 Cold alcoholic soda decomposes it into oxalic acid and di-methyl urea ($\text{M} + \text{H}_2$) —4 Zinc and dilute H_2SO_4 reduce it to di-methyl-glyoxyl urea $\text{CO} < \text{NMe} \cdot \text{CH} \cdot \text{OH}$, a crystalline body, melting below 100° decomposed by heating with baryta into glycolic acid, oxalic acid, methylamine, and CO_2 (Andreasch, *M* 3, 436) —5 On warming with water containing BaCO_3 it yields di-methyl-oxamide (Maly a Hinteregger, *M* 2, 88, 132)

Phenyl-parabanic acid $\text{CO} < \text{NH} \cdot \text{CO} < \text{NPh} \cdot \text{CO}$ [208°]

Made from phenyl-urea and $\text{ClCO} \cdot \text{CO}_2\text{Et}$ (Stojentun, *J. pr.* [2] 82, 11) Silky plates, v sol alcohol, ether, and hot water

Di phenyl-parabanic acid $\text{CO} < \text{NPh} \cdot \text{CO} < \text{NPh} \cdot \text{CO}$

[204°] Formed by boiling an alcoholic solution of diphenyl guanidine dicyanide (dicyanomel-aniline) $\text{C}_{12}\text{H}_{11}\text{N}_3$, or (a) tri phenyl-guanidine dicyanide with HClAq (Hofmann, *Pr* 11, 275, *B* 3, 764) Formed also by warming thiocarb-anilido thio oxanilide with alcoholic AgNO_3 , and by the action of $\text{ClCO} \cdot \text{CO}_2\text{Et}$ on di phenyl urea (Stojentun) Needles, insol water, v sol alcohol and ether. Decomposed by boiling KOH aq into oxalic acid, aniline, and CO_2 . Yields with fuming HNO_3 a di nitro di phenyl parabanic acid crystallising in needles

PARACONIC ACID v *Lactone of Oxytyro-BARTARIC ACID*

PARAFFIN Solid, wax like, fatty mixture of hydrocarbons of the $\text{C}_n\text{H}_{2n+2}$ series, very rich in carbon and characterised by a very marked indifference to most reagents

Analysis —

	1	2	3	4	5
C	85.15	84.9	85.81	84.86	85.5
H	14.85	14.31	14.44	15.02	14.9

1 Anderson, *J* 1857, 480, 2, 3 Brodie, *P. M.* [3] 83, 178, 4 Pawlewski, *B* 23, 327, 5 Gill a Meusel, *C. S. J.* 6, 466 Beilstein states (*Bn* 1, 189) that commercial paraffin usually contains 1 p.c oxygen, which can be removed by heating with Na in a tube Lippmann a Hawliczek (*B* 12, 69), from a sample of brown coal paraffin v sol alcohol [37°] by heating with PCl_5 , formed $\text{C}_{25}\text{H}_{52}\text{O}_4$, hence give formula $\text{C}_{25}\text{H}_{52}$. Bolley (*A* 106, 230), by heating hot paraffin with Cl_2 formed $\text{C}_{25}\text{H}_{52}\text{O}_4$ — $\text{C}_{25}\text{H}_{52}\text{Cl}_{11}$, hence gives formula $\text{C}_{25}\text{H}_{52}$. Gill a Meusel (*Z* 1869, 65) have formed from paraffin by oxidation with CrO_3 , cerotic acid $\text{C}_{25}\text{H}_{52}\text{O}_4$, hence formula $\text{C}_{25}\text{H}_{52}$. Pawlewski, by Reoult's method, using glacial acetic acid, finds $\text{C}_{25}\text{H}_{52}$ — $\text{C}_{25}\text{H}_{52}$, by dilute benzene: and *p*-xylene $\text{C}_{10}\text{H}_{12}$ — $\text{C}_{10}\text{H}_{12}$, and with saturated solutions in benzene and *p*-xylene

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($\text{C}_{25}\text{H}_{52}$)₂ — ($\text{C}_{25}\text{H}_{52}$)₃. It behaves as a colloid in most other solvents

Melting-point [45°–85°] By the action of solvents, small quantities of fractions [38°–45°] have also been obtained (Albrecht, *D. P. J.* 218, 280) Ozokerite paraffin [64°]

Boiling-point Above 300° In crude petroleum the portion (300°–400°) solidifies on cooling. On fractionating, Bolley and Tuschschmid found that the portion [53°] (300°) gave the following fractions (150°) [43°], (200°) [44.5°] and residue [53.5°]

Specific gravity varies from 0.872–0.912 at 17° (Albrecht) Ozokerite paraffin [64°] S.G. 22 0.917 [38°] S.G. 24 0.874 (Beilby, *C. J.* 43, 388) Paraffins expand considerably on melting

Occurrence —Occurs native as fossil wax, hatchettine, and ozokerite. The latter is the most abundant, and is found in Galicia, Roumania, on the island of Tschelen on the east coast of the Caspian Sea, and on the island of Swjatoi at Baku. When bleached it is called ceresine. According to Zawiecki, ozokerite contains a crystalline and amorphous paraffin. It is also found in Etna lava to the extent of 43 p.c in certain geodes (Silvestri, *G* 12, 9) It is obtained by the distillation of brown coal, turf, boghead coal, shales, schists, and natural bitumens, wood, wax, and wax and lime. It can also be obtained from brown coal tar. Details of various methods are given by von Boyen, *Z. f. Angew. Chem.* 1891, 261. Paraffin is also obtained in large quantities from American petroleum, being present in the high boiling portions which form a residue after the lighting oils have been distilled off. Other mineral oils contain appreciable quantities of solid paraffin, e.g. Rangoon oil of Burma, 6 p.c. For the preparation of commercial paraffin from these bodies v Thorpe's DICTIONARY OF APPLIED CHEMISTRY, and for a theory for its formation in nature by the decomposition of animal matter, vide Zawiecki, *D. P. J.* 280, 69, 85, and 133

Properties —Paraffin, when pure, is a solid, colourless, translucent substance, perfectly in odorous and tasteless, somewhat resembling spermaceti. It readily melts, forming a colourless oil, burns from a wick with a bright flame, but does not burn easily in the mass. It is insol water, sol hot alcohol, v sol ether and oils. The solubility of paraffin from ozokerite has been studied by Pawlewski a Fillmonowicz (*B* 21, 2973), who show that the liquid constituents are mostly sol glacial acetic acid, whereas vaseline, ceresine, ozokerite and paraffin are almost insoluble. Pawlewski (*B* 23, 327) also states that ozokerite paraffin is sol formic acid, acetic acid, benzene, *p*-xylene, and chloroform. Thorpe a Young (*A* 165, 1), by heating paraffin in closed vessels at a high temperature, have resolved it into a little gas and hydrocarbons of the $\text{C}_n\text{H}_{2n+2}$ series (C_2H_6 – $\text{C}_{12}\text{H}_{26}$), and others of the $\text{C}_n\text{H}_{2n+2}$ series (C_2H_6 – $\text{C}_{12}\text{H}_{26}$). Higher solid and liquid hydrocarbons were also obtained. Long heating at 150° causes an increase of weight, attributed by Bolley and Tuschschmid to oxygen absorption (*Z* 1868, 500, Jaznowitch, *B* 8, 768). Oxidised products can also be obtained by the action of oxidising agents. Champion (*C. R.* 75, 1576) has shown that nitrosulphuric acid slowly transforms it at

S F

90° into a liquid oil of the composition $C_{11}H_{22}NO_2$ (called paraffinic acid), from which ethereal salts have been obtained, and at the same time yields a white solid $C_{11}H_{22}NO_2$ sol Aq. Fuming nitric acid, according to Fouchet, also acts upon paraffin at 110°, forming a true paraffinic acid $C_{11}H_{21}O_2$ [46°] together with other fatty acids. The acid has a wax-like odour, is insol water, but sol alcohol, ether, and benzene, is easily decomposed by heat, and forms deliquescent salts of the alkaline metals, and yellow, cheesy plates with the earths and magnesia. It is carbonised by sulphuric acid (*Bl* 23, 111, *C R* 79, 320) and nitric acid converts it into nitro-compounds and suberic acid. Champion has also shown that chlorine is absorbed by paraffin in sunlight, producing hydrochloric acid. According to Gill & Meusel (*Z* 1869 65), CrO_3 and dilute HNO_3 oxidise paraffin to a mixture of cerotic $C_{26}H_{54}O_2$, succinic, and other fatty acids. According to Beilstein & Wiegand (*B* 16, 1548), ozokerite contains a solid hydrocarbon of the olefine series. It is formed by distilling the ozokerite *in vacuo*, and, after removing the first portions of only distillate, crystallising the solid portion from a solution of alcoholic benzene. It is called *tekene* [79°] *S G* 0 9392, and is a very stable compound, being unacted upon by CrO_3 , HNO_3 , and $KMnO_4$. It is, however, completely oxidised by acid $KMnO_4$, and slowly forms a bromo-compound when heated with Br and H_2O in a sealed tube. For methods of analysis of paraffin scale, and determination of impurities therein, *vide* B Redwood, *S C I* 3, 430, *Journ Soc Arts*, 1886 56, Sutherland, *S C I* 6, 123, 271, and Stuart Thomson, *ibid* 10, 342 *et seq*. S R

PARAFFIN HYDROCARBONS *v* HYDROCARBONS

PARAFFIN OIL *v* PETROLEUM

PARAGLOBIN *v* PROTEIDS

PARAHEMOGLOBIN *v* HEMOGLOBIN

PARALBUMIN *v* PROTEIDS, *Appendix C*

PARALDEHYDE *v* ALDEHYDE

PARANILINE $C_6H_4N_2$ [192°] A base found by Hofmann (*Pr* 12, 314) in the preparation of crude aniline on a large scale. Long, silky needles (from dilute alcohol) EtI yields $C_{12}H_{11}EtN$, and $C_{12}H_{11}EtN_2$. — $B''H_2Cl$ aq. six-sided plates (from conc. HCl aq.) converted by water into yellow needles of $B''HCl$ aq. sl sol water. — $B''H_2PtCl_6$ yellow prisms. — $B''HNO_3$, — $B''H_2SO_4$, — $B''H_2SO_4$.

Benzoyl derivative $C_{12}H_{11}BzN_2$ Needles

PARAPEPTONE *v* PROTEIDS

PARELLIC ACID $C_8H_8O_4$? An acid sometimes obtained in the preparation of lecanoric acid (Schunck, *A* 54, 274). Needles (containing aq.) v sl sol cold water, sol alcohol and ether. Tastes bitter. Reddens litmus. — PbA' ? white flocculent pp.

PARICINE *v* CINCHONA BASES

PARIDIN $C_{14}H_{24}O_{14}$ *S* 15, *S* (alcohol) 2. A neutral glucoside extracted by alcohol from the leaves of *Paris quadrifolia* (Walz, *Pharm Cent* 1841 690, *N Jahres Pharm* 13, 174, Delius, *ibid* 60, 25). Formed also, together with a sugar, by boiling paristypnin with dilute H_2SO_4 . Thin laminae (containing 4 aq.) (from water) or tufts of needles (from alcohol). Conc. H_2SO_4 turns it red. Boiling HCl aq. decomposes

it, in alcoholic solution, into a sugar and resinous paridol $C_{14}H_{24}O_{14}$.

PARILLIN $C_{16}H_{26}O_{16}$ or $C_{16}H_{26}O_{16}$ [210°]. *S* 008 in the cold, 5 at 100°. Ppd. by adding water to an alcoholic extract of sarsaparilla root (Flückiger, *Ph* [3] 8, 489). Plates or prisms, sol hot alcohol and chloroform. Conc. H_2SO_4 gives a yellow solution. Dilute HCl gives green fluorescence and splits it up into a sugar and parigenin $C_{16}H_{26}O_{16}$, which is insol boiling water.

PARISTYPHININ An amorphous body which accompanies paridin, and may be ppd. by tannin. Boiling dilute acids split it up into a sugar and paridin.

PARPEVOLINE Name given to the hexahydrate of di methyl ethyl pyridine, and to pyridine bases isomeric therewith.

PARSLEY The volatile oil of parsley contains a terpene (162° i v), *S G* 12 865 (Gerichten, *B* 9, 259, Sauer & Grunling, *A* 208, 75). The seeds contain apiol (*q v*). Water extracts *APIIN* (*q v*) from the plant.

PARSNEP The volatile oil of parsnep seeds contains octyl *n*-butyrate (Renesse, *A* 166, 80).

PARVOLINE $C_8H_{11}N$ (c 200°) A product of the putrefaction of horse flesh (Gautier, *Bl* [2] 48, 11). Oil, smelling like hawthorn blossom, v sol alcohol and ether. Resinifies in air. Its platinum chloride forms sparingly soluble flesh coloured crystals.

Parvoline $C_8H_{11}N$ (- 220°) A product of the distillation of cinchonine with potash (Oechsner de Coninck, *C R* 91, 296). — $B''H_2PtCl_6$ brownish yellow powder.

Parvoline $C_8H_{11}N$ (188°) A product of the dry distillation of the bituminous shale of Dorsetshire (Greville Williams, *C J* 7, 97).

Parvoline $C_8H_{11}N$ (188°) *S G* 21 986. Got by distillation of coal (Thenius, *J* 1861 502).

Parvolines of known constitution *v* DIETHYL-PYRIDINE, DI METHYL-ETHYL PYRIDINE, TETRA METHYL PYRIDINE, and METHYL PROPYL-PYRIDINE.

PASSIVE STATE OF METALS *v* Passivity of iron, art IRON, p 52.

PATCHOULI The volatile oil of patchouli, obtained from the leaves of *Pogostemon Patchouli*, contains a sesquiterpene (which yields $C_{15}H_{24} \cdot 2HCl$ [118°]), and a camphor $C_{15}H_{24}O$ [55°], (296°), *S G* 12 1051 (Gal, *C R* 68, 406, Montgolfier, *C R* 84, 88, *B* 10, 234, Wallach, *A* 238, 81). Patchouli camphor crystallises in hexagonal prisms, insol water, v sol alcohol and ether. It is levorotatory $[\alpha]_D = -118^\circ$. On distillation with $ZnCl_2$, or on heating with $HOAc$ and Ac_2O , it yields patchoulene $C_{15}H_{24}$ (254°), *S G* 2 946, $[\alpha]_D = -42^\circ$.

PATELLARIC ACID $C_{11}H_{20}O_{10}$ [above 100°] Occurs in the lichen *Patellaria* (or *Parmelia scruposa*, from which it can be extracted by ether (Knop, Weigelt, *Z* [2] 5, 298). Crystalline mass, insol water, v sol alcohol and ether. Tastes bitter. Decomposed on fusion or on boiling with baryta, with formation of orcin and oxalic acid. $FeCl_3$ gives a purple colour. Cold baryta water forms a salt with transient blue colour.

PAVIN *v* FRAXIN.

PAYTINE $C_{21}H_{21}N_2O$. [156°]. $[\alpha]_D = -49.5^\circ$ in a 45 p.c. alcoholic solution. An alkaloid extracted by alcohol from a white bark of an *Aspidosperma* from Payta in Peru (Hesse, *A* 154, 287, 166, 259, 211, 280, Wulfsberg, *Ph* [R] 11, 269, Arata, *G* 11, 246, *O* J 40, 622) Prisms (containing aq) sl sol water, sol ether, benzene, chloroform, and (unlike aspidospermine) lignin. Tastes bitter, is not poisonous. When heated with soda-lime it yields paytone, a non nitrogenous substance. Conc HNO_3 forms a colourless solution, changing through red to yellow. $FeCl_3$ and conc H_2SO_4 give no colour. Chloride of gold gives a purple pp. $HgCl_2$ gives a yellow amorphous pp. Bleaching powder produces, in an acid solution, a red colour changing through blue to yellow.

Salts— $BHCl$ S 6 at 15° . Prisms— $B \cdot H_2PtCl_6$, dark-yellow amorphous pp., dissolves in hot $HClAq$, forming a red solution changing to blue and depositing a blue pp.

Paytamine $C_{21}H_{21}N_2O$. An amorphous alkaloid accompanying paytine. Unlike paytine, KI does not ppt it from neutral solutions. Its salts are amorphous, and it does not yield paytone.

PECTOLACTIC ACID $C_8H_9O_6$. An acid got by boiling milk sugar with $NaOH$ aq and less CuO than is needful for complete oxidation (Boedeker, *A* 100, 281). Brownish syrup, drying up to a varnish (containing 2 aq). Miscible with water and alcohol, insol ether. Reduces ammoniacal $AgNO_3$, forming a mirror— BaA^{44} aq powder, insol alcohol.

PECTOUS SUBSTANCES. Unripe fleshy fruits and fleshy roots (e.g. pears, carrots, and turnips) contain a substance (pectose) insoluble in water, alcohol, and ether, transformed under the influence of acids into pectin, which is soluble in water, and is ppd as a jelly on adding alcohol or on boiling (Vauquelin, *A* Ch 5, 100, 6, 282, [2] 41, 46, Braconnot, *A* Ch [2] 28, 173, 30, 96, 47, 266, 72, 433, Guibourt, *J Chim Med* 1, 27, Mulder, *J pr* 16, 277, 16, 246, Fremy, *A* Ch [3] 24, 9, Soubeiran, *J Ph* [3] 11, 417, Poumarède a. Figuer, *J Ph* [3] 11, 458, 12, 81, Chodoff, *A* 51, 356, Scheibler, *Z* [2] 4, 433, Reichardt, *Ar Ph* [3] 10, 116, Giraud, *C R* 80, 477, Stude, *A* 131, 244, Rochleder, *Z* [2] 4, 381).

Pectin, obtained from the juice of ripe pears and apples by removing albuminoids by tannin and calcium by oxalic acid, and then ppg with alcohol, is, according to Bauer (*J pr* [2] 30, 370), a mixture of carbohydrates. Different analyses of pectin give C 39.5 to 46 p.c., H 5.5 p.c., O 48.5–55 p.c. Pectous substances differ from cellulose in being stained by methylene blue, but, unlike lignin, the colour is destroyed by acids (Mangun, *C R* 109, 579). By boiling with water pectin is converted into 'parapectin,' a solution of which, unlike pectin, gives a pp with lead acetate. Nitric acid yields mucic acid by oxidation. Boiling dilute acids convert pectin into 'metapectin,' which is acid in reaction and gives a pp with $BaCl_2$ (unlike pectin and parapectin). Boiling alkalis convert pectin into pectic acid.

Pectic acid $C_8H_7O_6$? (Fremy) C 41.4 to 44.9 p.c., H 4.71 to 5.4, O 49.7 to 53.8. Got by boiling an aqueous extract of carrots with Na_2CO_3 and ppg with HCl . Produced also from

pectin in fruit by the action of a ferment (pectase). Jelly, drying up to a horny mass. Acid in taste and reaction, insol. cold water, alcohol, and ether. Converted by boiling water, first into parapectic acid and then into metapectic acid. Boiling dilute acids yield arabic acid.

Salts— $Na_2C_8H_7O_6$, $PbC_8H_7O_6$, $Ag_2C_8H_7O_6$ (Chodoff)— $Ag_2C_8H_7O_6$ (Fremy). Arabic acid, when prepared by hydrolysis of pectic acid, is called metapectic acid, and the arabinose got by the action of $HClAq$ upon it is called pectinose. Pectinose or arabinose is tetra oxy-valeric aldehyde (q.v.).

References—ARABIN and CELLULOSE.

PELARGONIC ACID v. ENNOIC ACID.

PELLETIERINE $C_8H_{17}NO$ (195°). S.G. 2 988 V D 4 88 (calc 4 66) S. 5 Occurs, together with methyl pelletierine, pseudo pelletierine, and isopelletierine in the bark of the pomegranate (Tanret, *C R* 86, 1270, 87, 358, 88, 716, 90, 695, *J Ph* [4] 28, 168, 384). It may be extracted from the powdered bark by shaking with chloroform and milk of lime. If the chloroform is decanted and shaken with a dilute acid, the alkaloids will be taken up by the acid solution. The acid solution is mixed with $NaOH$ and shaken with chloroform, which extracts pelletierine and pseudo pelletierine, which may be separated by crystallisation of their sulphates, the pelletierine is finally distilled in a current of H .

Properties—Colourless liquid, partially decomposed at 195° , but may be distilled *in vacuo*. M sol water, miscible with alcohol, ether, and $CHCl_3$. The sulphate is levorotatory; $[\alpha]_D = -30^\circ$. Resinified by absorption of oxygen. Fumes with HCl .

Methyl pelletierine $C_8H_{17}NO$ (215°) S 4 at 12° . Extracted from an acid solution of the mixed bases by adding $NaHCO_3$ and shaking with chloroform. Liquid, forming very deliquescent salts. V sol alcohol, ether, and chloroform. Its hydrochloride is dextrorotatory $[\alpha]_D = +22^\circ$.

Pseudopelletierine $C_8H_{17}NO$ [46°] (when hydrated) (246°) S 40 at 0° S (ether) 11 at 10° . Crystallises from water in prisms (containing 2 aq), v e sol water, alcohol, and $CHCl_3$. Inactive as regards light. Powerful base, expelling NH_3 from its salts. Gives the alkaloidal reactions. Like pelletierine, it gives an intense green colour with H_2SO_4 and $K_2Cr_2O_7$ — $BHCl$ rhombohedra S 100 at 10° — $B \cdot H_2PtCl_6$ needles— $B \cdot H_2SO_4$ aq v sol water.

Isopelletierine $C_8H_{17}NO$. An inactive liquid, resembling pelletierine in other respects.

PELOSINE $C_{11}H_{21}NO$, 1 aq. An alkaloid contained in the root of *Parera brava* or *Cusampelos Parera* (Wiggers, *A* 27, 29, 33, 81, Bödker, *A* 69, 53, Flückiger, *N J P* 31, 257). Extracted by dilute H_2SO_4 , and ppg by Na_2CO_3 , and dissolved in ether or CS_2 . Amorphous powder, almost insol water. Strongly alkaline in reaction. Yields methylamine, dimethylamine, and a derivative of pyrrole on distilling with KOH (Williams, *Chem Gaz* 1858, 331)— $BHCl$ aq amorphous hygroscopic powder— $B \cdot H_2PtCl_6$ amorphous— $B \cdot H_2CrO_4$ aq. According to Flückiger, pelosine is identical with buxine and bebeerine.

Pellatella $C_{15}H_{11}NO$? A yellow base, insol. ether, either formed from pelosine by atmospheric oxidation or occurring in *Parera* — $B_2H_2PtCl_6$.

PENDECANAPHTHENE or **Pentadecanaphthene** $C_{15}H_{10}$ (247° cor) SG 11 829 Occurs in Russian petroleum (Markownikoff, *J R* 15, 339) Liquid

PENNYROYAL The oil of pennyroyal from *Mentha pulegium* contains pulegone, which may be purified by fractional distillation *in vacuo* (Beckmann, *A* 262, 3, cf Kana, *A* 32, 286)

Pulegone $C_{10}H_{16}O$ (181° at 60 mm) SG 22 982 $[\alpha]_D = +22.9^\circ$ Somewhat viscous oil, smelling like peppermint Darkens in colour even in closed tubes When reduced by Na it gives 40 p.c. of menthol — $C_{10}H_{17}BrO$ [40.5°] $[\alpha]_D = -33.8^\circ$ Made by passing HBr into pulegone dissolved in ligroin Whetstone shaped crystals v sol alcohol and ether This hydrobromide is converted by hydroxylamine into an oxim [38°] crystallising in four sided plates, which on standing lose Br and then melt at 110° This oxim also forms the hydrated oxim [157°] (*v infra*) by taking up water The hydrobromide is re-converted into pulegone by treatment with Ag_2O , while zinc-dust reduces it to pulegone hydride ($\alpha = -17.4^\circ$), which yields an oxim [83°], and can be converted, by the action of Na on its ethereal solution, into levorotatory menthol $[\alpha]_D = -21.3^\circ$, giving a benzoyl derivative [54°]

Hydrated oxim $C_{10}H_{18}O(NO H)$ [157°] $[\alpha]_D = -83.4^\circ$ in alcoholic solution Formed by the action of hydroxylamine on pulegone Long matted needles, sl sol ether, cold alcohol, and benzene Yields an acetyl derivative $C_{10}H_{17}AcNO_2$ [149°] and a benzoyl derivative [138°] — B^*HCl [118°] $[\alpha]_D = -32.4^\circ$ Trimeric crystals, $a b c = 605.1 \times 1048 - B^*HBr$ [111°]

Pulegonamine $C_{10}H_{15}NO$ Made by reducing the foregoing oxim with HI and P Amber coloured liquid, sl sol water, v sol alcohol and ether Tastes bitter Boiling Mel forms $C_{10}H_{15}MeNO$ Phenyl thiocarbimide forms $PhNHCSN(C_{10}H_{15}O)$ [198°] Yields a benzoyl derivative $C_{10}H_{15}BzNO$ [101°] — $(C_{10}H_{15}NO)HCl$ [117°] Long needles, v sol water

PENTADECANE $C_{15}H_{32}$ [10°] (270°) (136° at 10 mm) SG $\frac{1}{4}$ 772, $\frac{2}{4}$ 769, $\frac{12}{4}$ 718 Made by reduction of $C_{15}H_{22}CO_2$, CH_3 or of pentadecanoic acid with HI and P (Krafft, *B* 15, 1700) A pentadecane (114° at 15 mm) is got by distilling barium palmitate with NaOMe (Mal, *B* 22, 2134)

Pentadecane $C_{15}H_{32}$ (260°) SG 12 825 Occurs in American petroleum It yields decane when passed through red-hot tubes (Pelouze & Cahours, *Vohl*, *J* 1865, 841)

PENTADECENOIC ACID $C_{15}H_{28}O_2$? *D-ismylovaleric acid* (300°-308°) A product of the action of CO at 160° on a mixture of sodium isovalerate and NaOOC H_{11} (Geuther & Fröhlich, *A* 302, 304) Liquid

PENTADECENYL ALCOHOL? $C_{15}H_{32}O$? [78°] The portion of the wax of *Ficus gumiflua* that is readily soluble in ether (Kessel, *B* 11, 2114). Nodules (from ether)

PENTADECINENE $C_{15}H_{26}$ (230°-240°) Got by heating the acetyl derivative of di-oxy-penta-

decane (triamylene acetate) with potash (Bauer, *A* 137, 249) Thick liquid. Unites with bromine.

PENTADECOIC ACID $C_{15}H_{30}O_2$ [62°] Formed by oxidation of coccoeryl alcohol $C_{15}H_{30}(OH)_2$ or of coccoeric acid $C_{15}H_{28}O_2$ with CrO_3 and acetic acid, the yield is about 40 p.c. (Liebemann & Bergham, *B* 20, 964) Crystalline solid V sol alcohol, ether, acetic acid, and benzene, less readily in ligroin — A_2Ca and A_2Ba granular pps

Methyl ether A^*Me . [68°], crystalline solid

Pentadecic acid $C_{15}H_{30}O_2$ [51°] (257° at 100 mm) Made by oxidation of methyl penta decyl ketone with chromic acid mixture (Krafft, *B* 12, 1671, 19, 2993) Pearly scales — BaA^* — AgA^*

Pentadecic acid $C_{15}H_{30}O_2$ [70°] Occurs, together with mannite, in the fungus *Agaricus integer* (Thörner, *B* 12, 1636) Small needles, v e sol ether and boiling alcohol, insol water

Isomerides v CERIC ACID and LACTARIO ACID
PENTADECONENE $C_{15}H_{28}$ (220°) Made from $C_{15}H_{28}Br$, and alcoholic potash (Bauer, *A* 147, 255)

Pentadecanone (235°-245°) Made from santonic acid and HI (Cannizzaro & Amato, *B* 7, 1104)

DI PENTADECYL-CARBINOL $(C_{15}H_{31})_2CH(OH)$ [85°] Made by reducing palmitone in boiling alcohol by adding sodium (Kipping, *C J* 57, 987) Silky needles or plates (from alcohol), sol ether, insol water With boiling Ac_2O it yields $(C_{15}H_{31})_2CH O^*c$ [49°], a granular powder, v sol ether

DI-PENTADECYL KETONE v PALMITONE

n-PENTANE C_5H_{12} $\frac{1}{4}$ 6
 $CH_3CH_2CH_2CH_2CH_3$, Mol w 72 (87°) SG $\frac{127}{4}$ 6282 (Schiff), $\frac{11}{4}$ 6337, $\frac{33}{4}$ 6250 (Perkin), μ_2 1.3607 at 65°, $\mu_D = 1.3649$, $\mu_R = 1.3769$ (Gladstone, *C J* 59, 290) M M 5.638 at 17.2° Occurs in American and in Galician petroleum, in coal tar oil, and in resin oil (Greville Williams, *C J* 15, 130, Schorlemmer, *C J* 25, 1085, *Pr* 15, 131, Lachovitch, *A* 220, 190, Renard, *A Ch* [6] 1, 225) Oil. Yields on chlorination two amyl chlorides (106°) and (104°), and finally $C_5H_9Cl_3$ (230°-240°) (Bauer, *C R* 51, 572, Schorlemmer, Lachovitch) CrO_3Cl , followed by water, yields CH_3Cl CO C_4H_9 (Etard) At a red heat it is decomposed with formation of ethylene, propylene, and butylene (Norton & Andrews, *Am* 8, 1)

Isopentane $(CH_3)_2CHCH_2CH_3$, *Tri-methyl-ethane* VD 250 (calc 248) (30°) SG $\frac{11}{4}$ 6248, $\frac{33}{4}$ 6159 (Perkin), $\frac{50}{4}$ 6132 (Schiff, *A* 220, 87), $\frac{11}{4}$ 6375 (Just, *A* 220, 153) SV 117.2 (Schiff) M M 5.750 at 16.4° Occurs in American and in Galician petroleum (Warren, *Z* 1865, 668, Lachovitch) Formed by the action of water on zinc isoamyl, or by heating isoamyl iodide with zinc and water at 140° (Frankland, *A* 74, 47) It is also a product of the action of $ZnCl_2$ on hot isoamyl alcohol, and of zinc on dry isoamyl iodide

The same pentane can be obtained from active amyl iodide by treatment, in alcoholic solution, with zinc and HCl in the cold (Just). Oil with characteristic odour, unlike 'benzoline.' Not attacked by Br or HNO. Chlorine yields

an amyl chloride (100°) Isopentane (1 cc) is oxidised by $K_2Cr_2O_7$ (4 g) and CrO_3 (20 g) in water (2000 cc) in five weeks at 15°-20° to isovaleric acid (Berthelot, *C R* 79, 1435)

Pentane $C(CH_3)_4$, Tetra-methyl-methane [α -20°]. (95°) H.F.p 47.850 H.F.v 44.950 (Thomson, *Zn.*) Formed by the action of $ZnMe_2$ on *tert*-butyl iodide (Lwow, *Z* [3] 6, 520, 7, 257, Naumann, *B* 7, 173) Appears also to be formed from CH_3COI , CH_3I and $ZnMe_2$ (L) Oil or gas

References—DI-BROMO, DI-CHLORO, and IODO-PENTANE

PENTANE CARBOXYLIC ACID v Hexoic acid

Pentane dicarboxylic acid v BUTYL MALONIC, ETHYL-GLUTARIC, Di ETHYL MALONIC, Di METHYL-GLUTARIC, PIMELIC, and Propyl-succinic acids

n Pentane tricarboxylic acid
 $CH_3(CO_2H)_2CH_2CO_2H$ [148°] Got by saponifying its ether with alcoholic potash Slender needles, v sol water, alcohol, and ether Splits up at 175° into CO_2 and propyl-succinic acid

Ethyl ether Et_2A''' (280°) S.G. $\frac{1}{2}$ 1.052 Made from ethane tricarboxylic ether $CO_2EtCH_2CH(CO_2Et)_2$, NaOEt, and PrI (Waltz, *B* 15, 608, *A* 214, 58) Oil, miscible with alcohol and ether

Mono-nitrile of the diethyl ether
 $CH_3(CN)(CO_2Et)CH_2CO_2Et$ (205° at 45 mm) Formed from sodium cyano succinic ether and *n*-propyl iodide (Ba. the, *Bl* [8] 1, 805, *C R* 108, 297) Oil

Pentane tri-carboxylic acid, Ethyl ether
 $C_2H_5CH(CO_2Et)_2CMe(CO_2Et)_2$ **Methylbutenyl tricarboxylic ether** (281.5° cor) S.G. $\frac{2}{3}$ 1.0575, μ_D 1.4347 at 20° Formed by the action of Na on a mixture of a bromo butyric ether and sodium methyl-malonic ether (Bischoff, *B* 22, 1817, 8180, 23, 647) Apparently the same acid (281.8° cor), S.G. $\frac{2}{3}$ 1.0523, μ_D = 1.4340 at 20° is got from sodium butane tricarboxylic ether and MeI Both acids yield, on saponification by H_2SO_4 , two methyl ethyl-succinic acids [88°] and [168°]

Pentane tri-carboxylic acid Ethyl ether
 $C_2H_5C(CO_2Et)_2CHMeCO_2Et$ (272.8° cor) S.G. $\frac{2}{3}$ 1.0609 μ_D 1.4374 at 20° Formed by the action of Na on a mixture of a bromo propionic ether and sodium ethyl malonic ether (Bischoff a. Mintz, *B* 28, 648) The same, or an isomeric ether (282.8° cor), S.G. $\frac{2}{3}$ 1.0607, μ_D = 1.4373 at 20° is got from sodium propane tricarboxylic ether and EtI Both acids yield, on saponification with H_2SO_4 , methyl ethyl succinic acid [168°], the acid prepared by the second method yields also the isomeric acid [88°]

Pentane tri-carboxylic acid
 $CH_3(CO_2H)_2CH_2CH(CO_2H)CH_2CH_2CO_2H$ [107°] Formed from pentane tetra carboxylic ether by saponifying with alcoholic potash, and boiling the product with conc. $HClAq$ (Emery, *B* 24, 284) Mass of needles— Ag_2A''' white pp

Pentane tri-carboxylic acid
 $C_2H_5CH(CO_2H)_2CH(CO_2H)CH_2CO_2H$ [148°] Made by the action of fumaric ether on sodium ethyl-malonic ether, the product being saponified (Auer, *B* 24, 810) V sol. water and alcohol

Isopentane tri-carboxylic acid
 $(CH_3)_2CHCH(CO_2H)_2CH_2CO_2H$ [145°] (Hjelt, *B* 18, 2622). Formed from its ether, which is

made from ethane tricarboxylic ether, NaOEt and PrI (Waltz, *A* 214, 60) Yields isopropyl succinic acid on fusion

Ethyl ether Et_2A''' (270°-290°).

Isopentane tri carboxylic acid
 $(CH_3)_2CHCH(CO_2H)_2CH(CO_2H)_2$ **Carbopimelic acid** [160°] Got from its ether by cold alcoholic potash Monoclinic (?) crystals (from water), v sol water, alcohol, and ether Decomposes on fusion into CO_2 and isopropyl-succinic (pimelic) acid.— Ba_2A''' , sticky pp

Ethyl ether Et_2A''' (277°) Made from α -bromo isovaleric ether and sodium-malonic ether (W Roser, *A* 220, 274) Liquid, with bitter taste

Isopentane tricarboxylic acid Ethyl ether
 $CMe_2(CO_2Et)_2CMe(CO_2Et)_2$ (284.5° cor) S.G. $\frac{2}{3}$ 1.0524 μ_D = 1.4333 at 20° Made from sodium methyl malonic ether and α -bromo-isobutyric ether (Bischoff a. Mintz, *B* 23, 649) Yields, on saponification by H_2SO_4 , tri-methyl-succinic acid [105°] and a small quantity of a substance melting between 108° and 122° The same, or an isomeric ether ((277.8° cor), S.G. $\frac{2}{3}$ 1.0575, μ_D = 1.4341 at 20°) is got from sodium isobutane tricarboxylic ether and MeI It yields, on saponification, tri-methyl succinic acid [105°] and α di methyl succinic acid [140°]

Pentane tetracarboxylic acid
 $(CO_2H)_2CHCH_2CH_2CH_2CH(CO_2H)_2$ Oil, split up by heat into CO_2 and π pimelic acid $CO_2HCH_2CH_2CH_2CH_2CH_2CO_2H$

Ethyl ether Et_2A' (260° at 100 mm) Got from sodium malonic ether and trimethylene bromide (Perkin, jun, *C J* 51, 241, *B* 18, 3248) Disodium salt $CH_2(CH_2CNa(CO_2Et))_3$. Amorphous powder, converted by bromine into pentamethylene tetracarboxylic ether

Pentane tetra-carboxylic acid
 $CH_2(CMe(CO_2H))_3$ [164°] Got by boiling its ether with alcoholic potash (Dressel, *A* 256, 183) Crystalline mass, v sol. water, m sol. ether Converted by heating, by itself or in aqueous solution, into $CH_2(CHMeCO_2H)_3$ [90°]

Ethyl ether Et_2A' (191° at 12 mm) Made by the action of MeI on a solution of di sodium propane tetra-carboxylic ether in alcohol Oil

Pentane tetra-carboxylic acid Ethyl ether
 $C(CO_2Et)_2(CH_2CH_2CO_2Et)_2$ (215° at 13 mm) S.G. $\frac{2}{3}$ 1.1084 *A* by-product in the action of NaOEt on a mixture of β -bromo-propionic ether with malonic ether (Emery, *B* 24, 288) Colourless oil

PENTANE PHOSPHONIC ACID v ISOAETHYL PHOSPHINIC ACID

PENTANE SULPHONIC ACID $C_5H_{11}SO_3H$ Formed by the action of nitric acid on isomethyl mercaptan, disulphide, or sulphocyanide (Gerathewohl, *J pr* 34, 447, Henry, jun, *A Ch* [3] 25, 246, Medlock, *A* 69, 225) Deliquescent crystalline mass Chlorine acting on it in sunlight forms $C_5H_9ClSO_3H$ (Spring a. Wunsinger, *Bl* [2] 41, 801) ICl_3 at 180° forms the same body, together with tri- and tetra-chloro-pentanes and $C_5H_9Cl_2(SO_3H)$

Salts.— BaA' , (dried at 100°). Gyrate on water— PbA' , saq colourless laminae— CuA' ,— AgA' plates Double salts with chloropentane sulphonates $BaA'(C_5H_9ClSO_3)_2\text{aq}$.— $Ba_2A'_2(C_5H_9ClSO_3)_2Cl_2\text{aq}$
Chloride $C_5H_9SO_3Cl$. Liquid.

Pentane disulphonic acid $\text{OEt}_2(\text{SO}_3\text{H})_2$. Formed from COI , SO_3H and ZnEt_2 in ether (Ilse, *A* 147, 145) Syrup— $\text{K}_2\text{A}''$ 2aq needles (from alcohol).— BaA'' 2aq *S* 14 in the cold, *c.* 80 at 100° — PbA'' — ZnA'' 4aq plates

PENTA-THIONIC ACIDS *Thionic acids* under SULPHUR, OXYACIDS, or, vol. IV

α -PENTATRIACONTANE $\text{C}_{32}\text{H}_{66}$ [75°]. (381° at 15 mm) *SG* $\frac{1}{2}$ 7816, $\frac{2}{5}$ 777, $\frac{1}{10}$ 766. Formed by reduction of the dichloride of steatone with *H* and *P* (Kraft, *B* 15, 1715)

PENTENE *v* **AMYLENE** The name pentene is also given sometimes to pentamethenyl trihydride

Dipentene *v* **TERPENES**

PENTENOIC ACID $\text{C}_5\text{H}_8\text{O}_2$, *i.e.* $\text{CH}_2\text{CH}(\text{CH}=\text{CH}_2)\text{CO}_2\text{H}$ *Propyldiene acetic acid* (196°) *SG* $\frac{1}{2}$ 998 Made by heating propionic aldehyde with malonic acid, and HOAc at 100° (Kommenos, *A* 218, 166, Zincke a Kuster, *B* 32, 494) Oil— BaA' plates [270°]— CaA' , aq nodules, *v* sol water and alcohol— CuA' , green pp [91°] Yields Cu_2O at 175° — AgA' bulky pp The same acid (*S* $\frac{1}{2}$ 983) appears to be got by adding sodium-amalgam to an alkaline solution of COI , COI , COI , COI , COI derived from pyrocatechin or *o*-amido phenol It combines with bromine.

Pentenolic acid $\text{CMe}_2\text{CH}(\text{CO}_2\text{H})$ *Dimethyl-acrylic acid* [70°] (195)

Formation—1 From CH_3Pr , CO_2H by oxidising to $\text{CMe}_2(\text{OH})\text{CH}(\text{CO}_2\text{H})$ and heating the product with dilute H_2SO_4 (Neubauer, *A* 106, 62, W. von Miller, *A* 200, 261)—2 From *d*-methylallyl-carbinol by oxidising to β -oxy-valeric acid and distilling this acid with dilute H_2SO_4 or treating it with PCl_5 (Semljanitzin a Saytzeff, *A* 197, 72, Ustinoff, *J pr* [2] 34, 478, *Bl* [2] 45, 255)—3 *A* product of the action of alcoholic NaOEt on a bromo-isovaleric ether (Duvillier, *C R* 88, 913, 1209, 112, 1012, *A Ch* [5] 19, 428)—4 The chief product of the action of sodoform on sodium isobutylate (Gorboff, *J pr* [2] 41, 228)

Properties—Monoclinic prisms, *a b c* = 153.1 71, $\beta = 74^\circ 13'$ *Sl* sol water, *v* sol alcohol and ether Bromine combines, forming $\text{C}_5\text{H}_7\text{Br}_2\text{O}_2$ [108°] Converted into isobutylene and CO_2 by heating for 30 hours at 220°

Salts— NaA' — BaA' , 2aq groups of needles— CaA' ,— CaA' , 4aq— ZnA' , 4aq— CuA' , 2aq— PbA' , aq— AgA' .

Isomerides—**ALLYL ACETIC**, **ANGELIC**, **TETRAMETHYLENE CARBOXYLIC**, and **TRIGLIC ACID**

PENTENYL ALCOHOL C_5H_{10} , *i.e.* $\text{CH}_2=\text{CH}(\text{CH}_2)_3\text{OH}$ *Vinyl-ethyl-carbinol*. (114°). *SG* $\frac{1}{2}$ 856, $\frac{2}{5}$ 840 Formed from acrolein by successive treatment with ZnEt_2 and water (Wagner, *J R* 16, 819; *B* 21, 3349, *Bl* [2] 42, 830). Pungent liquid. Yields on oxidation by KMnO_4 propionic and formic acids, CO_2 , and tri-oxy-pentane

Acetyl derivative $\text{C}_5\text{H}_9\text{OAc}$ (182°) at 748 mm. *SG*, $\frac{1}{2}$ 918, $\frac{2}{5}$ 898 Liquid

Pentenyl alcohol $\text{CH}_2=\text{CH}(\text{CH}_2)_3\text{OH}$ *Methyl-allyl-carbinol*. (116°) Formed by the action of zinc on a mixture of allyl iodide and acetic aldehyde (Wagner, *B* 21, 3850) Yields on oxidation a tri-oxy-pentane, formic and acetic acids, and acetic aldehyde.

Pentenyl alcohol $\text{C}_5\text{H}_{10}\text{O}$ *Valerylène hydrate* (115°–120°) Got by treating its acetyl derivative with solid KOH

Acetyl derivative $\text{C}_5\text{H}_9\text{OAc}$ ($\approx 95^\circ$). *A* product of the action of AgOAc on $\text{C}_5\text{H}_{10}\text{O}$, obtained by combination of valerylène with HCl (Reboul, *Z* 1867, 174)

Pentenyl alcohol $\text{CH}_2=\text{CH}(\text{CH}_2)_3\text{OH}$ *Methyl propenyl-carbinol* (118°) Formed by the action of water on the corresponding pentenyl chloride, which is a product of chlorination of $\text{CHMe}(\text{CH}_2)_3\text{H}$ (Kondakoff, *B* 24, 931)

Pentenyl alcohol $\text{CH}_2=\text{CH}(\text{CH}_2)_3\text{OH}$ *Dimethyl-vinyl carbinol* *Isoprenic alcohol* (39°) Formed from $\text{C}_5\text{H}_9\text{Cl}$, which is obtained from isoprene and HCl (Gradziatsky, *Bl* [2] 47, 168) Smells like camphor Dilute H_2SO_4 at 100° forms a hydrocarbon

Acetyl derivative $\text{C}_5\text{H}_9\text{OAc}$ (121°) *A* pentenyl alcohol (125°–135°) is a product of reduction of tiglic aldehyde (Lieben a Zeisel, *M* 7, 57)

References.—**BROMO** and **CHLORO PENTENYL ALCOHOL**

PENTENYL AMIDO-PHENYL MERCAPTAN

$\text{C}_5\text{H}_9\text{N}(\text{S})\text{CH}_2\text{Pr}$ Liquid, formed from *o*-amido phenyl mercaptan and isovaleryl chloride (Hofmann, *B* 13, 8, 1223)— $\text{B}'\text{H}_2\text{PtCl}_5$ needles **PENTENYLAMINE** $\text{C}_5\text{H}_9\text{N}$, *i.e.* $\text{CH}_2=\text{CH}(\text{CH}_2)_3\text{NH}_2$ (85°) Got by reducing nitro-amylene with *azinc* and HCl (Gal, *J* 1873, 333)

Pentenylamine $\text{C}_5\text{H}_9\text{NH}_2$ *Valerylamine* *A* product of the action of NH_3 on chloro amyl alcohol (amylene chlorhydrin) at 100° (Wurtz, *A Suppl* 7, 89)— $\text{B}'\text{H}_2\text{PtCl}_5$ orange crystals

PENTENYL BENZENE $\text{C}_5\text{H}_9\text{C}_6\text{H}_5$ (173°–177°) *SG* 84 Formed, together with a polymeride $\text{C}_{22}\text{H}_{22}$ (*c* 210°), *SG* 23 96, by boiling bromo-amyl benzene $\text{C}_5\text{H}_9\text{CH}_2\text{CHBr}(\text{CH}_2)_3\text{CH}_3$ with water (Dafert, *M* 4, 153, 616) Yields benzene and acetic acids on oxidation

PENTENYL BROMIDE *v* **BROMO-AMYLENE**

PENTENYL THIOCARBIMIDE $\text{C}_5\text{H}_9\text{NCS}$ (190°) Formed from isoamylene bromide by successive treatment with alcoholic NH_3 and CS_2 (Hofmann, *B* 12, 990) Liquid

PENTENYL-THIO-UREA $\text{C}_5\text{H}_9\text{NHCSNH}_2$ [103°] Formed from pentenyl thiocarbimide and alcoholic NH_3 at 100° (Hofmann, *B* 12, 991)

PENTHIOPHENE The homologue of thiophene $\text{CH}_2=\text{CH}(\text{CH}_2)_3\text{CH}=\text{CH}_2$, known in some of its derivatives (Krekeler, *B* 19, 3270)

PENTIC ACID $\text{C}_5\text{H}_8\text{O}_4$, *i.e.* $\text{CH}_3\text{CH}(\text{CO}_2\text{H})_3$ [128°]. *A* product of the action of alcoholic potash on bromo ethyl-acetoacetic ether (Demarçay, *Bl* [2] 27, 483, *C. R* 88, 126) Got also by heating bromo-ethyl-acetoacetic ether at 100° (Wedel, *A* 219, 104) Trimetric crystals, *v* sol. hot CHCl_3

Salts— NaA' , aq— KA' , aq: *v.* *c.* sol water MgA' , 5aq— CaA' , aq— BaA' , 2aq— ZnA' , 5aq— MnA' , 4aq

Ethyl ether $\text{CH}_3\text{CH}(\text{CO}_2\text{H})_3$ *Mol wt* 153 (obs by Raoult's method); (calc 156) (Moscheles a Cornelius, *B* 21, 2607; 22, 244). Liquid.

Phenyl-hydrasido $C_6H_5O_2(N,HPH)$
Crystals (from alcohol)

PENTINENES C_5H_8 , *Pentylenes*, *Pentines*.
Mol w 68 Of the eight possible isomerides, three are derived from acetylene and precipitate ammoniacal silver and cuprous solutions, the remaining five, so far as they are known, give no metallic derivatives The following are known

1 **Propyl acetylene** $CH_3CH_2CH_2C \equiv CH$ (43°-49°) Methyl propyl ketone by action of PCl_5 gives a dichloride, from which alcoholic potash withdraws $2HCl$ (Friedel, *Z* 1869, 124) Forms liquid dibromide (190°) and tetrabromide (275°) (Bruylants, *B* 8, 411)

2 **Isopropylacetylene** $(CH_3)_2CHC \equiv CH$ (28°-29°) The chloride from isovaleric aldehyde (Bruylants, *B* 8, 413) or isopropyl-ethylene bromide, is decomposed by alcoholic potash (Eltekow, Flawitzky, Krylow, *B* 10, 707 and 1102) Oxidised by chromic liquor into acetone, acetic, and isobutyric acids Bromides liquid, C_5H_7Br , (175°), C_5H_7Br , (275°) (Bruylants, *B* 8, 407) The silver compound C_5H_7Ag is white, dissolves slightly in ammoniacal silver nitrate solution, and crystallises therefrom in small prisms Converted by a solution of iodine in KI into the compound C_5H_7I (140°)

3 **Eltekow** (*B* 10, 1904), by treating commercial amylene with sulphuric acid (dil with $\frac{1}{2}$ vol H_2O) at 0°, obtained an insoluble amylene, the bromide of which, by the action of KHO , yielded a mixture of two hydrocarbons, of which one, the smaller amount (35°), gave a pp with ammoniacal silver solution This he believed to be isopropylacetylene, though differing from the compound described above in b p

4 **Valerylene** This name was given to the liquid (44°-46°) obtained by Reboul (*A* 131, 238) from the dibromide prepared from commercial amylene This substance is a mixture from which Eltekow obtained the hydrocarbon described under 3, and a larger quantity of another (51°-52°), which is stated (*B* 10, 2057) to yield acetic and propionic acids by treatment with chromic acid Hence Eltekow ascribes to it the formula, C_5H_8 , $C \equiv CCH_2$, although it does not yield metallic derivatives Reboul's valerylene, heated to 250°-260°, yields a terpene $C_{10}H_{16}$, (180°) (Bouchardat, *Bl* [2] 33, 24) Strong sulphuric acid converts it into a hydrate $C_{10}H_{18}$, H_2O and polymerides $C_{10}H_{22}$, &c (Reboul, *A* 143, 372) Yields a dibromide, which on treatment with alcoholic potash gives C_5H_7Br , C_5H_7 (valylene), and valerylene (Reboul, *A* 135, 372) Valerylene tetrabromide $C_5H_7Br_4$ is liquid.

5 **Piperylene** (42°) (Hofmann, *B* 14, 665) Distinguished by the production of a crystalline tetrabromide, fusible [114.5°] and volatile without decomposition Gives no pp with solutions of silver or copper, and probably has the constitution $(CH_3)_2C \equiv CCH_2$, or β -dimethylallene

6 **Isoprene** (about 37°) Originally obtained (v Gr Williams, *Tr* 1860, 241), along with caoutchouc and heveene, by destructive distillation of caoutchouc (g v) Formed also in notable quantity by the action of very low red heat upon turpentine oil and its isomerides (Tilden, *C J* 45, 410) A colourless, limpid liquid, S G 6823 at 20°. Oxidises rapidly on exposure to air,

forming a syrupy compound which, on being quickly heated, often explodes Contact with strong acids in the cold converts it into a substance apparently identical with caoutchouc Combines violently with bromine, forming a tetrabromide which remains liquid at -20° and is decomposed by distillation Absorbs hydrochloric acid, forming a mixture of mono- and dihydrochlorides $C_5H_8 \cdot 2HCl$ (145°-150°), unstable liquid Oxidised by chromic acid to carbonic, formic, and acetic acids, by nitric acid to oxalic and a small quantity of undetermined acid

Heated to 280° isoprene is converted into dipentene $C_{10}H_{16}$, identical with the product from turpentine (v TERPENES) (Bouchardat, *C R* 87, 654 a 89, pp 361, 1117) W A T

PENTINENE CARBOXYLIC ACID v HEXANOIC ACID

Pentylene tricarboxylic acid $C_5H_8O_6$, *s.s.* $C_5H_8(CO_2H)_3$, CH_2CO_2H

Allyl ethane tricarboxylic acid [151°] Got from its ether, which is made from sodium ethane tricarboxylic ether and allyl bromide (Hjelt, *B* 16, 333) At 160° it splits up into allyl succinic acid and CO_2

Ethyl ether Et.A''' (c 282°)
PENTINOIC ACID $C_5H_8O_4$, [206°] A product of the oxidation of oil of turpentine by HNO_3 (Roser, *B* 15, 293) Tables (from water), sl sol water

PENTINYL ETHYL OXIDE $C_7H_{12}O$, *s.s.* $C_5H_7OCH_2CH_3$, (125°-130°) Formed from C_5H_7Br $EOEt$ and alcoholic potash at 155° (Reboul, *A* 133, 86) Light oil Combines with bromine

PENTOIC ACID v VALERIC ACID

PENTONENE C_5H_8 , (42.5°) S G 808 V D 2.45 Occurs in oil deposited by compressed gas derived from bituminous shale (Etard a Lambert, *C R* 112, 945) Oil polymerises in the cold, becoming $C_{10}H_{16}$, which forms crystal line $C_{10}H_{16} \cdot 2HSO_4$

PENTONYL ETHYL OXIDE $C_7H_{12}O$, *s.s.* $CH_3CH_2C(OEt)CH_2$, (155°) V D 3.87 Got by heating di chloro pentylene $CH_2(Cl)CH_2$, with alcoholic potash (Combes, *A Ch* [6] 12, 223) Colourless liquid, with unpleasant small

PENTOSE Name given to compounds resembling glucose but having only 5 atoms of carbon in the molecule, e.g. tetra-oxy-valeric aldehyde

PENTYL Another name for the radicle **AMYL**

PENTYLENE v AMYLENE and PENTINENE

PENTYLENE GLYCOL v DI-OXY-PENTANE

PENTYLIC ACID IS VALERIC ACID

PENTYLIDENE v AMYLIDENE

PENTYLIDENE ACETONAMINE v ACETONAMINE

PEPPER OIL S G 1.875 The oil of black pepper is levorotatory ($\alpha = -3.2^\circ$) and contains a terpene (165°) which is levorotatory ($\alpha = -7.8^\circ$) and gives a crystalline tetrabromide (Eberhardt, *Ar Ph* [3] 25, 516, cf Dumas, *A* 15, 159, Soubeiran, *A* 34, 327)

PEPPERMINT OIL The essential oil from *Mentha piperita* contains menthol, $C_{10}H_{18}O$, and a liquid terpene The oil obtained by distilling *M. arvensis* deposits crystals of menthol, and appears also to contain $C_{10}H_{18}O$, an inactive product

of oxidation of menthol (Moriya, *C J* 39, 82) The oil from *M. viridis* contains a terpene and $C_{10}H_{16}O$ [225°] (Gladstone, *C J* 17, 1)

PEPSIN The digestive ferment of the stomach (Schwann, Brucke, *Sitz W* 43, 601, Schmidt, *A* 61, 311) It also occurs sometimes in the urine (Stadelmann, *Zeit Biol* 25, 208)

Preparation—1 The glandular layer of the stomach is extracted with dilute phosphoric acid,

with alcohol (Wittich, *J* 1870, 894, Podwysosky, *Pf* 39, 62)

Properties—Amorphous Does not give the xanthoproteic reaction, and is not coloured by H_2SO_4 and sugar It renders insoluble proteins soluble, converting them into peptones, this action is greatly enhanced by the presence of 05 p c HCl Digestion is more rapid at 37° than at 24° Dry pepsin does not lose its power by heating to 100° (Huppe, *C J* 44, 101) According to Gautier (*C R* 94, 1192), pepsin contains some insoluble amorphous granules which also possess digestive power in presence of HClAq (cf Béchamp, *Ch R* 94, 970)

PEPTONES v PROTEIDS

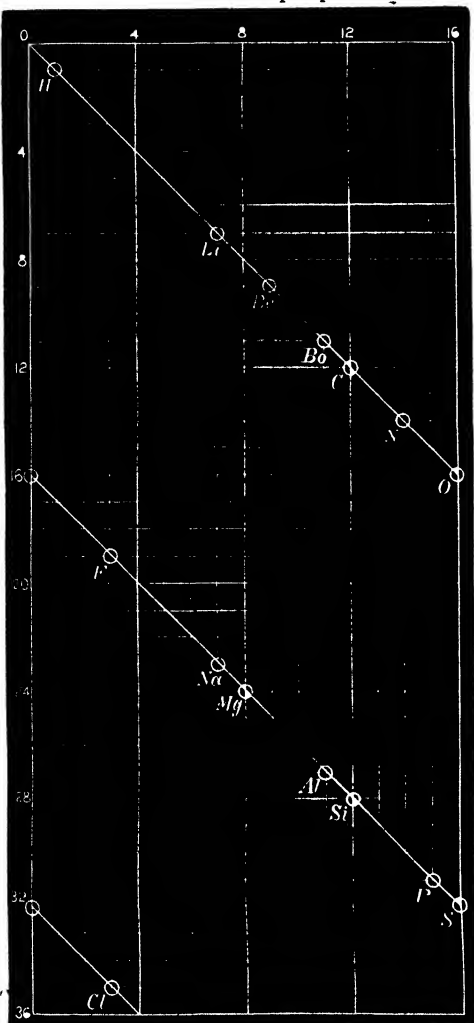
PER Use of this prefix applied to inorganic compounds, for per compounds and per salts v the element the per compound of which is sought for, or the salts to the name of which per is prefixed Thus peroxides are dealt with under OXIDES, peroxide of nitrogen under NITROGEN, oxides of, perchloric acid under CHLORINE, oxyacids of, permanganates under MANGANESE, OXYACIDS OF

PEREIRINE $C_{18}H_{24}N_2O$ [124°] An alkaloid, occurring with geissospermine in the bark of *Pao Pereiro* (*Geissospermum Vellosoi*, Peckolt) It is separated from geissospermine by ether (Hesse, *A* 202 147) Amorphous powder, nearly insol water, v sol alcohol and ether Conc H_2SO_4 gives a violet red and HNO_3 a purple red colour — $B^+H_2PtCl_6$ 4aq yellowish amorphous pp

PERIODIC LAW In the history of modern chemistry we find several attempts to trace connections between the properties of the elements and their atomic weights Dobereiner (*G A* 26, 331) was the first to show that a simple relation exists between the atomic weights of closely related elements Ca, Sr, and Ba are very closely related elements, and the atomic weight of Sr is the mean of those of Ca and Ba Again, Se has an atomic weight equal to the mean of the atomic weights of the closely related elements S and Te This doctrine of triad groups was further extended by Dumas (*J* 1851 291), Kremers (*P* 85, 56), and Lenssen (*A* 103, 121) Lenssen endeavoured to embrace all the elements in a classificatory system composed of 20 triads Pettenkofer (*A* 105, 188) pointed out that the atomic weights of analogous elements, e.g. the alkali metals, form simple arithmetical series, and he emphasised the analogy between such natural families of the elements and the homologous series of organic compounds Kremers (*P* 83, 56), Gladstone (*P M* [4]

5, 313), Cooke (*Am S* (2) 17, 387), and Dumas (*C R* 46, 709) followed with generalisations differing only in detail from that of Pettenkofer

These discoveries did not, however, herald any immediate great advance It was, of course, interesting to find that elements showing gradational similarity of properties possess atomic weights, which are also, so to speak, numerically similar, but the newly discovered relationships



the acid ppd by lime water, and the pp dissolved in dilute HClAq, the solution is then dialysed (Maly, *J pr* [2] 11, 104) — 2 The mucous membrane of pigs' stomachs is macerated with water containing HClAq and the pepsin separated from the filtrate by adding NaCl (Scheffer, *Ph* [8] 2, 761, 783, 843) — 3 The mucous membranes are kept 24 hours after death, and then extracted with glycerin The extract is ppd.

did not result in immediate practical applications, and did little more than emphasise pre-existing natural classifications. The investigations altogether failed to afford a generalisation capable of embracing the groups of related elements in a complete classificatory system. Instead of furnishing one inclusive scheme dealing with all the elements, they rather favoured a classification consisting of a number of isolated and partial systems, each system comprising only very closely allied elements, a classification similar in its imperfections to that which would have resulted in organic chemistry had chemists been content to differentiate empirically alcohols, aldehydes, and acids, without any knowledge of the relationships existing

oblivion from which it has but lately been withdrawn to the light of day (*v N* Dec 26, 1889; *C N* 63, 51).

Newlands was the first to look beyond the relations of analogous bodies merely, and to insist on a relationship of a higher order, connecting the properties and atomic weights of all the elements taken collectively.

This relationship, first roughly indicated by Newlands in 1864 (*C N* 10, 94), was further developed in 1865 under the title of the *law of octaves* as follows —

'If the elements are arranged in order of their equivalent, with a few slight transpositions, as in the accompanying table, it will be observed that elements belonging to the same group usually appear on the same horizontal line —

H	1	P	8	Cl	15	Co and Ni	22	Br	29	Pd	36	I	42	Pt and Ir	60
Li	2	Na	9	K	16	Cu	23	Rb	30	Ag	37	Cs	44	Tl	58
Be	3	Mg	10	Ca	17	Zn	25	Sr	31	Cd	38	Ba and V	45	Pb	54
B	4	Al	11	Cr	19	Y	24	Ce and La	33	U	40	Ta	46	Th	56
C	5	Si	12	Ti	18	In	26	Zr	32	Sn	39	W	47	Hg	52
N	6	P	13	Mn	20	As	27	Di and Mo	34	Sb	41	Nb	48	Bi	55
O	7	S	14	Fe	21	Se	28	Rh and R.J.	35	Te	43	Au	49	Os	51

between these distinct, but closely connected, classes of bodies.

The idea of arranging all the elements in the order of their atomic weights with a view to a more comprehensive classification based on the relations of these magnitudes and the salient properties of the elements, both chemical and physical, seems first to have suggested itself to M A E Déguyer de Chancourtois in the year 1862. His method of exhibiting the relationship was a geometrical one. On a right cylinder with circular base he traced his 'telluric helix' at a constant angle of 45° to the axis. On this curve lengths corresponding to the 'characteristic numbers' (atomic weights) were measured in terms of a unit equal to one sixteenth of the turn of the helix. The extremities of these lengths were the 'characteristic points' for the elements. De Chancourtois enunciated the principle that the relations between the properties of different bodies are manifested by simple geometrical relations between the positions of their characteristic points. For instance, elements with analogous properties fall together on vertical lines parallel to the generatrix. This relation becomes apparent when the cylinder is plane developed as in fig on p 808. It follows from this that the atomic weights of any natural group of elements are expressible in terms of the general algebraic formula $(n + 16m)$, where m is an integer, in other words, that the differences between the atomic weights of allied bodies are, in general, multiples of 16.

$$\begin{array}{cccc} \text{O} & \text{S} & \text{Se} & \text{Te} \\ 16, & 16+16=32, & 16+(4 \times 16)=80, & 16+(7 \times 16)=128 \end{array}$$

Other similar relations were developed by means of secondary helices.

De Chancourtois, evidently absorbed in the numerical relations exhibited by members of the individual groups, omitted to consider sufficiently the relations of the various groups, and so he failed to give clear expression to the periodicity of properties which his telluric helix implies. A mathematician and geologist, de Chancourtois expressed himself in terms not at all calculated to attract the attention of the chemical world at large, consequently his work fell into an

It will also be seen that the numbers of analogous elements generally differ by 7 or some multiple of 7, in other words, members of the same group stand to each other in the same relation as extremities of one or more octaves in music (*C N* 12, 83).

As will be seen, Newlands' law was retrospective rather than prophetic, it recapitulated and co-ordinated old facts, but did not suggest new ones. Though Newlands admitted the probability of the discovery of new elements, his rigid disposition into 8 octaves of the 62 elements then known neither admitted of interpolations nor suggested additions. In his development of the conception of chemical octaves there was a too evident subordination of fact, and chemists gave but little heed to a classificatory scheme which formed octaves at the cost of grouping together elements chemically dissimilar, and then only succeeded in producing octaves varying in range from 8 to 10 constituent elements.

In March 1869 Mendeléeff read a paper on the periodic law before the Russian Chemical Society. This periodic law was nothing else than a generalised and developed form of the law of octaves, the conceptions underlying both laws being practically identical. Yet Mendeléeff's law was the law of octaves so elaborated and expanded that it at once attracted general attention. Operative not only in the realm of the known, but boldly presuming to deal with the unknown, this periodic law clearly indicates methods of testing its own validity. The law states that the properties of the elements, as well as those of their compounds, are periodic functions of the atomic weights of the elements. That is to say, if in a co-ordinate system atomic weights be measured along one axis, and any measurable property along the other, a line joining the loci of the intersections of corresponding abscissae and ordinates will form a sinuous curve showing maxima and minima. As the atomic weight increases the property considered will not increase or diminish indefinitely, but will exhibit periodical waxing and waning in intensity.

The predictions to which Newlands lays claim in the preface to his pamphlet 'The Periodic Law' are founded on theoretical considerations at which he arrived independently of, and prior to, his law of octaves, and which, therefore, do not involve any idea of periodicity.

Contemporaneously with the enunciation of the periodic law by Mendeléeff, but quite independently, L. Meyer (*A Suppl* 7, 854) showed that such a periodic curve results when the relatively measurable property considered is atomic volume (at wt ÷ SG in solid state). In fact, nearly every measurable property—physical and chemical physical—has been investigated, and in every case the law has been substantiated.¹ On examination of the characteristic ultra-violet spectra of the elements, Hartley (*C J* 41, 84) found these to vary periodically in such a way that elements belonging to the same group have analogously disposed spectra, in fact, Hartley made use of this generalisation in assigning Be (*vide infra*) a place in the periodic system (*C J* 43, 816, see also Lecocq de Boisbaudran, *C N* 1896 (2) 4). Laurie (*P M* [5] 15, 42) obtained periodic curves when he plotted out the values of the heats of formation of haloid compounds of the elements in diagrams in which atomic weight was taken as the other variable. In this connection also Carnelley investigated the melting-points, boiling points, and heats of formation, of the halogen compounds of the elements (*P M* [5] 8, 1), the colours of corresponding compounds of the elements (*P M* [5] 18, 180), the occurrence of the elements in nature (*P M* [5] 18, 194), the salient physical properties of the compounds of the elements with alkyl radicals (*P M* [5] 20, 259), and finally the magnetic properties of the elements (*B* 12, 1958). Carnelley and Walker have examined the relation of the phenomenon of the complete dehydration of hydrated oxides to the periodic law (*C J* 53, 59). Prud'homme (*C R* 112, 236) found that the shades produced by using different metallic oxides to fix one and the same colouring matter varied periodically with the atomic weights of the metals. Brauner and Watts found confirmation of the law in studying the molecular volumes of the oxides (*B* 14, 48). Roberts Austen experimented with a like result on the tenacity of gold when alloyed with about 2 per cent of other metals (*Pr* 43, 425). Recently, Sutherland (*P M* [5] 80, 818) has asserted a periodicity of the vibration-periods of the atoms of elements at their melting-points.

For a fairly complete summary of such investigations showing that the atomic weights of the elements are the true variables which determine the properties of matter *v* Meyer's *Modern Theories of Chemistry*, pp 119-154.

At first sight it might seem that the best method of giving detailed expression to this periodic law would be the geometrical one of tracing periodic curves as above indicated, but in the present state of science this is impossible, for we have yet to learn methods of measuring chemical properties. Further, the periodicity which Mendeléeff asserts is peculiar in its discontinuity, and contrasts strongly with the continuous periodic curves of the geometer, such, for instance, as the curve of sines. Conclusions respecting *any* point on such a curve as the curve of sines, where the periodicity may be called continuous, are warranted, but unless geometrical analysis be modified in a special manner

the connecting by a continuous curve of the loci of intersection of ordinates and abscissae in a co-ordinate system, of which properties and atomic weights are the variables, amounts to a virtual negation of the indivisibility of atom, and to an assertion that the number of elements is infinite. No elements intermediate between, *e.g.*, silver and cadmium, and with atomic weights between 107.6 and 112, are known. Even if they existed, we could not for a moment suppose that they would form oxides with formulae containing a fractional number of oxygen atoms. Yet, as will be seen later on, if the periodic law were represented and interpreted geometrically it would certainly involve not only the existence of such intermediate elements, but also the existence of oxides and other compounds incompatible with the fundamental conception of modern chemistry—the conception of the atom.

For these and similar reasons it is not only desirable but imperative that the periodic law should have a non geometrical representation at present. Let the elements (hydrogen excepted) be arranged in order of their atomic weights. It will then be found that the properties vary gradually as the value of the atomic weight increases, that when the eighth element is reached we have reproduced in varying intensity many of the properties characterising the first. The same is true of the fifteenth element. Similarly, in the ninth and sixteenth elements we have the recurrence of the properties of the second in the series, and so on.

If now, instead of this linear disposition, we arrange the elements in two dimensions, placing elements with analogous properties in the same vertical lines, and if we suppose that certain elements exist which have not yet been isolated, we obtain the table on the following page (taken from Brauner (*C J* 41, 68), and differing in slight details from the one first proposed by Mendeléeff), consisting of twelve horizontal rows and eight vertical columns.

The horizontal rows are called 'series,' and consist of elements whose properties vary gradually from one member to the next. It will be seen that hydrogen, on account of the anomalous relations between the value of its atomic weight and those of succeeding elements, is regarded as constituting a series in itself. The vertical columns are called 'groups,' and comprise elements with similar properties, *i.e.* elements which would occur at comparable points on periodic curves, could such be traced as already indicated. The similarity in properties shown by members of a group is a regularly gradationed one, and while in a series the properties of the elements vary in kind, in a group the variation is, speaking widely, one of degree only.

It will be noticed that the perfect symmetry of the table and the analogy which for the first few series it shows with the notes of the diatonic scale disappear when the end of the fourth series is reached. The three elements Fe, Ni, and Co, following upon Mn, and showing no analogies with the elements of Groups I., II., and III. respectively (where, guided solely by considerations of symmetry, we should be inclined to locate them), inaugurate the eighth group of Mendeléeff's table, a group containing many of the more important industrial metals, all very

¹ It should be noted, however, that the specific heats of the elements in the solid state do not vary periodically with the atomic weights.

Series	GROUPS							
	I	II	III	IV	V	VI	VII	VIII
	R ₂ O	R ₂ O	R ₂ O	R ₂ O	R ₂ O	R ₂ O	R ₂ O	R ₂ O
1	H=1	—	—	—	—	—	—	—
2	Li=7	Be=9	B=11	C=12	N=14	O=16	F=19	—
3	Na=23	Mg=24	Al=27	Si=28	P=31	S=32	Cl=35.5	—
4	K=39	Ca=40	Sc=44	Ti=48	V=51	Cr=52	Mn=55	{ Fe=56 Ni=58.6 Co=59 (Cu=63)
5	Cu=63	Zn=65	Ga=70	Ge=72	As=75	Se=79	Br=80	—
6	Rb=85.5	Sr=87	Y=89	Zr=90	Nb=94	Mo=96	? 100	{ Ru=101 Rh=103 Pd=106 (Ag=108)
7	Ag=108	Cd=112	In=114	Sn=118	Sb=120	Te=125	I=127	—
8	Cs=133	Ba=137	La=139	Ce=140	Di=143	? 149	Sm=150?	? 152-156 4 Elements?
9	← ? 4 Elements 156 to 162 ?				Er=166	? 167	? 169	—
10	? 170	? 172	Yb=173	? 178	Ta=182	W=184	? 190	{ Os=190 Ir=192.5 Pt=194 (Au=197)
11	Au=197	Hg=200	Tl=204	Pb=207	Bi=208	← ? 2 Elements 212 to 220 ?		—
12	← ? 3 Elements 220 to 230 ?				Th=232	? 237	U=240	? 245

(In this Table the atomic weights are only approximate.)

closely allied from a chemical point of view (*A Suppl* 8, 147), and a group anomalous in that, even series only are represented, three elements occurring in each series. According to Mendeléeff, the metals Cu, Ag, and Au, similar in many respects to the elements of Group I, nevertheless show, in their higher oxidation forms and physical properties, such analogies with the members of Group VIII as to warrant their double representation in the table (*A* *lc* p 152). It should also be noticed that the table is so constructed as to indicate a subdivision of each of the groups I-VII into two sub groups or 'families,' one family in each case being constituted of members of even series only, the other of members of odd series only. This subdivision is more apparent in the following tabular ar-

the hydroxyl derivatives of the paraffins into the minor classes of primary, secondary, and tertiary alcohols—is that, although there is a general similarity between all the members of a group, yet there is a more pronounced similarity between the members of odd series and the members of even series respectively, in other words, alternate members of a group are in general more closely related than successive members.

Nevertheless the advisability of a rigid subdivision of all the groups as indicated is open to question. In the case of Group II this division is undoubtedly warranted. Mg, Ca, Zn, Sr, Cd, Ba, Hg have all certain properties in common. But the family Ca, Sr, Ba, on the one hand, and the family Mg, Zn, Cd, Hg, on the other, have respectively so many similarities that we are

Sub-groups	SERIES							
	1	2	4	6	8	10	12	
Ia.	Li=7.02	K=39.11	Rb=85.5	Cs=132.9	—	—	—	—
IIa.	Be=9	Ca=40	Sr=87.6	Ba=137	—	—	—	—
IIIa.	B=11	Sc=44	Yt=89.1	La=138.2	—	Yb=173	—	—
IVa.	C=12	Ti=48	Zr=90.6	Ce=140.2	—	—	—	Th=232.6
Va.	N=14.03	V=51.4	Nb=91	Di=142.3	—	Ta=182.6	—	—
VIa.	O=16	Cr=52.1	Mo=96	—	—	W=184	—	U=238.6
VIIa.	F=19	Mn=55	—	—	—	—	—	—
VIII.	—	Fe=56 Ni=58.7 Co=59 Cu=63.4	Ru=101.6 Rh=103.5 Pd=106.6 Ag=107.93	—	—	Os=190.8 Ir=193.1 Pt=195 Au=197.3	—	—
—	—	Cu=63.4	Ag=107.92	—	—	Au=197.3	—	—
Ib.	H=1.007	Na=23.06	Cu=63.4	Cd=112	—	Hg=200	—	—
IIb.	—	Mg=24.3	Zn=65.3	In=113.7	—	Tl=204.18	—	—
IIIb.	—	Al=27	Ga=69	Sn=119	—	Pb=206.95	—	—
IVb.	—	Si=28.4	Ge=72.3	Sb=120	—	Bi=208.9	—	—
Vb.	—	P=31	As=75	Te=125	—	—	—	—
VIb.	—	S=32.06	Se=79	I=126.85	—	—	—	—
VIIb.	—	Cl=35.5	Br=79.96	—	—	—	—	—
VIIIb.	—	—	—	—	—	—	—	—
	1	3	5	7	9	11	13	

(In this Table O=16 is the starting point of atomic weights. See F W Clarke, *C N* 63, 76.)

angement of the elements which Mendeléeff suggests as a truer representation than the foregoing table of the peculiarly compounded periodicity which obtains.

The reason for this subdivision—which may not inaptly be compared with the subdivision of

justified in making the subdivision in this case (*U CLASSIFICATION*, vol II p 204).

But in the case of Group V. the 'family characteristics' do not so predominate over those of the group. N, P, V, As, Sb, &c., are all so much alike in properties that here a resolution

into families is by some regarded as superfluous. In this group, it is said, is more conveniently studied as a whole, for now the 'group characteristics' predominate over those of the families (v. vol. ii p. 210)

The elements constituting the eighth group have so many characteristic properties in common that subdivision of the group in any way would be artificial and unnecessary

There have been various other methods (di- and tri-dimensional) suggested for representing the periodicity in properties of the elements, but none of them perhaps sums up the facts known more simply and clearly than Mendeléeff's table (v. Meyer's *Modern Theories*, p. 120, Reynolds, N 84, 428, Crookes, *C J* 53, 503, Bayley, *P M* [5] 13, 26, Kremers, *Physikalisch-chemische Untersuchungen*, Wiesbaden, 1869-70, Baumhauer, *Die Beziehungen zwischen den Atomgewichte und der Natur der Elemente*, Braunschweig, 1879)

On inspecting his table in the light of known facts, Mendeléeff was led to make the following generalisations.—

(i) Excluding Series 2, the most markedly non-metallic elements occur in odd series¹

Types		La	Be	B	C	N	O	F	—	—	—	Na	Mg	Al	Si	
P	S	Cl	K	Ca	Sc	Ti	V	Cr	Mn	Fe	Co	Ni	Cu	Zn	Ga	—
As	Se &c															

(ii) Omitting Series 2, only members of odd series form compounds with organic radicles (the organo metallic compounds)² Just as the hydrides of Pd, Cu, and Nb contrast strongly with the hydrides of elements belonging to odd

		Li	Be	B	C											
N	O	F	Na	Mg	Al	Si										
P	S	Cl	K	Ca	Sc	Ti	V	Cr	Mn	Fe	Co	Ni	Cu	Zn	Ga	—
As	Se	Br	&c													

series, so organic compounds of members of even series (should such be hereafter discovered) will possess properties very different from those of similar compounds with which we are at present acquainted

Mendeléeff expressly omits Series 2 in making these generalisations. If, excluding the very incomplete Series 10 and 12, we tabulate the differences between the atomic weights of corresponding elements of Series 2 and 4, 4 and 6, 6 and 8, we find that, whereas the mean difference of corresponding elements of consecutive series is about 42, the mean difference for Series 2 and 4 is only 84.5. Since properties and atomic weights are dependent, we should expect from the above relations that the properties of the elements of Series 2 would contrast with the general properties exhibited by the other members of the families which they head. Such is actually the case to a greater or less extent with all the members of Series 2; e.g. Li differs from the other alkali metals, and approaches the alkaline earths in its insoluble phosphate, its easily soluble

bicarbonate, its difficultly soluble carbonate, and in the formation of diglycollic acid by the interaction of LiOH and monochloroacetic acid. N is more closely related to P than to V; O to S than to Cr. Again F and Mn contrast very strongly. On this account Mendeléeff has styled the elements composing Series 1 and 2 *Typical Elements*. Though the differences (averaging about forty) between the atomic weights of Na and Mg (Series 3) and those of the corresponding elements Cu and Zn (Series 5) are greater than those derived from a consideration of Series 2 and 4, yet the properties of Na and Cu on the one hand, and of Mg and Zn on the other, contrast so strongly, in Mendeléeff's opinion, as to induce him to enrol Na and Mg also among the typical elements. It is open to question whether the differences between Mg and Zn are sufficient to justify the classification of the former as a typical element, most chemists would probably find in Na the limit of the typical elements. Meyer, however, insists that the differences between Mg, Al, Si, and the other members of the corresponding groups is such as to justify the enrolment of these three substances among the typical elements, as follows.—

Yet the typical elements, after nitrogen exhibit so little analogy with the groups placed below them in the above scheme that Meyer proposes the following as perhaps a more scientific disposition of the typical elements —

Some go still further than Meyer, and regard H to Cl inclusive as typical elements, but such extreme views can scarcely be regarded otherwise than as unwarranted subordination of fact to considerations of symmetry suggested by Mendeléeff's table

Mendeléeff compares his so called typical elements to the lower members of homologous series in organic chemistry, which, as is well known, possess many properties peculiar to themselves and unrepresented in the higher members. In this connection contrast H_2O and CH_3O with the higher alcohols of the $C_nH_{2n+2}O$ series

While some disapprove of the title 'typical elements,' maintaining that this nomenclature connotes exactly the opposite of what it should, seeing that the typical elements are by no means types (as this word is generally understood) of the groups or even of the families which they head, others attempt to justify Mendeléeff's nomenclature. The former would rechristen the typical elements as anomalous elements, the latter would either find in the typical element of the group a combination of individual characteristics of each of the constituents of the sub-groups (vide vol. ii. p. 207), or in the typical elements as a class a representation of the gamut of variations of chemical properties.

(iii.) The passage from even to odd series is

¹ To render these generalisations unconditional Walker (*C F* 63, 351) has proposed that H to F be regarded as constituting Series 1, Series 2 being regarded as wanting. If this proposal is adopted, it becomes hardly necessary to differentiate typical elements (*C f. p. 2*)

² Recently it has been shown by Winkler (*B. 24*, 872, 1886) that the same generalisation can scarcely be applied to the hydrides also.

accompanied by a more gradual change in the properties of the elements than is the passage from an odd to an even series. This fact is in a manner emphasised in Mendeleëff's second table

(iv.) As the at wt increases in each group —

(a) The elements become more electropositive, and their oxides become more basic. This is well exemplified by the oxides of the types R_2O_3 and R_2O_5 formed by the elements of Gr V

(β) Corresponding compounds of the elements become more easily reduced. This is well instanced in the triplets S, Se, Te, and Cu, Ag, Au

(γ) There is a tendency to form stable oxides (and the salts corresponding to these) containing less O than the maximum salt-forming oxide characteristic of the group (*vide infra*). Thus, the characteristic oxide of Gr V is of the type R_2O_5 , and the haloid and oxyhaloid salts of the lower members of the group P, V, Nb, belong to this form, e.g. PF_5 , $VOCl_3$, $NbCl_5$, but Bi, which is the highest known member of Gr V, forms salts corresponding to the oxide R_2O_3 . Again in Gr IV no oxide of Si having the formula SiO_3 is known, SnO is known but is unstable, while PbO figures as the stable oxide of lead

(v.) Periodicity of valency and limiting forms. It has long been admitted that the atoms of those elements which form hydrides are limited with respect to the maximum number of H atoms which they can hold in combination, but it was only with the introduction of the periodic law that it was noticed that this limited power of combining with H atoms varies in a regularly periodic manner with the atomic weights. Little is as yet known of metallic hydrides, so that we have to look to Groups IV–VII, consisting of what may broadly be called the non-metals, for a confirmation of this periodicity

For instance —

Gr IV	furnishes us with	CH_4 , SiH_4 ,
Gr V	" "	NH_3 , PH_3 , AsH_3 , SbH_3 ,
Gr VI	" "	OH_2 , SH_2 , SeH_2 , TeH_2 ,
Gr VII	" "	FH , ClH , BrH , and IH

In the same way, but more perfectly, the law teaches us that the maximum number of oxygen atoms with which the elementary atoms can severally combine to form definite salt-forming oxides—either acidic or basic, as the case may be—also varies periodically with the atomic weights of the elements. The oxides characteristic of Gr IV are of the type R_2O_3 (or RO_2), those of Gr V of the type R_2O_5 , and so on regularly till we come to the eighth group, which has maximum salt-forming oxides of the type R_2O_7 (or RO_3). In carrying back this regularity to the groups I–IV, consisting for the greater part of metals, we find that it assigns to each of these groups exactly that type of oxide which we know to be characteristic of the individual members of the group. Hence we are justified in broadly asserting that the types of the hydrides (so far as they are already known), as well as those of definite salt-forming oxides, are periodic functions of the atomic weights of the elements. At the present time they are merely considerations of a regular periodicity, and not facts actually known, that lead us to assign B_2O_3 to Group VII as its characteristic oxide. Fluorine, so far as we know, forms no oxides at all; and

the highest halogen oxide is of the type R_2O_7 . In fact, Mn is the only member of the group that has an oxide conforming to the type R_2O_7 . Yet, having regard to the successes (to be detailed later) attendant on other predictions of the law, founded only on considerations of symmetry and general plan, all this is simply tantamount to admitting that the conditions of the existence of such oxides as Cl_2O_7 , I_2O_7 , &c., are legitimate subjects for research. Again, in Gr VIII the only oxides of the form RO , yet isolated are RuO , and OsO ,¹ but chemistry is by no means a completely worked out science wanting nothing, and the periodic law would be at fault did it fail to mirror forth such shortcomings and imperfections as still exist. The forms of complex oxyacids are determined by the forms of the maximum oxides, and may be derived from these by replacing O by its equivalent (OH), or H . Thus from SO_3 we can derive the chief oxyacids of S $SO_3(OH)_2$, $SO_2H(OH)$, and SO_2H_2 .

The highest compounds of the elements with halogens also correspond in type to the maximum oxide forms, and therefore exhibit a regular periodicity. In many cases, however, e.g. $TeCl_4$, ICl_3 , $AsCl_3$, only haloids lower in type than the maximum oxides are known, but in no case is a simple haloid known of higher type than the maximum oxide of the element considered

Against the doctrine of a maximum oxide form peculiar to all the members of each group, and of the types assigned (Table, top p 811), the existence of such oxides as K_2O_3 , BaO_3 , &c., has been adduced. In reply it has been pointed out that these oxides belong to the class of peroxides,² all of which are distinguished by their instability in the presence of the so-called 'contact agents,' as well as by their inability to form corresponding salts with a given basic or acidic oxide, as the case may be, and that a generalisation explicitly extending only to salt-forming oxides, strictly so called, cannot be impugned by considerations originating in the study of a class of bodies lying wholly without the pale of its jurisdiction

Yet this reply is scarcely satisfactory, for in certain cases we are bound, in order to justify the principle advanced, to recognise such bodies as PbO_2 and Bi_2O_5 —oxides currently regarded as peroxides. But PbO_2 is undoubtedly an acidic oxide, forming with basic oxides the corresponding plumbates, and Bi_2O_5 would also seem to have the power of forming with strong basic oxides unstable bismuthates—so unstable, in fact, that their true composition is still very doubtful. Hence Mendeleëff (*B* 15, 242) asserts that the oxides in question are *not* peroxides, for, according to him, true peroxides such as BaO_2 , Cr_2O_3 , ThO_2 , H_2O_2 , cannot form corresponding salts,³ and must of necessity contain relatively

¹ Some of the recently isolated carbonyl compounds of the elements of Group VIII. conform to type MO_2 (*O J W*, 748, 59, 1090, *C R* 119, 1481)

² It seems more than probable that a wider study of the higher forms of oxidation will result not only in the discovery of several new peroxides, but also of a distinct periodicity of type amongst this class of bodies also.

³ The recent isolation of persulphates, M_2SO_8 , by Berthelot (*C R* 112, 1481) and Marshall (*O J W*, 771) does not harmonise with Mendeleëff's views. Nor does Péchard's work (*O R* 112, 730, 1080) on higher oxidised compounds of Mo and W

more O than do the maximum salt-forming oxides (acidic or basic) as defined by the periodic law (v also Piccini, *B* 18, *Ref* 255). Still the case of CuO (which cannot be regarded as a peroxide) presents a difficulty, if Cu is rightly placed in Gr. I, for, in accordance with this position of Cu among the elements, the principle under discussion would lead us to expect a maximum salt-forming oxide of the formula Cu_2O .

In conclusion it may be said that although there undoubtedly is some connection between the at wts of the elements and the types of oxides they form, yet with the imperfect methods which we have of representing the periodic law, and in the absence of any firmly grounded and generally recognised classification of oxides, it is impossible to give a hard-and-fast enunciation of this connection.

We may assert that, so far as we are aware, a single atom of an element combines with at most four atoms of O or four atoms of H. The periodic law teaches us that the hydrogen holding power of the non-metals decreases regularly with increasing at wt, while the oxygen-holding power, as measured by the group oxides, increases regularly in such a way that the sum of the number of equivalents of both is equal to eight. For instance, S combines with two equivalents of H to form H_2S , it cannot, therefore, form a higher salt-forming oxide than SO_2 , which contains six equivalents of O.

Some chemists have proposed to extend this principle to the metals, i.e. to Groups I, II, and III. Since Na combines with one equivalent of O to give a salt-forming oxide, it must, they argue, combine with not more than seven equivalents of H or its equivalent. Similarly Ba combining with two equivalents of O must combine with six equivalents of H, and so on. But Brauner (*Sitz W*, 'Math-naturwiss Classe,' 84, 1165) would go still further. Let X denote a single equivalent of any element or radicle ($\text{X} = \text{H}, \frac{\text{O}}{2}, \frac{\text{S}}{2}, \frac{\text{N}}{3}, \text{OH}, \&c$), then he asserts

that the whole of the compounds of Na will be constituted on the types NaX or NaX_2 , or on types intermediate between these two, that all the compounds of Ba will be constituted on some of the types included in the scheme $\text{BaX}_2, \text{BaX}_3$, and so on. In short he would make out that there are certain limiting forms defining the complexities not only of the binary, but of all the compounds formed by each element, and that the range of these limits varies regularly and periodically with the atomic weights. In order to substantiate this view, recourse has been had to formulae not yet authorised by Avogadro's law, and sweeping assumptions are made as to the dispositions of the constituent atoms or atomic groups in the molecules or reacting masses of the bodies taken to exemplify this doctrine of limiting forms. Thus, out of all the numerous compounds of the alkali metals, the only one that can be found conforming to the type RX , is the substance NaOH $\frac{8\text{H}_2\text{O}}{2}$, and then only provided we manipulate the constituent atoms as follows: $\text{Na}(\text{OH})(\text{OH})\frac{\text{H}_2}{2}$. Si belongs to Group IV, therefore, according to Brauner, all its compounds ought to be of the type RX ,

its oxide SiO_2 , certainly conforms to the rule, but the highly characteristic body SiH_4F_2 , does not. Again, can such compounds of Pt as $\text{PtCl}_2\frac{8\text{H}_2\text{O}}{2}$, $\text{PtCl}_2\frac{2\text{HCl}6\text{H}_2\text{O}}{2}$, &c, be regarded as belonging to the type RX_2 ? These are one or two examples, out of very many, of the kind of difficulties that stand in the way of a full recognition of Brauner's extension of Mendelëff's views (v CLASSIFICATION, vol ii pp 210-13).

The idea that the valencies of the atoms vary regularly and periodically with their weights is one which has often been mooted, but in the absence of any recognised definition of valency it is an idea difficult to criticise.

If, as was originally done, the valency of an element be referred back to some constant objective attribute of the atom, and measured by the number of H atoms with which the given atom can combine, a partial periodicity involving only groups IV-VII is undoubtedly apparent.

Group	IV	V	VI	VII
	CH_4 SiH_4	NH_3 PH_3 AsH_3	OH_2 SH_2 SeH_2	FH ClH BrH
Valency	IV	III	II	I.

This conception of constant valency not only lacks in generality (since out of the 70 or so elements only some 17 combine with H), but for the establishment of its validity demands a division of compounds into the two classes of atomic and molecular—a division altogether too artificial to be admitted. If we forgo the idea of constant valency, and merely attempt to define the valency of an element as a number expressing the maximum number of monovalent atoms (H, F, Cl, Br, I) with which a single atom of the element in question combines to form true gaseous molecules, i.e. if we give a mere name to the maximum number of 'equivalents' represented in an atom, the alleged periodicity is still far from being perfect, as the table on the following page, embracing the latest results, shows.

Regarding this question of valency in the light of what has been said of limiting hydride and oxide forms, it is at least apparent that our crude conceptions of valency do not admit of a satisfactory quantitative interpretation. The sulphur atom fixes 2 equivalents of H and 6 of O, the arsenic atom fixes 3 equivalents of H and 5 of O. Why, we may ask, should we decide to overlook these essential relations exhibited by the oxides, to say nothing of the peroxides, and regard the valencies as deduced from the hydrides and haloids merely as fundamental properties of the atoms?

Mendelëff himself regards the theory of valency as extreme and not very valuable. Originating in the study of carbon compounds, valency finds in the domain of organic chemistry its only legitimate application, for the carbon atom, in accordance with its position in the periodic scheme, combines with *equal* numbers of equivalents of H and O, and, further, carbon compounds do not form so-called molecular compounds. He maintains that the forms of chemical compounds (including here isomerism, and therefore structural formulae) are fully explained, without the spurious aid of a statical

Series	GROUPS							
	I	II	III	IV	V	VI	VII	VIII
1	H ^I							
2	Li	Be ^{II}	B ^{III}	C ^{IV}	N ^V	O ^{VI}	F ^{VII}	
3	Na	Mg	Al ^{III}	Si ^{IV}	P ^V	S ^{VI}	Cl ^{VII}	
4	K	Ca	Sc	Ti ^{IV}	V ^V	Cr ^{VI}	Mn ^{VII}	Fe ^{VIII} Ni Co
5	Cu	Zn ^{II}	Ga ^{III}	Ge ^{IV}	As ^V	Se ^{VI}	Br ^{VII}	
6	Rb ^I	Sr	Y	Zr ^{IV}	Nb ^V	Mo ^{VI}		Ru Rh Pd
7	Ag ^I	Cd ^{II}	In ^{III}	Sn ^{IV}	Sb ^V	Te ^{VI}	I ^{VII}	
8	Cs ^I	Ba	La	Ce	Di			
9					Er			
10			Yb		Ta ^V	W ^{VI}		Os Ir Pt
11	Au	Hg ^{II}	Tl ^I	Pb ^{IV}	Bi ^{III}			
12				Th ^{IV}		U ^{VI}		

(The index numbers express valencies.)

theory of valency, in terms of the periodic principle together with more generalised views on substitution involving the recognition of Newton's third law, which states that action and react on are equal (*v A Suppl* 8, 211, *N* 40, 354)

Uses of the Periodic Law—An induction of any value should be suggestive of deductions admitting in their turn of experimental confirmation. Judged from this standpoint, the periodic law is well worthy the exalted position accorded it among the principles of chemistry, for it has opened up immense fields of research which deduction has already, to a small extent, successfully explored.

In order to maintain a general identity of properties in the vertical columns or groups of Mendeléeff's table, it was found necessary to make gaps here and there, to leave certain series unrepresented in the various groups. It was asserted that these empty places in the scheme were the positions of undiscovered elements for which Mendeléeff proposed a provisional and temporary nomenclature. Thus, in the year 1869, the element next to Ca = 40 with a higher atomic weight was Ti = 48, but Ti could not find a place in Group III Series 4, as its properties resemble those characterising Group IV and show no analogies with those of the other members of Group III. Ti was accordingly placed in Group IV Series 4, and the vacancy in Group III Series 4 was assigned to a then unknown element provisionally styled *ekaboron*. The principles of this nomenclature are very simple. The predicted element takes its temporary name from the one immediately above it in the group family, the Sanskrit prefixes *eka*, *dui*, *tri*, &c. being prefixed according as the unknown element is one, two, &c. removes lower down in the family than the name determining element.

Those elements of a family which stand immediately above and below a gap, together with those which immediately precede and succeed it in the series, are called the *atom-analogues* of

the element to which the gap corresponds. Thus B, Yt, Ca, and Ti are the atom analogues of ekaboron. Now it follows from the very nature of the law that the properties of any given element must be determinable from those of its atom analogues, that the properties of ekaboron, for instance, must be similar to, but intermediate in intensity between, those of B and Yt, and at the same time while dissimilar from those of Ca and Ti, they must show an intermediacy in their dissimilarity. Hence it becomes possible to predict the properties of still undiscovered elements, the mean of means of the properties of the atom-analogues forming the basis of the prophecy. How closely the properties of ekaboron thus predicted by Mendeléeff tallied with the properties of Sc experimentally investigated ten years later by Nilson is shown in the article ATOMIC AND MOLECULAR WEIGHTS (vol 1). Even were this case of ekaboron an isolated one, the wonderfully exact concordance between prediction on the one hand and experimental realisation on the other would go far to establish the periodic law as a generalisation of the highest order. But ekaboron is not an isolated example of the prophetic infallibility of the law which has as strikingly asserted itself in connection with the two recently discovered elements Ga and Ge (*q v* vol 11 597, 610).¹ The periodic law is and will be to the science of chemistry what Newton's law of gravitation was and is to the science of astronomy. Neptune had its place assigned in the worlds around us before it was seen, before they were discovered. Sc, Ga, and Ge had their properties detailed and their places assigned them among the elements, which by means of the periodic law have been raised from the level of 'mere fragmentary and incidental facts in nature'.

The validity of Mendeléeff's generalisation has also been confirmed in connection with the question of atomic weights. Before the enun-

¹ It yet remains to be seen whether the new element, aurtiacum, separated by Branner from tellurium ores, will identify itself with dvi tellurium, whose properties have recently been specified by Mendeléeff.

elation of the periodic law the values for the atomic weights formed a series of isolated and purely empirical numbers, the atomic weight of an element once ascertained, there was nothing in the actual numerical value itself, even when considered along with the properties, either to confirm or cast doubt on it as the true atomic weight. But the periodic law teaches that the atomic weights are by no means disconnected quantities, but that, taken in connection with the properties of the elements, their values constitute a series of mutually corrective numbers; in short, the law gives significance to the expression 'the probability of an atomic weight'.

In dealing with this aspect of the law it will be well to distinguish two sets of cases —

α . Those in which the law has actually fixed certain atomic weights indeterminable at the time by other means

β . Those in which the law has merely served to correct the values of atomic weights inaccurately determined by the usual methods

As illustrative of α we may consider the case of Be¹. The equivalent of Be having by accurate experiment been fixed at 4.51, it follows that the atomic weight must be numerically equal to $n \times 4.51$, where n is an integer. At first chemists were led to assign the formula Be_2O_3 to the oxide of Be on account of its isomorphism and many points of similarity with Al_2O_3 . This taken in conjunction with the analysis of the oxide makes $n=8$, and consequently fixes the atomic weight as 13.5. But Brauner repeatedly emphasised the view that the oxide of Be has the formula BeO , and that Be has therefore the atomic weight 9. The keynote of the many arguments advanced by Brauner in favour of his views was the incompatibility of the existence of an element with an atomic weight of 13.5, and having the properties of Be, with the system of the elements as arranged and classified by the periodic law.

In this system he argued that, not only was there no room for an element at wt 13.5, but that the proved existence of such an element would be totally subversive of the law. On the other hand, he pointed out that a vacancy existed in Series 2, Group II, for an element with an atomic weight equal to 9, and a careful examination of the relations of the members of Series 2 as a whole to those of other series, taken along with the known relations of beryllium to the magnesium group, absolutely demanded in his opinion the filling up of this vacancy with the element beryllium. Brauner's views were fully confirmed by Nilson and Pettersson, who succeeded in taking the vapour density of beryllium chloride, the application of Avogadro's law to their results gave the atomic weight of Be equal to 9 and the formula of the oxide BeO .

We are inclined to wonder that the question of the atomic weight of so rare and comparatively unimportant an element as Be has originated such a large amount of work and stimulated such lively discussion, unless we remember that a question of much greater import than the atomic weight of Be was at issue, the validity of the periodic law itself was at stake. And it is of interest to note that Mendeléeff regards the substantiation of Brauner's views on Be as a confir-

mation of his law of the same order as the discovery by Nilson of Sc, the ekaboron of prophecy. In a similar way the suggestions thrown out by the periodic law as to the atomic weights of U, Ce, and In have all met with corroboration. Up to the date of the periodic law, Pélitot's value 120 (= three times the equivalent 40) was received as the atomic weight of U, but Mendeléeff (*l.c.* 178) suggested six times the equivalent, or 240, as the correct atomic weight, thus conferring on U the distinction, which it is now universally admitted to hold, of being the element with the highest known atomic weight.

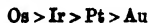
In 1870 the recognised atomic weight of Ce was 92 (2×46). Mendeléeff (*l.c.* 163), however, showed that no place existed in the system for an element with this atomic weight, and showing the properties of Ce, but that a suitable vacancy existed in Group IV Series 8, if Ce was regarded as having an atomic weight one and a half times the then accepted value. Mendeléeff's proposal has been fully justified by later work on the cerite metals (*C J* 41, 68).

The equivalent of In is 87.8, and formerly its atomic weight was taken equal to twice this, or 75.6. But Mendeléeff (*l.c.* 174) showed the necessity of trebling the equivalent, thus assigning the value 113.4, which is now accepted, to the atomic weight of this metal. Similarly Mendeléeff suggested, what has not yet been very satisfactorily settled, that the atomic weight of yttrium is three times its equivalent number, 29.87 (*v. EARTHS*).

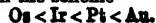
The investigations on tellurium are fairly illustrative of the cases summed under the heading β . The adoption of the value 128 for the atomic weight of Te as determined by Berzelius would necessitate iodine (at w 126.5) taking precedence of Te in Series 7 of the periodic scheme, iodine being thus separated from the rest of the halogens and falling into Group VI with such elements as S and Se, with which it shows no analogies, and Te falling into Group VII with such elements as Cl, Br, and Mn, with which it in turn shows no kinship. This violation of the principle of identity of chemical behaviour in the groups suggested some grave error in the accepted atomic weight of Te, an error causing this element to succeed iodine in the series instead of preceding it, as the general plan of the law requires. The subject has recently been investigated by Brauner (*C J* 55, 382), who for a second time has vindicated the law in a most striking manner by showing that the *ce-devant* tellurium is probably not elemental, and that the atomic weight of unalloyed tellurium is considerably lower than that of iodine, being equal to about 125.

In much the same way the law has led to the correction of the previously accepted atomic weights of osmium, platinum, and gold.

Formerly the accepted atomic weights of Os, Ir, Pt, and Au were in accordance with the scheme



But from the analogies existing between Os, Ru, and Fe, and the disposition of the first two series represented in Group VIII Mendeléeff (*A. l.c.*) predicted alterations of the atomic weights in accordance with the scheme



¹ In what immediately follows the values given for atomic weights and equivalents are only approximate.

This prediction has been fully confirmed by recent researches, it has not, however, yet been found possible to verify and substantiate the results now accepted for the last three metals by the application of Avogadro's law

As will be seen from the case of tellurium, the periodic law in its relation to atomic weight determinations is broadly suggestive rather than accurately definitive. It may be that ere long the discovery of the exact character of the periodicity, which at present we comprehend only vaguely, will raise Mendeléeff's law to the rank of an instrument for the absolute evaluation of atomic weights.

Mendeléeff has repeatedly emphasised the great advantage accruing to students and chemists generally from an adoption of the periodic classification as a working basis—the

	I	II	III	IV	V	VI	VII	VIII
a	$\left\{ \begin{array}{l} C_2H_4 \\ C_2H_2 \end{array} \right.$	C_2H_2	C_2H_4	C_2H_2	C_2H_4	C_2H_2	C_2H_4	C_2H_2
b	$\left\{ \begin{array}{l} NH_2C \\ \text{or} \\ NH_2(CH_3) \end{array} \right.$	NH_2C	NH_2C	NH_2C	NH_2C	NHC	NC	

advantage lying in the great mnemonic power of the law, which, introduced into the disheartening chaos of accumulated chemical fact, at once resolved the complexities of the apparent and rearranged them so as to exhibit the simplicities of the real. With one mental effort we commit to memory a large mass of facts which previously demanded so many independent but severally no less taxing mental exertions. No longer is phosphorus studied as an element apart from arsenic, but the general scheme of properties of the whole fifth group is learnt, and the facts about P, As, V, &c., are easily mastered by remembering their small individual deviations from this general scheme.

The law is also useful in that it points out the directions which should be taken by future research on the as yet unknown compounds of well known elements. It is well called 'the finger post of modern chemistry.' Examples of its utility in this respect have been indicated when treating of the law in its relation to limiting forms and to atomic weight determinations. In fact, every generalisation made in connection with the law suggests new work, organo-metallic compounds of In and Tl, and of the form MR_n (where R = organic radicle) must be forthcoming (*v. supra*, ii), a suboxide of Os having the formula Os_2O is looked for (*v. iii*), while various new peroxides of Mo and W, also a true peroxide of lead (Pb_2O_3) are very probable realisations of the future (*v. v*).

Theories having their origin in the Periodic Law—Although, according to Mendeléeff (*C J* 55, 684), the periodic law, solely founded as it is on the solid rock of well-ascertained chemical fact, in no way indicates the nature of the elements nor predicates the existence of a unique matter, yet many have made it a peg whereon to hang theories respecting either the compound nature of our elements or the existence of a primordial matter.

The simple relations holding between the

* The existence of Curium's azotimide N_2R was foreseen by Mendeléeff (*B* 29, 2464).

atomic weights of elements constituting some of the natural families, and the analogy between the relations so obtained and those found between the molecular weights of members of homologous series, led Dumas (*C R* 45, 46, 47) and Gladstone (*P M* [4] 5, 313) to suggest that the so called elements are not primary, but are composed of two or more simpler elements.

Pelopidas (*B* 16, 1868) called attention to the fact that the residues of hydrocarbons and nitrogenous organic bodies can be arranged in a periodic system, exhibiting in the number of members constituting the period, as well as in the gradual passage in each period from basic to acidic characters, a close analogy to Mendeléeff's periodic arrangement of the elements. The following are examples of the periods arranged by Pelopidas.—

The monovalent radicles $NH_2(OH_2)$ and C_2H_5 , undoubtedly show many analogies with the alkali metals, and CN has always been compared with the halogens of Group VII. Sulphur, Group VI, forms the acid $SO_2(OH_2)$, so the radicle C_2H_5 forms the acid $C_2H_5O_2(OH_2)$, and so on.

Carnelley (*C N* Nos 1875 and 1878), from a consideration of the fact that the salient physical properties of the normal halogen and alkyl compounds of hydrocarbon radicles exhibit relationships similar to those of corresponding compounds of the elements, develops the view that elements are analogous to the hydrocarbon radicles both in form and function. On the supposition that C and H are true elements, he then attempts to draw up a scheme of compounds of two primary elements, A and B, corresponding to Mendeléeff's scheme of the alleged elements, the compounds possessing the same 'atomic weights' and showing the same periodicity as the accepted elements. In this scheme the elements appear as bodies of the type A_nB_{m+n-2} , where n is the group and m the homologous series to which the element belongs, A is a tetrad element identical with carbon, and B, 'probably the ethereal fluid of space,' is a monad element with atomic weight = -21. These views of Carnelley are as much in advance of the earlier ones of Dumas and Gladstone as is the periodic law in advance of their disconnected schemes of classification, but they cannot be regarded otherwise than as ingenious and bold speculations indicating the directions in which investigations on the *rationale* of the periodic law, and on the nature of the elements, will probably be prosecuted before we arrive at anything approaching the truth, either concerning the law on the one hand or the elements on the other.

The attempt to discover some kind of unity in the sea of apparent diversities by which we are surrounded, or, rather, the natural inclination to assume such a unity, is as old as philosophy itself. Prout merely gave scientific definiteness to the views of the early Grecian philosophers when he suggested that the atomic

weights of all the elements were whole multiples of that of hydrogen, which he identified with the primary matter, and in later years many have in turn thought to find in the periodic law a warranty for resuscitating the principle underlying Prout's law, at the same time either shifting the primary matter lower down in the scale, so that hydrogen itself appears as a highly condensed form of matter, or not attempting in any way to particularise concerning the primary matter.

The application of recent developments in spectroscopy, combined with improved methods of fractionation, has resulted in the view that many of our so called rare elements at least are not strictly elemental (*v* METALS, RARE, and ELEMENTS). At first sight the large increase in the number of recognised elements which this recent work involves would seem to militate against the indications of the periodic law. But Crookes (*C J* 53, 487) has attempted to show that the new views may be brought into complete harmony with the periodic law if we will but admit into chemistry the principle of evolution. Granted that the different forms of matter have been cyclically evolved from a primitive formless fluid (protyle), and that the units of Mendeleeff's generalisation are rather 'elementary groups' than true elements, then the periodic law with its limited accommodation necessarily follows (*v* also Mills, *infra*).

Some attempts have recently been made to express the atomic weights of the elements by means of general algebraic formulæ. While it must be allowed that we are more likely to arrive at clear views regarding the periodicity of the elements through methods founded on the theory of numbers than through methods of a purely geometrical nature, yet many of the researches so far undertaken in this direction do not even promise to help towards the final solution of the periodic law. In a few cases expressions have been obtained which, as they involve constants numerically equal to some of the important constants occurring in the principles of chemistry, may on a more detailed study culminate in a clearer understanding of the nature of the elements and the rationale of the law. Mills (*P M* [5] 18, 393, 21, 151) arranges all the elements according to their atomic weights, and then subdivides this arrangement into sixteen groups. He then finds that all atomic weights are involved in the logarithmic expression $15 p - 15 (9375)^x$, in which x is an integer and p is the group number of the element considered. This expression arrived at empirically is, according to Mills, such as we should expect if we regard the elements as stable, but more or less incomplete, stages in the polymerisation of the primary matter as it cooled. It is also noteworthy that the above expression, in that it points to the existence of an upper limit to our existing system of atomic weights, confirms views originally expressed by Mendeleeff.

Carnelley's attempt (*P M* [5] 29, 97) to give a general expression to the atomic weights of the elements is of more interest in that it is founded, not on a conveniently assumed and artificial

periodicity, but on the natural periodicity as this finds expression in Mendeleeff's schemes.

Carnelley's equation is $A = c(m + \sqrt{v})$, where A = approximate atomic weight, m = a member of an arithmetical progression depending on the series to which the element belongs, v = number of the group of which the element is a member, and c is a constant.

Excluding the elements of the first three series (typical elements), the expression for the atomic weights of all the other elements may be thrown into the more convenient form

$$A = c(3\frac{1}{2}a - 9 + \sqrt{v}),$$

where a is the number of the series to which the element belongs. In his paper, Carnelley notes the following points of interest: the mean value of $c = 6.64$ is nearly identical with the mean value of the atomic heats of the elements. Assume that c in the above formula actually represents atomic heat, then it follows that the specific heats of the elements should be

given by the expression $\frac{1}{m + \sqrt{v}}$, and this is

found actually to be the case. It should be noted, however, that these coincidences may be purely accidental and meaningless, being simply the result of the particular units adopted. *D C*

PERSEITE C_8H_8O , [188°] 8 5 at 18° Mol *w* (by Raoult's method) 179 (calc 212) A body resembling mannite, occurring in the fruit and leaves of *Laurus persca* growing in the tropics (Muntz a Marcant, *C R* 99, 38, *A Ch* [6] 3, 279, Maquenne, *A Ch* [6] 15, 5, *C R* 106, 1235, 107, 583, 658, Dehérain, *C R* 108, 101, cf Avequin a Melsens, *A Ch* [2] 72, 109). Extracted by water at 60°, the extract treated with lead acetate, filtered, freed from lead by H_2S , concentrated, and mixed with $MeOH$. Formed by reduction of mannoheptose by sodium amalgam (E Fischer a Passmore, *B* 23, 2228). Small needles (from alcohol), sl sol cold, *v* sol hot water and alcohol. Its solution becomes dextro rotatory when borax is added. Does not reduce Fehling's solution or undergo alcoholic fermentation. On oxidation by nitric acid it yields mannoheptose $C_8H_{16}O$, and finally oxalic acid. Boiling HI and P give C_8H_{12} (c 104°) and C_8H_{10} (190°–200°). HCl yields C_8H_{11} (92°). An alcoholic solution treated with HCl and benzoic aldehyde yields $C_8H_{12}(CHPh)_2O$, [c 219°].

Acetyl derivative $C_8H_8(OAc)_2$, [119°] Crystalline powder, insol water, sol alcohol. Butyryl derivative $C_8H_8(O C_4H_9)_2$, (c 300° *in vacuo*) Syrup.

Nitrate $C_8H_8(NO_3)_2$, [138°] Made from perseite (1 pt), fuming HNO_3 (5 pts), and conc H_2SO_4 . White needles, insol water, sol hot alcohol. Explodes when struck.

PETROCENE The highest boiling portion of American petroleum contains, according to Hemlin (*B* 9, 1604), a hydrocarbon $C_{12}H_{22}$ [above 300°] crystallising from benzene in yellow laminae. Prunier (*A Ch* [5] 17, 28) found carbopetrocene $C_{12}H_{22}$ [268°] which yielded $C_{12}H_{18}C_6H_5N_2O$, [185°] and $C_{12}H_{18}C_6H_5N_2O$, [135°]. Prunier also found, in petroleum, petrocine $C_{12}H_{22}$? [102°] yielding $C_{12}H_{18}C_6H_5N_2O$, [98°] and a hydrocarbon $C_{12}H_{22}$? [119°]. *V* also PETROLOX.

PETROLOX. *Earth oil, Rock oil, Naphtha, Mineral oil, Erdöl, Steindl, Petrole, Huile de*

* G J Stoney has also arrived at a logarithmic expression for the atomic weights (*G J. Proc.* 1888, 68).

Pierre, Bitume liquide Petroleum is the general name given to the natural oily liquids occurring in the earth at different depths and in many localities. The natural oil is a mixture of several hydrocarbons, has a strong bituminous smell, but differs very considerably in its physical properties and chemical composition. It is found in large quantities in the United States, in the Caucasus, and in the country situated at the mouth of the Danube, in considerable quantities in Burmah and in Galicia, and has been found in Persia, the West Indian islands, Italy, parts of Germany, Switzerland, China, India, France, and England. Its specific gravity ranges from 77 to 11, and its colour from pale yellow to brown, dark green, and black. The colour of crude petroleum not only varies with the locality, but varies from time to time in the same district. B. Redwood has recorded the colours of various kinds of crude petroleum (*Journ Soc Arts*, 34, 823, 878), and Lovibond's tintometer is used for determining the colour of commercial samples. Crude petroleum varies considerably in its composition, that from North America consisting chiefly of hydrocarbons of the C_nH_{2n+2} series, while that from Baku (Caucasus) contains in addition the hydrogenised aromatic hydrocarbons of the general formula C_nH_m (Beilstein & Kurbatow, *B* 13, 1818, 2028) and small quantities of other hydrocarbons (Markownikow & Ogobine, *J R* 15, 237, 307). Galician petroleum contains hydrocarbons of the series C_nH_{2n+2} and C_nH_{2n-4} (Freund, *A* 115, 19), and also the hydrogenised aromatic hydrocarbons of the formula C_nH_m (Lachowicz, *A* 220, 168). Crude petroleum also contains small quantities of compounds containing sulphur, nitrogen, and oxygen. 0.23 per cent of nitrogen has been found in petroleum from Ohio, and 1.11 per cent in that obtained from California. Beilby has also found 0.5 per cent of nitrogen in Baku oil and 0.188 per cent in Galician ozokerite. Crude American petroleum contains at least 0.008 per cent of nitrogen (*S C I* 10, 120). Sulphur amounts to 1.87 per cent in certain Russian petroleum, and is present in that obtained from California, Lima, Ohio, and Canada, but is absent from Pennsylvanian and West Virginian oils (Vohl, *D P J* 216, 47, Woodman, *A C J* [1891] 13, 179). From the variable composition of crude petroleum it follows that its volatility will be very different, the several constituents of crude petroleum extending from the gaseous members of the marsh gas series to solids which boil at 400°. As a rule, the petroleum with lowest specific gravity is the most volatile and inflammable. Crude petroleum also may contain water and sediment, and is tested commercially for its specific gravity, odour, colour, its feel when rubbed between the fingers, and the percentage of naphtha (or portion volatile below 150°) yielded on fractional distillation (Allen, 'Commercial Organic Analysis,' 11, 869, Nawratne, *D P J* 246, 323, 423). The flashing-point and burning point of petroleum are also important factors for ascertaining its commercial value, but these tests are usually applied only to the refined petroleum or kerosene, which consists of the more volatile portions of the crude oil which can be burned with a wick (*Petroleum Act*, 1871, 84 and 85 *Visc. cap.*

105, *Petroleum Amendment Act*, 1879, and for variation of flashing point with climate *S C I* 1, 471, *C N* 40, 305, 49, 196). For calorific values of petroleum, see Deville, *C R* 72, 195, and 68, 848. Co-efficient of absorption of oxygen is higher for petroleum than for water (*Zett Phys Chem* 1, 70). When crude petroleum is distilled, the various fractions are separately collected and used for a variety of commercial purposes. In America the distillation is conducted in large stills, and the first fractions, known as 'cymogene' and 'rhigolene,' are condensed by artificial cold and pressure. The fraction which next comes over, having a density increasing from 636 to 725 or 750, is separately collected, and when re-distilled is termed 'gasoline,' 'naphtha,' and 'benzine.' The next fraction consists of oils of *S G* 75 to 84, and is known as 'kerosene' or 'photogene', while the residue is distilled in other retorts for lubricating oils, *S G* 84-9, and solid paraffin, until coke is left as a residue combustible with difficulty. Petroleum ether is the fraction containing hydrocarbons of *S G* 625-665, and consists of 'rhigolene' and 'gasolene,' while petroleum spirit or benzoline consists of the 'naphtha' and 'benzine' fractions, *S G* 68-745.

In Russia the petroleum is distilled in a series of stills in a continuous process, and yields three fractions: light benzene, *S G* 754, 'gasolene' or heavy benzene, *S G* 787, and kerosene, *S G* 825. The residue on further distillation yields (1) lubricating oils, (2) solar oil, and (3) 'astatki,' which last is either used as fuel or distilled at a red heat, yielding benzenes and anthracene. Baku petroleum contains only traces of solid hydrocarbons, while that from the Caucasus yields about 6 per cent of paraffin (*B Redwood, S C I* 4, 74).

In Germany the crude naphtha is refined into four distinct products:

Petroleum ether (40°-70°) *S G* 0.640-0.650
Benzene (70°-120°) *S G* 0.700
Ligroin (120°-135°) *S G* 0.730
Cleaning oil (130°-160°) *S G* 0.745-0.750
(Schenkel, *Chem Ind* 13, 512)

Further details on the petroleum industry, and the properties and uses of these commercial products, will be found in Thorpe's *DICTIONARY OF APPLIED CHEMISTRY*.

Early Literature—Before 1864 very little accurate knowledge of the chemical constitution of the various petroleum oils was known, the earlier investigations chiefly recording the localities in which the oil had been found, its physical properties, and its behaviour on distillation.

On the next page is a list of the more important of these examinations.

AMERICAN PETROLEUM Pelouze & Cahours (*C R* 54, 124, 56, 505, 57, 62) made a systematic investigation of the Pennsylvanian oil in 1864, and showed that it contained hydrocarbons of the formula C_nH_{2n+2} , including representatives of every member of the series from C_2H_6 to $C_{10}H_{22}$. Ronalds added C_2H_4 , C_3H_6 , and nC_4H_{10} to this list (*C J* 18, 529). Above 280°, the boiling-point of the highest of these hydrocarbons, the oil yields 3 fusible paraffins (45°-85°), and they when distilled in closed vessels were resolved into other C_nH_{2n+2} and C_nH_{2n-4} hydro-

Date	Description	Author	Reference
1788	Black Hungary	Wintere	<i>Crell's Chem Annal</i> 1, 498
1791	Galicia	Martinovich	" " 1, 72
1817	Naphtha of Miano, Parma	Saussure	<i>Bibl Univ</i> 1832, 160
1829	Persia	Unverdorben	<i>S</i> 57, 243
1831	Bangoon	Christison & Gregory	<i>T E</i> 13, 118, 124
1833	Persia	Blanchet & Sell	<i>A</i> 6, 309
1836	Bavaria	Kobell	<i>J pr</i> 8, 805
—	Galicia	Torowiewicz	<i>R P</i> 55, 15, 61, 398
—	Baku	Hess	<i>P</i> 84, 417
1840	Miano	Pelletier & Walter	<i>J Ph</i> 26, 549
1848	Derbyshire	Ure & Mansfield	<i>O J</i> 1, 249
1849			<i>Ph</i> 7, 435
1857			<i>Pr</i> 8, 221
1858			<i>A</i> 113, 151
—			<i>A</i> 114, 279
—	Galicia	Freund & Pebal	<i>A</i> 115, 19

carbons Schorlemmer found small quantities of benzene and its homologues, and Beilstein a Kurbatoff a hydrogenised aromatic hydrocarbon of the C_nH_{2n} series, which they isolated and determined to be hexahydro metaxylene.

The paraffin hydrocarbons above alluded to have also been examined in detail, and in many cases their constitution determined.

In the neighbourhood of the North American oil wells methane is evolved, and is associated with ethane in the gases which issue from the De la Mater well at Pittsburg (Sadler, *Am Ch* 1876, 98, Fouqué, *C R* 67, 1015). Ethane, propane, and n butane exist dissolved in the petroleum. Of the higher paraffins the following have been isolated.

C_5H_{12} , n pentane, dimethyl-propane,

C_6H_{14} , n hexane, ethyl iso butane,

C_7H_{16} , n -heptane and an isomeride;

either dimethyl-diethyl methane or methyl-ethyl isopropyl methane (Schorlemmer, *C J* 26, 319),

C_8H_{18} , n octane and an isomeride,

$C_{10}H_{22}$, two isomerides exist (Lemoine, *Bl* [2] 41, 164)

α (136°) SG 1.0742

β (130°) SG 1.0734

$C_{15}H_{32}$, constitution not known (160°) Pelouze & Cahours, (156°) Wurtz, *Bl* 1863, 300, (160°) SG 1.0753 Lemoine. The solid paraffins $C_{22}H_{44}$, $C_{24}H_{50}$, and $C_{26}H_{54}$ have also been separated (*V* also Greville Williams, *Tr* 1857, 737, *O J* 15, 180, Pelouze & Cahours, *A Ch* [4] 1, 5, *J* 1862, 410, Warren, *Z* 1865, 668, *J* 1868, 380, Warren & Storer, *Z* 1868, 228, Lefebvre, *Z* 1869, 185, Schorlemmer, *Tr* 182, 111, Ronalds, *Z* 1865, 523, *O J* 18, 54, *J pr* 95, 421, Wurtz, *A* 96, 372; and for the occurrence of the same hydrocarbons in boghead coal and seal oil, Schorlemmer, *A* 125, 109, Williams, *A* 125, 107, and for a comparative table showing the C_nH_{2n+2} hydrocarbons found in petroleum, or by distilling paraffin and reducing the fatty acids, Beilstein, vol i 186). Warren also obtained the olefines $C_{14}H_{28}$, $C_{16}H_{32}$, and $C_{18}H_{38}$ from American petroleum, and traces of volatile benzene hydrocarbons and the hydro benzene compounds present in Canadian oil have also been detected in this oil. Canadian petroleum contains a somewhat greater percentage of aro-

matic hydrocarbons, and is further characterised by its large amount of sulphur derivatives.

The action of heat on American petroleum products has been studied by many observers. When the crude oil is heated alone the lower members of the C_nH_{2n+2} series of hydrocarbons are evolved as gas. As the temperature increases the higher paraffin hydrocarbons distil over, but at the same time vapours are evolved which are absorbable by bromine. Le Bel examined this product, and found it to consist of bromides of the olefines C_2H_4 , C_3H_6 , C_4H_8 , C_5H_{10} , C_6H_{12} , and that there are isomeric compounds of the higher members of the series. They have been separated either by fractionating the bromides obtained in this way, or by their different behaviour towards cold HCl. Acetylene and crotonylene are also formed, and other less hydrogenised hydrocarbons. When the vapour of petroleum ether, or the fraction (50°-80°) and containing chiefly C_5H_{10} , C_6H_{12} , and C_7H_{14} , is passed through a red hot tube it yields C_2H_4 and C_3H_6 , and hydrocarbons absorbable by bromine. These, according to Prunier, are acetylenes consisting of crotonylene C_3H_4 , and traces of C_4H_6 , C_5H_8 , and C_6H_{10} .

Petroleum spirit (70°-120°) contains the hydrocarbons C_5H_{10} , C_6H_{12} , and C_7H_{14} , and the rectified oil (150°-280°) the paraffins from C_8H_{18} to $C_{16}H_{34}$, the crude fraction, however, contains hydrocarbons which are dissolved by sulphuric acid. The heavy oil (above 400°), used for lubricating purposes, has not been carefully examined. Vaseline is the next product obtained on distilling the crude oil, the distillation for heavy oil being stopped as soon as acid vapours begin to be evolved. Little is known as to the constitution of its hydrocarbons. When vaseline is not manufactured the distillation is continued until solid paraffin distils over. The solid residue, rich in carbon, also contains hydrocarbons, and in it or in the solid paraffin have been found, besides the higher paraffins, anthracene, phenanthrene, chrysene, pyrene, chrysogen (Fritzsche, *O R* 54, 910), parachrysene (Raschack), benzerythrene (Schultz), and fluoranthrene (Prunier, *Bl* [2] 81, 293). A hydrocarbon $C_{12}H_{24}$ isomeric with acenaphthalene, has also been isolated, and there are indications of the presence of quinones. From the coke Prunier and Varetine have also succeeded, by the action of solvents, in extract-

ing small quantities of hydrocarbons containing a very high percentage of carbon

From a green solid (190° - 240°) obtained from Pittsburg, and called 'petrocene' or the 'new product,' formed by the distillation of the residue after the ordinary paraffin had come over, Pruner isolated hydrocarbons which are called carbozene, carbopetrocene, and thallene. Analysis shows that they contain 96-97.7 p.c. of carbon, and have formulae, therefore, ranging from $(C_6H_4)_n$ to $(C_6H_2)_n$, where n is variable but higher than 4.

Gustavson has studied the action of Al_2Br_6 and HBr on the different fractions of petroleum ether. The fraction (67° - 70°), and chiefly hexane, gave by this treatment a solid from which he obtained an orange liquid $C_6H_5AlBr_2$, which decomposes above 120° and on addition of water it is insoluble in the hydrocarbons from which it is derived, and in CS_2 , but soluble in $EtBr$. The other fractions give similar results, but not such a good yield.

Beilstein and Kurbatoff, by the action of HNO_3 on the fraction (95° - 100°), obtained a nitro-compound $C_6H_5(NO_2)$ (195°), and soluble in KOH . The fraction (115° - 120°) similarly treated also gave acid nitro-products, called by the authors trinitro isoxylene.

Russian petroleum. The Baku oil has been the subject of much investigation, and it apparently is a more complicated mixture than the American product. Its density, according to Mendeleeff, varies from 881-886 at 15° , and its variation with temperature is given by the

$$\text{equation } \frac{d\Delta}{dt} = -[0.00635d - 0.000015d^2 - 1.44],$$

since the co-efficient of variation of density with temperature can be considered constant for any given hydrocarbon. The densities d range between 750 and 900.

Baku petroleum gives off a large volume of inflammable gas, and leaves on distillation a vaseline having all the properties of the American product save that its density is higher. The low boiling portions of the oil contain some hydrocarbons of the C_nH_{2n-2} series, and marsh gas is evolved in the neighbourhood of the Caspian Sea, but Schutzenberger and Ionnine found that the major portion of the hydrocarbons present were characterised by great inertness, and had a composition represented by the formula C_nH_{2n-2} , being isomerides of the olefine series of hydrocarbons. The hydrocarbons were called paraffenes ($C R$ 91, 823), and were found not to form addition products with Br , fuming HNO_3 , nor H_2SO_4 . They found that the vapours, when passed through a red hot tube, produced aromatic hydrocarbons, and at a dull redness gave products which unite energetically with Br , and are converted into resins by H_2SO_4 . Chlorine and a little iodine convert them into unstable chloro compounds, which cannot be distilled without decomposition. Of the several hydrocarbons present they isolated two, (221°) and (231°), and found that the latter had a V.D. corresponding to the formula C_6H_4 . Subsequent investigators have concluded that these paraffenes are the hexahydrides of the benzene series of hydrocarbons which are now called naphthenes, similar to those obtained syn-

thetically by Wreden (A . 187, 161), who described the following—

C_6H_{12} , hexahydrobenzene, S.G. 76 (69°);
 C_6H_{10} , hexahydrotoluene, S.G. 772 (97°);
 and C_6H_8 , hexahydroisoxylene, S.G. 771 (117°)
 Beilstein & Kurbatoff have found the isomeric hexahydrometaxylene (B 13, 1820, $C J$ 40, 159, also Markownikoff & Spady, B 20, 1850) in Russian petroleum, and subsequent investigations have established the presence of a series of naphthenes from C_6H_{12} to $C_{10}H_{18}$. Markownikoff (A 234, 89-115) has also obtained similar results, and has shown that the naphthenes are the chief constituents of the oil boiling below 300° . The fraction distilling (210° - 330°) under 20 mm also contains a large percentage of these C_nH_{2n-2} hydrocarbons. In addition, he isolated in the fraction (35° - 250°) the following aromatic hydrocarbons C_6H_6 , $C_6H_5(CH_3)$, C_6H_4 , isoxylene, p -xylene, C_6H_4 , pseudo cumene and mesitylene, C_6H_4 , durene, isodurene and another (diethylbenzene?), $C_{11}H_{16}$, diethyl toluene and isomerides, and hydrocarbons of the formulae $C_{11}H_{14}$ (styrene?), $C_{11}H_{12}$, $C_{11}H_{10}$, and $C_{11}H_8$. Markownikoff and Oglobine, from the fraction (210° - 330°) under 20 mm obtained evidence of the presence of oxygenated compounds of both acid and neutral characters (B [2] 41, 258), and obtained a denser naphthene $C_{11}H_{16}$. Engler has also confirmed the presence of mesitylene and pseudo cumene (B 18, 2234) in Caucasian petroleum (v also Le Bel, $C R$ 103, 1017-1019).

Aschan has isolated from the Baku oil, acids of low carbon percentage derived from hexa-, hepta-, and octo naphthenes. From the heptanaphthene carboxylic acid by P and HI he has prepared an octonaphthene (117°) under 742 mm S.G. ≈ 0.7706 , identical with that obtained by Markownikow from the Caucasian oil (B 24, 2710, and Zaloziecki, B 24, 1806). On the formation of acid compounds in petroleum v Zaloziecki, $Z f$ *Angew C* 1891 410.

The oil of still higher boiling-point consists of naphthines C_nH_{2n-2} , or hydrocarbons probably derived from the naphthenes by the high temperature, together with C_nH_{2n-4} hydrocarbons and about 10 p.c. of true benzene compounds, which are homologues of styrene. When the naphtha residues are distilled at temperatures above 400° , dissociation of the high boiling products takes place, and fresh low-boiling hydrocarbons are formed. According to Lissenko a Rosenblatt, the best yield of low-boiling products is obtained at a temperature 434° - 437° (Berg, $J R$ 1897, 349), but Nobel gives 400° as the best temperature for maximum dissociation ($D P J$ 266, 226). The Nobel brothers obtain benzene, naphthalene, and anthracene from their petroleum residues at Baku ($D P J$ 246, 429-432). Beilstein & Kurbatoff have studied the action of nitric acid upon Russian petroleum. They find that the fraction (95° - 100°) is oxidised to succinic and several volatile acids, and at the same time a nitro compound $C_6H_5(NO_2)$ (219°) is formed. The corresponding fraction of American petroleum under like treatment yields a nitro-body $C_6H_5(NO_2)$ (195°). The fraction (115° - 120°) gives crystals of α -nitro-isoxylene. Markownikoff & Oglobine isolated the above-mentioned

hydrocarbons chiefly by treating the several fractions of the oil with concentrated H_2SO_4 , and separating the sulphonates produced. About 15-20 per cent of the fraction (180° - 280°) is converted into these sulphonates. Chiefly mono and di-acids derived from the hydrocarbons C_8H_{18} , C_9H_{20} , $\text{C}_{10}\text{H}_{22}$, $\text{C}_{11}\text{H}_{24}$, and $\text{C}_{12}\text{H}_{26}$. The naphthenes are not attacked by conc H_2SO_4 , but fuming acid converts them into resins. The principal sulphonates isolated are $\text{C}_{11}\text{H}_{22}\text{SO}_3\text{H}$, 2 isomers $\text{C}_{10}\text{H}_{20}\text{SO}_3\text{H}$, $\text{C}_9\text{H}_{18}\text{SO}_3\text{H}$, $\text{C}_8\text{H}_{16}\text{SO}_3\text{H}$, $\text{C}_7\text{H}_{14}\text{SO}_3\text{H}$, and $\text{C}_6\text{H}_{12}\text{SO}_3\text{H}$. Schutzenberger has shown that a bright red heat dissociates the low boiling naphthenes into benzene derivatives, while a dull heat determines the formation of butylene, and crotonylene and its homologues. The oil obtained from Tiflis appears to be intermediate in character to that of the American and Baku products. Beilstein a Kurbatoff (*C J* 40, 1020) isolated from samples of this petroleum, n pentane, isopentane, and a butane in the (30° - 35°) fraction, a hexane and naphthenes in the (70° - 75°) fraction, and a heptane, benzene, and toluene in the (95° - 100°) fraction. They also obtained from this oil, by the action of HNO_3 , a dinitro-compound $\text{C}_8\text{H}_6(\text{NO}_2)_2$, [95°] (*v* also Le Bel, *C R* 103, 1018).

The petroleum found in Germany and Galicia is characterised by a large amount of aromatic hydrocarbons, and Kraemar & Bottcher hold that the hydrocarbons of German petroleum and Baku oil differ from coal tar and shale tar oils only in the relative proportions of those hydrocarbons which are attacked by conc H_2SO_4 and HNO_3 (*B* 20, 595-609).

Engler has made a very systematic investigation into the composition of the German oils, and has isolated or identified the following substances: gases CH_4 , CO , CO_2 , N , C_2H_4 , and homologues, in the first fraction (below 150°) C_2H_6 , C_3H_8 , and C_4H_{10} , in the fraction (160° - 182°) pseudo-cumene and mesitylene, and in the higher-boiling portion solid paraffin. The oil from Oelheimer and Wietzer contains saturated hydrocarbons, naphthenes, but no solid paraffin nor volatile products below 150° . The Tegernsee oil is especially rich in aromatic hydrocarbons (Engler, *D P J* 267, 550-570, 592-597, 268, 76-90). Engler has also detected phenols and fatty and oleic acids in many specimens. Boussingault has also isolated from Alsatian petroleum, petrole, and a black colouring matter similar to that found in Alsatian bitumen and in Galician oil (Le Bel, *Bi* 1888, 359).

GALICIAN PETROLEUM contains hydrocarbons of the $\text{C}_n\text{H}_{2n+2}$ and $\text{C}_n\text{H}_{2n-6}$ series (Freund, *A* 115, 91). It has also been examined by Lachowicz (*A* 220, 168), who finds in the low boiling fraction (80° - 125°) the following hydrocarbons: isopentane (30°), n pentane (37°), hexane (60°), n heptane (99°), ennane (148°), SG Ψ 742, two decanes (152°), SG Δ 7187 and (162°) SG Δ 7324, benzene, toluene, xylene, and mesitylene, but no olefines. From the high SG of the benzene fractions he also infers the presence of naphthenes (hexahydro toluene and hexahydro isoxylene). Pawlewski has also found 2 p.c. of aromatic hydrocarbons, principally benzene and p xylene, in Galician oil. According to Bandrowski, this

petroleum contains a small quantity of a basic body resembling the alkaloids (*M* 8, 225), and Weller has also detected the presence of bases in the yellow oil (SG 85) obtained from Saxon petroleum (*B* 20, 2098). On the Galician petroleum industry *v* Redwood, *S O I* 1892.

BURMESE PETROLEUM has only been perfectly examined Romanis has found gas, benzene, and solid paraffin in the crude oil and 5 p.c. of solid paraffin in that refined at Rangoon. It solidifies at 24° , and has a SG 85-9. From another district in Burma (Arracan) a mineral oil is obtained which contains benzene and its homologues, but does not solidify. Its SG is also lower, 825 (*C N* 59, 292).

The origin of petroleum—Mendeléeff has given considerable attention to this subject, and has advanced strong reasons for believing that mineral oils have not been produced like coal from the decomposition of past vegetation. He believes that it is formed in the depths of the earth beneath the very site on which it is found, since it cannot be water borne. The absence of any large masses of organic matter in the oil districts negatives the vegetable origin of petroleum. In Europe the oil wells belong to Tertiary and late geological periods, but in America and Canada the oil bearing sands are found in the Devonian and Silurian formations, and hence below the carboniferous beds. The oil beds also always run parallel to mountain ranges, and Mendeléeff believes that water has found its way through the fissures formed at the upheaval of these ranges to the heated metallic carbides below, resulting in the formation of metallic oxides and hydrocarbons, the chemical composition of which depends upon the conditions of temperature and pressure under which they are formed. This origin of petroleum is supported by the frequent presence of sulphur in crude oils, by the asphaltene containing a mineral ash consisting of oxides of Fe, Al, Cu, Ca (Ag), and by the non saturated nature of many of the hydrocarbons. The occurrence of petroleum in the lavas of Etna lends additional support to this theory (Silvestri, *G* 1877, 1). Hoefer, Engler, Leopold *v* Buch, Sterry Hunt, and many geologists believe that petroleum has been formed by the decomposition of organic matter of animal origin, and advance the presence of nitrogen compounds and direct experiments with animal fats in support of their view. On the origin of petroleum see *Neues Wörterbuch*, iii 39, Byasson (*M* 8 1876, 1077), Mendeléeff (*Revue Scientif* 1877, 409), Anderson (*B A* address, 1889), Engler (*B* 21, 1816-1827), Hoefer (*Mineralindustrie* in *Boiley's Technologie*), Zoloziecki (*D P J* 280, 69, 85, 138), Watson Smith (*S O I* 10, 979) S R

PETROLEUMIC ACID *v* HENDECENOIC ACID.
PETTENKOPFER'S REACTION A blood red colour on warming with cane sugar and conc H_2SO_4 . The colour is produced by the bile acids, cholic acid, and many other bodies. Furfural aldehyde may be used instead of sugar (Mylus, *H* 11, 492, *v* also vol i p 508).

PEUCEDANIN *v* IMPERATORIN.

PEWTER An alloy of Pb and Sn; *v* this vol p 125.

PHACONIN *v* PROTEINIS.

PHASEOMANNITE *v* INOSITE.

PHASOL $C_{15}H_{24}O$ [190°] $[\alpha]_D -30.6^\circ$ in a 4 p c chloroform solution. Found in the husks of peas (*Pisum sativum*) (Lukernik, *B* 24, 188). Groups of tables (from alcohol), insol water, v sol hot alcohol and ether. Gives a purple colour on shaking its chloroform solution with H_2SO_4 of SG 1.74.

PHELLANDRENE A dextrorotatory modification of this terpene occurs in oil of elemi, a levorotatory variety in Australian eucalyptus oil (Wallach, *A* 246, 234). The dextrorotatory variety occurs also in the seeds of *Phellandrium aquaticum*, and in oil of fennel (v TERPENE).

PHENACETURIC ACID $C_{11}H_{11}NO_3$, $\pm C_6H_5CH_2CO.NH.CO_2H$ [143°] S 7 at 11°. Occurs in horses' urine (Salkowski, *B* 17, 3010). Found in urine after taking phenyl acetic acid (Salkowski, *H* 7, 162). Prepared by digesting phenyl acetic anhydride with glycochol and benzene (Hotter, *J pr* [2] 38, 97, *B* 20, 84). White laminae (from water) or cubes (from alcohol). On nitration it gives *p* nitro phenaceturic acid [173°] — CaA' , 2aq S (of CaA') 3 at 11° — CuA' , aq — AgA' amorphous insoluble pp.

Methyl ether MeA' [86.5°] Needles

Ethyl ether EtA' [79°] Prisms

n Propyl ether PrA' [31°] Plates

Amide [174°] Pearly tables

PHENACONIC ACID A name given by Carius to a mixture of maleic and fumaric acids.

PHENACYL The radicle $C_6H_5.CO.CH_2$.

DI PHENACYL ACETIC ACID v DI BENZOYL-ISOBUTYRIC ACID

DI-PHENACYL-ACETOACETIC ETHER

$(CH_2Bz)_2Cac.CO_2Et$ [83°] Made from acetoacetic ether, NaOEt, and phenacyl bromide (Paal & Hoermann, *B* 22, 3225). Monoclinic crystals, sl sol cold alcohol, v sol CS. Alcoholic NH₃ at 125° forms two bodies [136°] and [192°].

Mono-oxim [63°] Flakes

Di-oxim [63°] Flakes

Tri-oxim [68°] Flakes

Di-phenyl di hydrazide $C_{18}H_{15}N_3O_2$ [88°–92°] From the ether and phenyl hydrazine

PHENACYL- α -AMIDO BENZOIC ACID

Formyl derivative

$CH_2Bz.N(CHO).C_6H_4.CO_2H$ [184°] Got by oxidising quinoline phenacylo bromide with $KMnO_4$ (Bamberger, *B* 20, 3342). Tables (from alcohol), sl sol cold water.

PHENACYLAMINE $C_6H_5.CO.CH_2.NH_2$.

ω Amido acetophenone Got by decomposing phenacyl phthalamic acid [160°] with conc HClAq (Goedeckemeyer, *B* 21, 2687). Converted by NH_3 into di-phenyl pyrazine — $B'HCl$ [188°] — $B'H_2PtCl_2$ [c 210°] — $B'C_6H_5N_2O$ [176°] Yellow needles.

PHENACYL ISOAMYL MALONIC ACID

$C_6H_5.CO.CH_2.C(OH)(CH_3)(CO_2H)$ [160°] Made from sodium isoamyl malonic ether and phenacyl bromide, the product being saponified (Paal & Hofmann, *B* 23, 1500). Needles, v sol alcohol. Yields $CH_2Bz.CH(OH)(CH_3).CO_2H$ [103°] on heating — $NH_4C_6H_5O_2$ [185°] Needles.

Ethyl ether EtA'' Oil

Amide Crystalline

PHENACYL-BENZOYL ACETIC ETHER

$CH_2Bz.CHBz.CO_2Et$ *Di-benzoyl-propionic ether* [c 58°] Made from ω -bromo acetophenone and sodium benzoyl-acetic ether (Kapf

a Paal, *B* 21, 1485, 8058). Crystals (from ether), insol water. With aqueous potash it gives benzoyl propionic acid, while alcoholic potash yields $C_6H_5C.CHBz.CO_2H$ [185°] and $CH_2Bz.CH_2Bz$ [145°].

PHENACYL BROMIDE v ω -BROMO ACETOPHENONE

PHENACYL CHLORIDE v ω -CHLORO ACETOPHENONE

PHENACYL CYANIDE v BENZOYL-ACETO

NITRILE

DI-PHENACYL-MALONIC ACID v DI BEN

ZOYL DI METHYL MALONIC ACID

PHENACYL PHTHALMIC ACID

$CO_2H.C_6H_4.CO.NH.CH_2.CO.C_6H_5$ [160°] Made by the action of alcoholic potash on phenacyl-phthalimide (Goedeckemeyer, *B* 21, 2686). Needles, insol water. Decomposed by boiling HClAq into phthalic acid and amido acetophenone.

Phenacyl-phthalimide $C_6H_5.C_6H_4.N.CH_2Bz$ [167°] Made by heating ω -bromo acetophenone with potassium phthalimide at 150° (Goedeckemeyer, *B* 21, 2685). Dimetric plates, sl alcohol and ether, almost insol water and ligroin.

Phenyl-hydrazide

$C_6H_5O.N.CH_2.C(N.HPh).C_6H_5$ [155°] Orange needles, insol water.

PHENACYL SULPHIDE $S(CH_2.CO.C_6H_5)_2$

[77°] Made by adding ω -bromo acetophenone (100 pts) in alcohol (400 pts) to a solution of Na (12 pts) in alcohol (400 pts) saturated with H_2S (Tafel & Moritz, *B* 23, 3474). Prisms (from hot alcohol). Reduces Fehling's solution. Yields a *di oxim* [161°] and a diphenylidyl-drazide [147°].

PHENACYL SULPHOCYANIDE

$C_6H_5.CO.CH_2.S.CN$ *Sulphocyanoacetophenone* [74°] Made by mixing alcoholic solutions of barium sulphocyanide and ω -bromo acetophenone (Arapides, *A* 249, 10). Needles or prisms, v sol ether, insol water. Hot HCl converts it into $CH_2Bz.S.CONH_2$ and finally oxy phenylthiazole.

PHENACYL THIOCARBAMATE

$CH.Bz.S.CONH_2$. A very unstable body got by boiling phenacyl sulphocyanide with conc HClAq until crystallisation begins (Arapides, *A* 249, 12). Yields oxy phenyl thiazole on boiling with HClAq — $B'HCl$ [c 177°] Silky needles — $B'H.PtCl_2$ [c 200°] Yellow crystalline powder.

PHENACYL TOLUIDINE v TOLYL-AMIDO-ACETOPHENONE

Di phenacyl p-toluidine $C_6H_4.Me.N(CH_2Bz)_2$ [255°] Formed from *p*-toluidine and ω -bromo acetophenone in alcohol (Lellmann & Donner, *B* 23, 168). Needles, v sol alcohol.

PHENAMYLAMINE v AMIDO-AMYL-BENZENE

Diphenamylamine $NH(C_6H_5.C_6H_4)_2$ *Di-isoamyl di phenylamine* (320°)

Formed, together with amido isoamyl-benzene $C_6H_5(C_6H_4)_2NH_2$ (260°) by heating $C_6H_5(C_6H_4)_2OH$ with ammoniacal $ZnBr_2$ or with $ZnCl_2$ and NH_4Br or NH_4Cl , the yield being 18 to 25 p c (Lloyd, *B* 20, 1257) — $B'H_2PtCl_2$. *Acetyl derivative* $NaAc(C_6H_4)_2$ [81°]. White glistening plates.

PHENANTHRAQUINONE $C_{14}H_8O$, \pm

$C_6H_5.CO$ or $C_6H_5.CO$ Mol. w. 208 [203°]

(Hayduck, *A* 167, 184) (above 860°) Formed by oxidising the residue left after evaporating the alcoholic washings from the distillate got in preparing anthracene from α -bromo benzyl bromide and Na (Jackson & White, *Am* 2, 392)

Preparation—1 By warming phenanthrene (1 pt) with $K_2Cr_2O_7$ (1 pt), water (8 pts) and H_2SO_4 (1½ pt) and crystallising the product from diluted HOAc (Fittig & Ostermayer, *B* 5, 933, *A* 166, 365)—2 By adding a solution of CrO_3 in HOAc to a hot solution of phenanthrene in HOAc (Graebe, *B* 5, 861, *A* 167, 139) The product is purified by solution in aqueous $NaHSO_4$, followed by ppn with HCl and crystallisation from HOAc—3 By oxidising crude phenanthrene (310°–340°), the following bodies being obtained at the same time, methyl anthraquinone, diphenic acid, carbazole, acridine, and diphenylene-ketone (Anschütz & Schultz, *A* 196, 32)

Properties—Orange needles or prisms (from solvents) or tables (by sublimation), almost insol cold water, sol hot water and cold alcohol, m sol hot alcohol, ether, HOAc, and benzene Conc H_2SO_4 forms a dark green solution but does not sulphate it, even at 100° A solution of phenanthraquinone (1 g) in HOAc (20 cc) mixed with toluene (4 cc) containing methyl-thiophene gives on shaking with H_2SO_4 (16 cc) a bluish-green liquid which, when poured into water and extracted with ether, imparts a purple colour to the ether (Laubenheimer, *B* 8, 224, *V* Meyer, *B* 16, 1624) The colouring matter $C_{12}H_{10}SO$ formed in this reaction gives anthraquinone on distilling with $PbCrO_4$ (Oderheimer, *B* 17, 1338) A solution of phenanthraquinone in wet ether when exposed in closed tubes to direct sunshine is readily reduced to phenanthrahydroquinone, with formation of aldehyde (Klinger, *B* 19, 1869)

Reactions—1 Oxidised by chromic acid to diphenyl di- α -carboxylic acid—2 Alkaline $KMnO_4$ gives oxy-diphenylene acetic acid and diphenylene-ketone (Anschütz & Japp, *B* 11, 212)—3 Reduced to dihydride by warming with aqueous SO_2 —4 Sodium-amalgam acting on its alcoholic solutions forms diphenic acid [228°] (*A* & *S*)—5 Boiling aqueous NaOH yields $C_{12}H_8$ $C(OH)CO_2H$ (*A* & *J*) Boiling baryta-water forms, besides oxy-diphenylene acetic acid, diphenylene ketone and fluorene alcohol—6 Alcoholic potash gradually forms diphenic acid, the solution meanwhile giving out phosphorescent light on shaking (Lachovitch, *B* 16, 332)—7 By passing the vapour over heated lead oxide di-phenylene ketone is formed (Wittenberg & Meyer, *B* 16, 502)—8 Distillation with dry soda-lime gives diphenyl When the soda-lime is moist, fluorene, fluorene alcohol, and diphenylene ketone are also formed—9 Distillation with quick-lime gives fluorene and diphenylene ketone (Anschütz & Schultz, *B* 9, 1400)—10 Distillation with zinc-dust yields phenanthrene—11 An ethereal solution of $ZnEt_2$ decolorises it On adding alcohol, boiling, and filtering, crystals of $C_{12}H_{10}O_2HOEt$, (77°), are formed. These are rectangular plates, insol water Ac_2O converts it into $C_{12}H_{10}O_2Ac$, [108°] (Japp, *C* *J* 85, 526)—12 With PCl_5 it gives $C_{12}H_8$ CCl_2 , C_6H_5 CO , benzene may be used as diluent. The

product 'di-chloro-phenanthrone' may be re-crystallised from benzene It melts at [166°], although at 140° it begins to get brown Alkalis convert it into phenanthraquinone Shaken with alcoholic potash, it is oxidised to diphenic acid, the solution phosphorescing meanwhile Iron and acetic acid reduce it first to chloro-phenanthrone, $C_{12}H_8$ $CHCl$ [128°], and then to

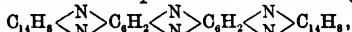
phenanthrone $C_{12}H_8$ CH_2 , [149°] Chloro-

phenanthrone may be recrystallised from glacial acetic acid, and is not decomposed even by boiling alkalis, but HNO_3 (S G 13) converts it into nitro-phenanthraquinone Phenanthrone may be crystallised first from glacial acetic acid, then from butyl bromide (B Lachovitch, *J* *pr* [2] 28, 168)—18 By exhaustive chlorination with $SbCl_5$ it yields perchloro-diphenyl together with a very small quantity of per chloro benzene (Merz & Weith, *B* 16, 2870)—14 Benzoic aldehyde (free from HCl) at 260° forms benzalquin of phenanthrene $C_{12}H_8O$ [329°], which crystallises from CS_2 in rectangular plates and yields benzoic acid and phenanthraquinone on oxidation It gives off no gas with $ZnEt_2$, hence contains no hydroxyl (Japp & Wilcock, *C* *J* 37, 661)—15 Heating with PCl_5 followed by addition of water forms $C_{12}H_8$ CO $C(OH)PO(OH)_2$, crystallising

in flesh coloured plates, sol water, alcohol, and ether (Fossek, *M* 7, 36)—16 Acetone at 200° forms acetone phenanthraquinone $C_{12}H_8O_2$ (vol 1 p 33) By the action of H_2SO_4 on this body a small quantity of $C_{12}H_8O_2$ [238°] is formed (Wadsworth, *C* *J* 59, 105)—17 Ammonia passed into an alcoholic solution forms phenanthraquinonimide $C_{12}H_8NO$ On heating with alcoholic NH_3 in sealed tubes there are formed diphenanthrylene azotide $C_{12}H_8N_2$, two compounds $C_{12}H_8N_2O$ [282°] and [over 300°] and a compound $C_{12}H_8N_2$ [above 285°] (Schmidt, *B* 7, 1365, Anschütz & Schultz, *A* 196, 49, Zincke, *B* 12, 1641, Sommaruga, *M* 1, 146, Japp, *C* *J* 49, 845, 51, 98)—18 Benzoic aldehyde and aqueous NH_3 at 100° quickly forms $C_{12}H_8NO$ or $C_{12}H_8$ O_2 $\langle N \rangle \supset CPh$

This 'benzenyl amido phenanthrol' crystallises from benzene in tufts of silky needles, [202°], v sl sol alcohol, sol conc $HClAq$ and conc H_2SO_4 without change It yields benzoic acid and phenanthraquinone on oxidation $HClAq$ at 200° forms benzoic acid It does not act on $ZnEt_2$ (Japp, *C* *J* 87, 666, 89, 925)—19 Cumic aldehyde and NH_3 forms, in like manner, cumenyl-amido-phenanthrol $C_{12}H_8$ O_2 $\langle N \rangle \supset C_6H_5$ C_6H_5 , [186°], crystallising from benzene in silky needles, and forming in conc H_2SO_4 a yellowish green fluorescent liquid (Japp & Wilcock, *C* *J* 89, 226)—20 Furfuraldehyde and aqueous NH_3 , give $C_{12}H_8$ O_2 $\langle N \rangle \supset C_6H_5O$ [231°], crystallising from isomyl alcohol in needles (*J* & *W*)—21 Salicylic aldehyde (1 mol) and conc NH_3Aq form, on warming, an oxy benzenyl di-amido-phenanthrene $C_{12}H_8$ CNH $C-N \supset C_6H_5$ OH (Japp & Streatfield, *C* *J* 41, 146) This body crystallises from HOAc in slender needles [270°–

276°], and is sl sol alcohol, v sol HOAc. It dissolves in boiling KOHAq, and is pppd by CO₂. Boiling alcoholic potash gives salicylic acid. HClAq at 200° has no action. BzCl yields a benzoyl derivative [220°]—22 *o-Methoxy-benzonic aldehyde* (15 g) heated with phenanthraquinone (30 g) and excess of NH₄Aq at 100° deposits yellow crystals of C₁₄H₈N₂H C₆H₄OMe [208°], while the mother liquor contains C₁₄H₈<N>^OC₆H₄OMe, which crystallises in white needles, v sol hot benzene (Japp a Streatfield, *C J* 41, 154)—23 *p-Oxy benzoic aldehyde* and aqueous 1° H₂ form *p* oxy benzenyl-di amido phenanthrene [ab. ca 350°], crystallising from HOAc in slender needles. It yields an acetyl derivative [205°–210°] (J a S)—24 *Aldehyde ammonia* yields an amorphous base (J a W)—25 *Alcoholic methylamine* forms, on warming, yellow crystals. The mother liquor deposits C₁₄H₈N₂ or C₁₄H₈(NMe)₂, crystallising in colourless prisms [186°], and yielding B'HCl, v sol water, and B'HNO₃, B'H₂SO₄, B'H₂O₂, all crystallising in needles (Zincke, *B* 12, 1643)—26 *Ethylene diamine* and HOAc form the azine C₁₄H₈<N>^NC₂H₄, which crystallises from alcohol in needles [181°]. It yields a platinochloride B'H₂PtCl₄, nearly insol alcohol (Mason, *B* 19, 112, 20, 268)—27 *Propylene diamine* yields C₁₄H₈N₂, crystallising in needles [128°], and yielding B'H₂PtCl₄ (Strache, *B* 21, 2362)—28 *Trimethylene diamine* forms C₂₂H₁₆NO₂, a lemon yellow powder, not melted at 250°. Its alcohol solution is coloured violet by acids—29 *Tetra amido phenazine* and HOAc give

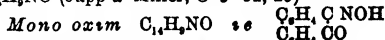


a green crystalline pp, forming a bluish green solution in H₂SO₄, changing on dilution through red to orange (Nietzki a Muller, *B* 22, 450)—30 *Phenylene o-diamine* gives phenanthrazine C₁₄H₈<N>^NC₆H₄, [217°], insol water, v sol alcohol (Hinsberg, *A* 237, 340)—31 *Tolylene-o-diamine* forms C₁₄H₈<N>^NC₆H₃Me [213°]—32 Heating with *acetamide* and HOAc forms the azine C₁₄H₈<N>^NC₆H₄, crystallising in yellowish brown flat needles, [400°], sol aniline, nitro benzene, and phenol (Mason, *C J* 55, 108)—33 *Naphthylene (1,2)-diamine* yields C₁₄H₈<N>^NC₁₀H₆ [264°] (Leuckart, *B* 19, 174)—34 *Phenyl-naphthylene-(1,2)-diamine* boiled with HOAc and phenanthraquinone forms, on adding HNO₃, a pp of C₂₆H₁₈(NO)₂N₂O, whence boiling potash separates C₁₄H₈<N>^NC₁₀H₆ (NPh(OH)) which forms a yellow solution in ether. It forms a blue solution in H₂SO₄, turned orange by dilution (Witt, *B* 20, 1185)—35 *Nitro-o-phenylene-diamine* and HOAc give, on heating, C₂₆H₁₈N₂O₂ [261°] (Heim, *B* 21, 2301)—36 *Isobutyl phenylene-diamine* in HOAc forms C₁₄H₈<N>^NC₆H₄C₂H₅ [147°], crystallising in pale-yellow needles, coloured cherry red by H₂SO₄ (Gelzer, *B* 20, 3253, 21, 2951). The compound C₁₄H₈N₂C₆H₃BrC₂H₅ [154°] also

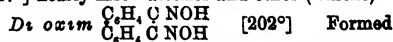
forms yellow needles (B) *Isobutyl phenylene-diamine* gives an isomeric azine [144°]—37 (6,4,2,1) *Bromo tolylene diamine* in HOAc forms C₁₄H₈N₂C₆H₄MeBr [210°] crystallising in yellow needles (Hartmann, *B* 23, 1050)—38 *1-Tri amido-benzene* (from chrysoidine) yields C₁₄H₈N₂C₆H₄NH₂, as brown crystals [179°] (Witt, *C J* 49, 402, Heim)—39 *Acetoacetic ether* forms, in presence of KOH, phenanthroxylene acetoacetic ether C₁₄H₈EtO, [185°] crystallising in white needles (Japp a Streatfield, *C J* 43, 27)—40 *Acetone* and NH₄Aq form C₁₄H₈NO₂, crystallising in colourless laminæ [c 130°]. This body forms a yellow solution in HClAq, which deposits a dark blue substance and gives, on addition of water to the filtrate, needles of C₁₄H₈O₂ (Japp a Streatfield, *C J* 41, 272)

Combinations—C₁₄H₈O₂NaHSO₄ 2aq. Small colourless plates, v e sol water, which slowly decomposes it—(C₁₄H₈O₂)₂HgCy₂ [223°] Red crystals with green lustre, deposited from a hot saturated solution of phenanthraquinone and HgCy₂ in acetone (Japp a Turner, *C J* 57, 7)—(C₁₄H₈O₂)₂HgCl₂ [223°] Red prisms (from boiling acetone)—C₁₄H₈O₂ZnCl₂. Dark reddish-brown needles got by adding a hot solution of ZnCl₂ in HOAc to a hot solution of phenanthraquinone in HOAc (Japp a Turner, *C J* 57, 5)

Hydrocyanide C₁₄H₈O₂(HCN), tufts of needles, formed by action of 80 p c HCN solution. Decomposed by heat into its components. Conc HCl decomposes it, giving off CO₂ and forming C₁₄H₈NO [241°] and C₁₄H₈NO₂ [183°]. The latter separates from benzene in slender needles. Both dissolve in sodium carbonate, expelling CO₂, the compound [241°] producing C₁₄H₈NaNO₂ 4aq, and also the corresponding (C₁₄H₈NO)₂. Ba7aq whence HCl liberates C₁₄H₈NO (Japp a Miller, *C J* 51, 29)



[158°] Formed by boiling phenanthraquinone with alcoholic hydroxylamine hydrochloride for an hour (Goldschmidt, *B* 16, 2178). Small yellow needles, v sol hot alcohol. Forms coloured pps with metallic salts (Kostanecki, *B* 22, 1347). Boiling NaOHAq forms a green liquid. Conc H₂SO₄ forms a blood red solution and at 100° converts it into diphenylene ketone carboxylic amide HCl, Ac₂O, and HOAc at 100° convert it, by intra molecular change, into a weak base [217°], probably $\begin{smallmatrix} C_6H_4 \\ C_6H_4 \end{smallmatrix} \begin{smallmatrix} \text{CO} \\ \text{CO} \end{smallmatrix} \text{NH}$, and also form prisms [92°] (Wegerhoff, *A* 252, 17). Ac₂O forms C₂₆H₁₈N₂O crystallising in small brownish tables [247°] nearly insol alcohol and ether (Zincke)



by heating an alcoholic solution of phenanthraquinone (1 mol) with hydroxylamine hydrochloride (4 mols) for 30 hours at 100° (Auwers a V Meyer, *B* 22, 1993). Minute yellow prisms (from alcohol or HOAc), insol water, sl sol hot alcohol and ether. Conc H₂SO₄ forms a blood-red solution. NaOHAq forms a yellow solution, depositing the Na salt as pearly plates. Alcohol at 150° forms the anhydride $\begin{smallmatrix} C_6H_4 \\ C_6H_4 \end{smallmatrix} \begin{smallmatrix} \text{C} \\ \text{C} \end{smallmatrix} \begin{smallmatrix} \text{N} \\ \text{N} \end{smallmatrix}$ [183°] which crystallises in long yellow needles, insol. NaOHAq. A solution of the dioxim in HOAc

and Ac_2O saturated with HCl in the cold gives the acetyl derivative $\text{C}_{14}\text{H}_8\text{O}(\text{NOAc})_2$ separating from alcohol in minute crystals [184°]

Phenyl-hydrazide $\text{C}_{20}\text{H}_{14}\text{N}_2\text{O}$ [165°] Formed by warming an alcoholic solution of phenanthraquinone with aqueous phenyl-hydrazine hydrochloride (Zincke, *B* 16, 1564) Red needles or plates (from alcohol) Gives a violet solution in H_2SO_4

Di bromo-phenanthraquinone $\text{C}_{14}\text{H}_8\text{Br}_2\text{O}_2$ [230°] (*H*), [238°] (Ostermayer, *B* 7, 1090) Made by heating the quinone with Br and a little water for six hours at 180° (Hayduck, *A* 167, 185) Yellow nodules (from HOAc), sl sol alcohol Yields di bromo diphenyl dicarboxylic acid on oxidation by chromic acid mixture

Nitro-phenanthraquinone

[1 2] $\text{C}_{14}\text{H}_7(\text{NO}_2)_2\text{O}$ Formed from the quinone and boiling HNO_3 (S G 14) (Anschütz a Schultz, *B* 9, 1404, Strasburger, *B* 16, 2346) Orange plates (from HOAc) yields nitro diphenyl dicarboxylic acid [217°] on oxidation By oxidation of (α), (β), and (γ) nitro phenanthrene with CrO_3 and HOAc there are formed (α), (β), and (γ) nitro-phenanthraquinones [215° - 220°], [260° - 266°], and [263°] respectively (Schmidt, *B* 12, 1156)

Nitro-phenanthraquinone [282°] Made by warming chloro-phenanthrone with nitric acid (S G 13) (Lachovitch, *J pr* [2] 28, 172) Orange plates, sl sol HOAc

Di nitro phenanthraquinone

$\text{C}_{14}\text{H}_6(\text{NO}_2)_4\text{O}$ [294°] Made by boiling phenanthrene with fuming HNO_3 or with a mixture of HNO_3 and H_2SO_4 Obtained also, together with a more soluble isomeride, by nitration of phenanthraquinone and of nitro phenanthraquinone [257°] (Graebe, *A* 167, 144, Schultz, *A* 203, 108, Strasburger, *B* 16, 2346) Yellow plates, v sl sol alcohol Yields di nitro diphenyl dicarboxylic acid [258°] on oxidation

Amido-phenanthraquinone

[1 2] $\text{C}_{14}\text{H}_8(\text{NH}_2)_2\text{O}$ Formed by reduction of nitro-phenanthraquinone [257°] with tin and HCl (Anschütz a Meyer, *B* 18, 1943) Violet-black needles, sl sol hot water, forming a reddish violet solution— B^*HCl yellowish red feathery needles

Di amido phenanthraquinone

[1 4 2] $\text{C}_{14}\text{H}_6(\text{NH}_2)_4\text{O}$ Formed by reduction of the di nitro-compound with tin and HCl (A a M., Kleemann a Wense, *B* 18, 2168) Violet black needles, not melted at 310° The hydrochloride forms yellow plates.

Oxy-phenanthraquinone $\text{C}_{14}\text{H}_8\text{O}_2$ Formed by oxidation of phenanthrolquinone

Formed by the action of nitrous acid on amido phenanthraquinone Brownish red needles May be sublimed. Forms a yellowish-green solution in NaOH aq

Acetyl derivative

[200°-210°] **Di-oxy-phenanthraquinone** $\text{C}_{14}\text{H}_6(\text{OH})_2\text{O}_2$ Formed by the action of nitrous acid on di-amido-phenanthraquinone (A a M.) Minute dark-brown needles Yields a di acetyl derivative crystallising in yellowish-red needles.

Hydro phenanthraquinone $\text{C}_{14}\text{H}_{10}\text{O}_2$ $\text{C}_{14}\text{H}_8\text{O}_2$ $\text{C}_6\text{H}_5\text{COH}$ *Phenanthraquinone dihydride* *Phenanthrenehydroquinone* Mol w 210 Formed by heating phenanthraquinone with aqueous SO_2 , or by passing SO_2 into a warm alcoholic solution of the quinone (Graebe, *A* 167, 146) Colourless needles, m sol hot water, v e sol alcohol, ether, and benzene Absorbs oxygen when moist, or in aqueous solution, forming the quinhydrone $\text{C}_{22}\text{H}_{18}\text{O}_2$, which crystallises in black needles [169°], and finally phenanthraquinone FeCl_3 , HNO_3 , and CrO_3 oxidise it in the same way

Mono acetyl derivat

$\text{C}_{14}\text{H}_9(\text{OH})(\text{OAc})$ [170°] Got by boiling phenanthraquinone with HOAc , HI , and red P (Japp a Klingemann, *C J Proc* 6, 31) Formed by the action of sunlight on phenanthraquinone in aldehyde (Klinger, *A* 249, 138) Flat needles

Di-acetyl derivative

$\text{C}_{14}\text{H}_8(\text{OAc})_2$ [202°] Formed from the dihydride and Ac_2O Colourless plates (from benzene), not oxidised by boiling chromic acid mixture Not attacked by heating with KOH aq, unless the solution has a higher SG than 1.3

Benzoyl derivative

$\text{C}_{14}\text{H}_9(\text{OH})(\text{OBz})$ [178°] Got by action of sunlight on phenanthraquinone and benzoic aldehyde (K) White needles (from HOAc)

Valeryl derivative

$\text{C}_{14}\text{H}_9(\text{OH})(\text{O C}_4\text{H}_7\text{O})$ [149°] Got by using isovaleric aldehyde

Ethyl derivative

$\text{C}_{14}\text{H}_{10}(\text{OH})(\text{OEt})$ [80°] On treatment of phenanthraquinone with ZnEt_2 and alcohol successively the compound $\text{C}_{14}\text{H}_{11}\text{O}_2\text{EtOH}$ [77°] is got, from which EtOH can be removed by standing for some months *in vacuo* over H_2SO_4 (Japp, *C J* 37, 408) Gives a mono acetyl derivative $\text{C}_{14}\text{H}_9\text{AcO}$ [103°]

Isomeride of hydro phenanthraquinone

$\text{C}_{14}\text{H}_8(\text{OH})_2$ [143°] Got by heating its diacetyl derivative with alcoholic NH_3 (Fischer a Gerichten, *B* 19, 792) Needles, very oxidisable, its alkaline solution turning green and finally red

Acetyl derivative

$\text{C}_{14}\text{H}_9(\text{OAc})_2$ [159°] Made by boiling morphine methyl iodide with Ac_2O , adding dry AgOAc , filtering, and heating the filtrate at 180° Needles (from ether)

Di amido-hydro phenanthraquinone

$\text{C}_{14}\text{H}_8(\text{NH}_2)_2(\text{OH})_2$ Formed by reduction of di nitro phenanthraquinone with SnCl_2 (Kleemann a Wense, *B* 18, 2168) Very readily oxidised by air, FeCl_3 , or CrO_3 to violet black needles of di-amido phenanthraquinone— $\text{B}^*\text{H}_2\text{Cl}_2$ 3aq

Tetra-acetyl derivative

$\text{C}_{14}\text{H}_6(\text{NHAc})_4(\text{OAc})_2$ Colourless needles, solid at 800° , sl sol alcohol and HOAc

Phenanthraquinone carboxylic acid

$\text{C}_{14}\text{H}_7(\text{CO}_2\text{H})\text{O}$ [315°] Made by oxidation of phenanthrene carboxylic acid with CrO_3 in HOAc (Japp a Schultz, *B* 10, 1861, *A* 196, 14) Orange substance, sol NaHSO_4 aq

Phenanthraquinone sulphonic acid

$\text{C}_{14}\text{H}_7(\text{SO}_3\text{H})\text{O}_2$ Formed from phenanthraquinone and SO_3 (Graebe) Gives a colouring matter resembling alizarin when fused with potash.

Phenanthrone $\text{C}_{14}\text{H}_8\text{O}$ [149°] Formed as above (Reaction 12). Brownish red plates, v.

sol alcohol and ether, does not combine with NaHSO_4 . Its alkaline solution is green

Diphenanthrylene azotide $\text{C}_{14}\text{H}_8\text{N}_2\text{C}_{14}\text{H}_8$ [above 400°] Formed from phenanthraquinone and alcoholic NH_3 at 100°, and also by heating *terphenyl* zinc with soda lime to a red heat (Japp & Bun. on, *C J* 49, 845, 51, 98) Yellow crystalline powder or yellow needles (by sublimation), v sol ether Its solution in H_2SO_4 is deep blue, becoming orange on dilution

Isophenanthraquinone $\text{C}_{14}\text{H}_8\text{O}_2$ [156°] Formed by the further action of CrO_3 and HOAc on an oil formed in the oxidation of phenanthrene, which remains in the alcoholic liquid from which phenanthraquinone has separated (Hayduck, *A* 167, 185) Yellow crystals (from alcohol), sl sol water, v sol hot alcohol

PHENANTHAZINE v PHENANTHRAQUINONE, Reaction 29

PHENANTHRENE $\text{C}_{14}\text{H}_{10}$ *ts*

[1 2] $\text{C}_{14}\text{H}_{10}$ CH Mol w 178 [100°] (Graebe, Schiff), [103°] (Reissert, *B* 23, 2244) (340° i V) S (alcohol) 2 62 at 16°, 10 08 at 78° S (toluene) 33 at 16 5° (Bechi, *B* 12, 1978) S V S 167 05 (Schiff) S V 186 2 (Lossen, *A* 254, 54), 196 7 (Ramsay) H.F (from diamond) -39,400 (Berthelot & Vieille, *A Ch* [6] 10, 446, *Bl* [2] 47, 864), -32,500 (Stohmann, *J pr* [2] 40, 94) H C v 1,699,000 (*B A V*), 1,692,000 (*S*) H C p 1,700,400 (*B A V*), 1,693,500 (*S*) Occurs in coal tar c l (Graebe, *B* 5, 861, *A* 167, 181, Fittig & Ostermayer, *B* 5, 933, *A* 166, 361, Hayduck, *B* 6, 532, *A* 167, 177) and in 'idryl' got by distillation of an Idrian ore of mercury

Formation.—1 Together with toluene by passing s di phenyl ethylene through a red hot tube (Graebe) —2 From s di phenyl ethane, from toluene (Graebe, *B* 7, 49), from a mixture of diphenyl and ethylene, from di methyl di phenyl, and from phenyl tolyl methane (Barbier, *C R* 79, 121) by passing the vapours through red hot tubes Other hydrocarbons are also formed in these reactions —3 Together with anthracene in the action of sodium on [1 2] $\text{C}_6\text{H}_5\text{Br} \cdot \text{CH}_2\text{Br}$ (Jackson & White, *Am* 2, 391) —4 By heating coumarone and benzene to a high temperature (Kraemer & Spilker, *B* 23, 85)

Preparation—By fractional distillation of the portion of hydrocarbons (310°-360°) from coal tar oil, followed by crystallisation from alcohol A solution of crude phenanthrene (3 pts) in hot toluene mixed with picric acid (1 pts) deposits on cooling the picric acid compound in golden needles, which may be subsequently decomposed by alkalis The mixture of phenanthrene and anthracene may also be crystallised from toluene, when anthracene separates first (Wense, *B* 19, 761) Chromic acid mixture attacks anthracene more readily than phenanthrene, so that a small quantity of anthracene may be removed by its means (Anschutz & Schultze, *A* 196, 35)

Properties—Small colourless plates (from alcohol), sl sol cold alcohol, v sol ether, benz, ne, HOAc , and CS_2 May be sublimed Exhibits slight blue fluorescence Its absorption spectrum in the ultra-violet has been studied

by Hartley (*C J* 39, 164) Not reduced in alcoholic solution by sodium amalgam

Reactions—1 Oxidised by chromic acid mixture and by CrO_3 in HOAc to phenanthraquinone —2 HIAq and P at 200° form a tetrahydride —3 Bromine added to its solution in ether or CS_2 forms unstable $\text{C}_{14}\text{H}_8\text{Br}_2$, crystallising in four sided prisms (*F A O*, Hayduck, *A* 167, 180) This dibromide melts with effervescence at 98°, and its alcoholic solution gives a pp of AgBr on adding AgNO_3 By alcoholic KCy it is reconverted into phenanthrene (Anschutz, *B* 11, 1217) —4 By exhaustive chlorination with SbCl_5 it yields per-chloro benzene (Merz & Weith, *B* 16, 2869)

Combinations— $\text{C}_{14}\text{H}_{10}\text{C}_6\text{H}_5\text{N}_3\text{O}$, [143°] S (95 p.c alcohol) 2 7 at 15° Golden prisms — $\text{C}_{14}\text{H}_{10}\text{C}_6\text{H}_5\text{Cl}(\text{NO}_2)_2$ [1 2 4] [44°] Orange needles (Willgerodt, *B* 11, 604) — $\text{C}_{14}\text{H}_{10}\text{C}_6\text{H}_5\text{Cl}(\text{NO}_2)_2$ [88°] Lemon yellow needles (Liebermann, *B* 8, 378)

Tetrahydride $\text{C}_{14}\text{H}_{14}$, [0°] (810° i V) SG 19 1 067 Got by reducing phenanthrene with HIAq and P, or by isomyl alcohol and sodium (Bamberger & Lodter, *B* 20, 3076) Liquid, v sol hot alcohol Yields phenanthraquinone on oxidation by CrO_3 and HOAc

Octohydride $\text{C}_{14}\text{H}_{18}$ (below 300°) Got by heating phenanthrene with HIAq and P at 240° (Graebe) Liquid

Perhydride $\text{C}_{14}\text{H}_{24}$, [-8°] (270°-275°) SG 2° 933 Made, together with a hydride (290°), by heating phenanthrene with HIAq and red P at 250° (Liebermann & Spiegel, *B* 22, 779) Yields phenanthrene and anthracene when distilled with zinc dust. Not attacked by H_2SO_4 , HNO_3 , or Br

Chloro phenanthrenes By passing chlorine into a solution of phenanthrene in HOAc the compound $\text{C}_{14}\text{H}_9\text{Cl}$, [170°] is formed, together with oily $\text{C}_{14}\text{H}_9\text{Cl}$ and $\text{C}_{14}\text{H}_9\text{Cl}_2$ Alcoholic potash converts $\text{C}_{14}\text{H}_9\text{Cl}$ into $\text{C}_{14}\text{H}_8\text{Cl}$ By heating phenanthrene with SbCl_5 , the compounds $\text{C}_{14}\text{H}_8\text{Cl}$, [172°], $\text{C}_{14}\text{H}_7\text{Cl}_2$, [250°], and $\text{C}_{14}\text{H}_6\text{Cl}_3$, [270°-280°] may be obtained (Zetter, *B* 11, 165), the final products being C_6Cl_6 and CCl_4

Bromo phenanthrenes Bromine acting on phenanthrene dissolved in ether forms $\text{C}_{14}\text{H}_9\text{Br}_2$, which at 100° splits up into HBr and $\text{C}_{14}\text{H}_8\text{Br}$, $\text{C}_{14}\text{H}_7\text{Br}_2$

[63°], (above 360°) Bromo phenanthrene crystallises in thin prisms, v sol HOAc and CS_2 , and is converted by oxidation into phenanthraquinone (Zetter, Hayduck, Anschutz) Bromine (4 mols) acting on phenanthrene in ethereal solution also forms two di-bromo phenanthrenes $\text{C}_{14}\text{H}_8\text{Br}_2$, [148°] and [156°] A third di-bromo-phenanthrene [202°] is a crystalline powder insol. ether By heating phenanthrene with bromine the compounds $\text{C}_{14}\text{H}_8\text{Br}_2$, [126°] and $\text{C}_{14}\text{H}_7\text{Br}_3$, [185°] may be got By heating phenanthrene with bromine and iodine the compounds $\text{C}_{14}\text{H}_8\text{Br}_2$, [245°] and $\text{C}_{14}\text{H}_7\text{Br}_3$, [above 270°] are obtained All these bromo-phenanthrenes crystallise in needles

(a) - Nitro - phenanthrene $\text{C}_{14}\text{H}_9\text{NO}_2$, [75°]. Made, together with the two following isomerides, by nitration of phenanthrene (Schmidt, *B* 12, 1153) Yellow needles Gives (a) nitro phenanthraquinone [215°-220°] on oxidation.

(β) Nitro-phenanthrene. [127°] Less soluble than the (α)-isomeride. Yields nitro-phenanthraquinone [260°-266°] on oxidation

(γ)-Nitro-phenanthrene [171°] Small yellow leaflets. Less sol. alcohol than either of its isomerides. Yields on oxidation a nitro-phenanthraquinone [268°]

Di-nitro-phenanthrene $C_{14}H_8(NO_2)_2$. [150°-160°] Made by prolonged action of HNO_3 on phenanthrene (Graebe). Yellow crystals

Bromo-nitro-phenanthrene $C_{14}H_7Br(NO_2)$. [196°] Made by nitration of bromo-phenanthrene (Anschütz, *B* 11, 1218). Long spikes

(α)-Amido-phenanthrene $C_{14}H_9NH_2$. Got by reduction of (α)-nitro-phenanthrene (Schmidt, *B* 12, 1156). Small leaflets, insol. water — $BHCl$ crystalline pp — $B'H_2SO_4$ powder

(β) Amido-phenanthrene. Small leaflets — $BHCl$ sl sol. water

(γ)-Amido-phenanthrene. Got, like the preceding bodies, by reduction of the corresponding nitro-phenanthrene — $BHCl$ glistening needles

Oxy-phenanthrene *v* Phenanthrol

Di-oxy-phenanthrene *v*. Hydro-phenanthraquinone

Pseudophenanthrene $C_{14}H_{12}$. [115°] Occurs in crude anthracene (Zeidler, *A* 191, 295). Large white plates, yielding on oxidation a quinone [170°] which is v sol. alcohol and benzene. The picric acid compound [147°] separates in bright-red needles on mixing saturated alcoholic solutions of pseudophenanthrene and picric acid

A hydrocarbon $C_{14}H_{10}$ or $C_{14}H_{12}$. [104°], possibly a mixture of phenanthrene with pseudophenanthrene, was got by Zeidler (*A* 191, 292) from crude anthracene. It yields an orange picrate [142°], a quinone [205°-209°], insol. $NaHSO_4$ (difference from phenanthraquinone), and a bromide [95°] which changes on fusion to a compound [83°], and on boiling with alcoholic potash forms yellow needles [250°]. The quinone gives Laubenheimer's reaction

(α)-PHENANTHRENE CARBOXYLIC ACID CO_2H $C_{14}H_7$ CH $C_{14}H_7$ CH [266°] Formed by the action

of alcoholic potash on the nitrile which is got by distilling sodium phenanthrene (α) sulphonate (2 pts) with K_2FeCy_4 (3 pts) (Japp & Schultz, *B* 10, 1861, *C* *J* 37, 86). Colourless curved blades (from $HOAc$) or fern shaped leaves (by sublimation), almost insol. water. Yields phenanthraquinone carboxylic acid on oxidation by CrO_3 and $HOAc$ — NaA' 4aq. *S* (of NaA') 68 at 20° — BaA' 7aq. *S* (of BaA') 066 at 20°, 56 at 100°

Phenanthrene (β)-carboxylic acid

$C_{14}H_7$ C CO_2H $C_{14}H_7$ CH [c 252°] Formed in like manner from a calcium phenanthrene sulphonate contained in the mother-liquid from which its (α)-isomeride has crystallised (Japp). Stellate groups of straight needles (from $HOAc$), insol. water, sol. alcohol, ether, and $HOAc$. Yields phenanthraquinone on oxidation — NaA' 5aq. laminae. *S* (of NaA') 62 at 20°, v e. sol. boiling water — BaA' 6aq. rectangular laminae. *S* (of BaA') 27 at 20°, 37 at 100°.

PHENANTHRENE (α) SULPHONIC ACID SO_3H $C_{14}H_7$ CH $C_{14}H_7$ CH Made, together with a (β)-acid

which forms more soluble salts, by heating phenanthrene (1 pt) with H_2SO_4 (1 pt) at 100° (Graebe, *A* 167, 152, Japp, *C* *J* 37, 93, *B* 11, 218). Crystalline mass, m sol. wa' r. Yields phthalic acid on oxidation — CaA' 4aq. small plates, v sol. hot water — PbA' 2aq. crystalline

Phenanthrene sulphonic acid $C_{14}H_7$ SO_3H Got by heating phenanthrene (8 pts) with H_2SO_4 (2 pts) at 170° (Morton & Geyer, *A* *C* *J* 2, 203, *B* 13, 1870). Pearly needles, sol. water and alcohol. Its salts are less soluble than those of the (α)-acid — KA' — BaA' 3r. — PbA' 3aq.

Phenanthrene disulphonic acid $C_{14}H_6(SO_3H)_2$ Formed by heating phenanthrene with fuming H_2SO_4 (E Fischer, *B* 13, 314, Hazura & Julius, *M* 5, 188). Syrup — K_2A' 2aq. powder, v sol. water — BaA' — Ag_2A' yellowish powder

Bromo-phenanthrene sulphonic acid $C_{14}H_6Br(SO_3H)$ Made by sulphonating bromo-phenanthrene (Anschutz & Siemensky, *B* 13, 1179) — KA' needles, sl sol. water — BaA' , in soluble pp — AgA' glistening needles

PHENANTHRIDINE $C_{14}H_9N$ *v* $C_{14}H_8$ CH $C_{14}H_8$ N

V D 65 [104°] (360°) Made by passing the vapour of benzylidene-aniline through a red hot tube (Pictet & Ankersmit, *B* 22, 3339). White needles, v e sol. alcohol, sl sol. hot water. Its aqueous solution shows blue fluorescence. $NaNO_2$ gives a bulky pp of the nitrite. Tin and HCl give a hydride crystallising in needles [100°] — $BHCl$ needles, v sol. water — $B'H_2PtCl_4$ needles — $B'HAuCl_4$ — $B'HHgCl_2$ [190°] — $B'C_6H_5N_3O$, needles — $B'HCrO_3$

Methylo-iodide $B'MeI$ [201°]

Oxy phenanthridine $C_{14}H_7$ CO $C_{14}H_7$ NH Got by reducing *o*-nitro-*o* phenyl benzoic acid with zinc dust and ammonia. It yields phenanthridine on distillation with zinc dust

PHENANTHROL $C_{14}H_9$ OH [112°] Formed by potash fusion from phenanthrene sulphonic acid (Rehs, *B* 10, 1253). Thin laminae (from benzene ligroin), with bluish fluorescence. V sol. alcohol and ether, sl sol. water, v sol. alkalis

Acetyl derivative $C_{14}H_8$ OAc [118°]

PHENANTHROLINE $C_{12}H_8N_2$ *v* CH CH C CH C N CH CH C N CH [78°] (above 360°)

Prepared by heating *m* phenylene diamine or (*B* 3) amido quinoline with nitro benzene, glycerin, and H_2SO_4 (Skraup, *B* 15, 895, *M* 3, 578, 5, 532, La Coste, *B* 16, 674). Got also by heating its carboxylic acid [209°] (Gerdesen, *B* 22, 252). Colourless anhydrous four-sided tables [78°] or needles (containing aq) [66°]. M sol. hot water, v sol. alcohol, v sl sol. ether. Bromine gives in a solution of its hydrochloride a crystalline pp $C_{14}H_8N_2Br$ [149°], converted by hot alcohol into $C_{14}H_7N_2Br$ [178°]. $KMnO_4$ oxidises it to dipyrindyl dicarboxylic acid

Salts — $B'H_2Cl_2$ 2aq. colourless prisms. Its aqueous solution yields on evaporation long prisms of $B'HCl$ 1aq — $B'HBz$ 1aq [280°] — $B'H_2CrO_4$ golden needles — $B'C_6H_5N_3O$ [288°] — $B'H_2PtCl_4$ 1aq — $B'HNO_3$ prisms, m sol. hot water

Methylo-iodide $BMeI_{2aq}$ Golden prisms, sol water, sl sol alcohol, insol ether

Oxy-phenanthroline $C_{12}H_8N_2O$ [160°]. Formed in small quantity, together with phenanthroline, by heating *m* nitro aniline with glycerin, nitro-benzene, and H_2SO_4 (La Coste). Needles (fr *m* benzene) — $B'H.PtCl_2aq$

Methyl-phenanthroline *v* p 355

Di-methyl-phenanthroline

$CH_3CH_2C(CH_3)CH_2CH_2C(CH_3)CH_2CH_2N(CH_3)CH_2CH_2N(CH_3)CH_2CH_2$ [98°] Formed

by the action of paraldehyde and HCl_{aq} at 100° on *m* phenylene-diamine or (*B* 3) amido (*Py* 3)-methyl pyridine (Von Miller & Niederlander, *B* 24, 1740, *Scrifft*, *B* 24, 2127) Colourless needles (from water). Melts at 78° when containing water of crystallisation. *V* sol alcohol, volatile with steam. Yields on reduction a base whence $BzCl$ forms $C_{11}H_{11}BzN_2$ [164°]. — $B'H.PtCl_2$

Di hexyl di amyl-phenanthroline

$C_6H_{11}CH_2CH_2CH_2CH_2CH_2CH_2N(CH_2CH_2CH_2CH_2CH_2CH_2)CH_2CH_2CH_2CH_2CH_2CH_2N(CH_2CH_2CH_2CH_2CH_2CH_2)CH_2CH_2CH_2CH_2CH_2CH_2$ [51°]

Formed by the action of cantharol on an alcoholic solution of *m* phenylene diamine or of amido-hexyl-amyl quinoline at 100° (Von Miller & Gerdeissen, *B* 24, 1731, *cf* Schiff, *A* 253, 322) Needles. Its alcohol solution does not fluoresce — $B'HCl$ — $B'H.PtCl_2aq$ [201°–210°] Orange powder — $B'O.C_6H_5N_2O$, [104°] Needles

Pseudo phenanthroline $C_{12}H_8N_2$, *±* c

$CH_3N(CH_3)CH_2CH_2CH_2CH_2CH_2N(CH_3)CH_2CH_2CH_2CH_2CH_2CH_2$ [173°] Made from

p phenylene diamine, glycerin, H_2SO_4 , and nitro benzene (Skraup & Vorkmann, *M* 4, 569), and from PhN , $C_6H_5NH_2$, glycerin, and H_2SO_4 (Lellmann & Lippert, *B* 24, 2623) Obtained also as a by-product in the preparation of (*B* 3)-nitro-quinoline from *p* nitro aniline (Bornemann, *B* 19, 2377) Crystallises from water in needles (containing 4aq), *v* sol alcohol, sl sol ether. Oxidised by $KMnO_4$ to dipyrindyl dicarboxylic acid

Salts — $B'HCl_{2aq}$ plates — $B'H.Cl_2$ monoclinic prisms — $B'H.PtCl_2$ 2½aq — $B'H.Br$ — $B'H.Br$ — $B'Br$ — $B'I$ — $B'HI$ — $B'H.Cr_2O_7$ 2½aq orange needles, sl sol cold Aq

Methylo-iodides $B'MeI_{2aq}$ lemon yellow needles — $B'MeI_{2aq}$ red tables

PHENANTHROLINE (*B*) CARBOXYLIC

ACID $C_{12}H_7(CO_2H)N_2$ [277°] Formed by oxidation of (*B*) methyl phenanthroline with chromic acid (Skraup & Fischer, *M* 5, 527) Minute needles, sl sol water and alcohol, sol alkalis and acids — Ca_2HA 10aq needles Yields phenanthroline on distillation with alkalis

Phenanthroline- (*Py* 3)-carboxylic acid

$C_{12}H_7(CO_2H)N_2[N(CO_2H)=12]$ [209°] Formed by oxidation of (*a*)-methyl phenanthroline with $KMnO_4$ and H_2SO_4 (Gerdeissen, *B* 22, 250) Pale-yellow needles (containing aq), *v* sl. sol. cold water

Phenanthrone *v* PHENANTHRAQUINONE

PHENANTHROXYLENE - ACETOACETIC

ACID *Ethyl ether* $C_{20}H_{16}O_4$, *±* c . $C_6H_5CO_2CO_2C_2H_5$ Mol *w* (by Raoul's method) C_6H_5CO

287 (calc. 320) [185°] Formed by heating phenanthraquinone with acetoacetic ether and NEt_{3aq} or $KOHAq$ (Japp & Streetfield, *O* J 48, 27, Japp & Klingemann, *O* J 59, 3) White silky needles, *v* sol. hot benzene and alcohol.

Reactions — 1. **Alcoholic potash** forms $C_{11}H_{12}O_2$ crystallising from alcohol, after solution at 100° in sealed tubes, in needles [259°] — 2 **Alcoholic ammonia** at 100° yields lustrous yellow laminae [188°] of $C_{10}H_{12}O_2$ or N_2O_3 — 3 **Alcoholic HCl** yields $C_{12}H_{12}ClO_2$ [146°], whence alcoholic NH_3 produces $C_{12}H_{12}NO_2$ — 4 Heating with $HOAc$ forms three compounds, $C_{12}H_{12}O_2$, decomposing at 285° without melting, $C_{12}H_{12}AcO$, [165°–171°], and $C_{14}H_{16}O_2$, [227°] — 5 **Propionic acid** at 140° yields silky needles $C_{12}H_{12}O_2$, and crystalline $C_{12}H_{12}O_2$, which body is also got by heating with propionic anhydride at 150° — 6 **HI** and **P** yield $C_{12}H_{12}O_2$ [128°], whence potash forms an acid $C_{11}H_{11}O_4$ [255°], which gives BaA''_{2aq} and AgA'' — 7 Heated on the water bath with alcohol containing a few drops of H_2SO_4 , it forms $C_{11}H_{11}O(OEt)(CO_2Et)$ [144°], which yields, with phenyl-hydrazine, $C_{11}H_{11}(N_2HPh)(OEt)(CO_2Et)$ [220°] and, on hydrolysis, $C_{11}H_{11}(OEt)(CO_2H)$, [203°], crystallising in colourless needles — 8 Alcohol containing a large quantity of H_2SO_4 forms $C_{12}H_{12}O_2$, [144°], sol boiling alcohol, and $C_{11}H_{12}O_2$ [277°], insol alcohol

Isophenanthroxylene acetoacetic acid $C_{15}H_{12}O_4$, [269°] Formed by saponifying its ether with $NaOH$ (Japp & Klingemann, *O* J 59, 14) Flat needles (from alcohol) Conc $KOHAq$ at 150° yields $C_{15}H_{12}O_4$ Ac_2O at 150° forms $C_{15}H_{12}O_4$, insol alcohol, and $C_{15}H_{11}O_4$, [226°] — CaA' 9aq

Ethyl ether EtA' [177°] Formed from phenanthroxylene acetoacetic ether by heating with formic acid (*S* G 1 22) at 130° for an hour, or by boiling with H_2SO_4 diluted with twice its weight of water (Japp & Klingemann, *O* J 59, 3) Colourless triclinic prisms (from $EtOAc$), a b c = 964 1 507, α = 78° 6', β = 72° 0', γ = 83° 18', sol alcohol, benzene, and light petroleum

Reactions — 1 Ac_2O at 150° forms the acetyl derivative $C_{15}H_{12}AcO_4$, [165°–170°], crystallising from benzene in needles — 2 **Phenyl hydrazine** in alcoholic solution at 135° yields the compound $C_{20}H_{16}(N_2HPh)O$, crystallising in yellow needles [212°] — 3 **Zinc** and HCl yield a crystalline compound [165°–170°], and also $C_{15}H_{12}O_4$, [123°], a body which yields a phenyl hydrazine $C_{20}H_{16}(N_2HPh)O_2$ — 4 **Bromine** in $CHCl_3$ yields $C_{15}H_{12}BrO_4$, [212°] crystallising in yellow prisms and tables — 5 $HIAq$ at 100° forms $C_{11}H_{12}O$ [215°] crystallising in slender flat needles

PHENAZINE $C_{12}H_8N_2$, *±* c C_6H_5 $\langle N \rangle$ C_6H_5 , **Asophenylene** [171°] (above 360°) *S* (alcohol) 2 in the cold

Formation — 1 By distilling *m*- or *p* azobenzoic acid with excess of lime (Claus, *B* 5, 867, 610, 6, 723, 8, 39, 600, 10, 1303, *A* 168, 1) — 2 By passing aniline over red-hot PbO (Schichutzky, *J* R 6, 248), or merely through a red-hot tube (Bernthsen, *B* 19, 3256) — 3 By heating equal weights of pyrocatechin and *o*-phenylene diamine at 205°, followed by atmospheric oxidation of the resulting dihydride (Ris, *B* 19, 2206) — 4 By heating *o*-amido-phenol and oxidising in the same way (Ris) — 5 From di-amido phenazine by the diazo-reaction (*O* Fischer & Hepp, *B* 22, 358; Nietzki, *B* 23, 1855)

Properties — Long yellowish needles (by sub-

limation), ν sol hot alcohol and ether, ν sl sol. hot water H_2SO_4 forms a blood red solution, becoming yellow on dilution. Volatile with steam. Combines with bromine, forming $C_{12}H_8N_2Br_2$, which separates from benzene in yellow needles, and with chlorine, forming $C_{12}H_8N_2Cl_2$ as unstable red crystals (from alcohol).

Salts— B^+HCl . Tables (from $HClAq$), decomposed by hot water— B^+HAuCl_4 crystals— $B^+H_2HgCl_4$ — $B^+H_2PtCl_6$, 2aq yellow plates— B^+HI Dark-green needles— B^+HBr Brown crystals— $B^+C_6H_5N_3O$, $[180^\circ-190^\circ]$ Long yellow needles, sol cold alcohol— $B^+Hg(NO_3)_2$ ruby-red crystals (from HNO_3)— B^+AgNO_3 .

Dihydrate $C_{12}H_{10}N_2 \cdot 2 H_2O$ $C_6H_5 \begin{smallmatrix} \text{NH} \\ \diagup \quad \diagdown \\ \text{N} \end{smallmatrix} C_6H_5$

Formed by reduction of phenazine with alcoholic NH_3 and H_2S (Olaus, A 168, 8). Trimetric laminae, almost insol water and benzene, ν sl sol cold alcohol. Conc H_2SO_4 forms a green solution, turning red on further addition of H_2SO_4 , and green on cautious addition of water, in this reaction an unstable base $C_{12}H_{10}N_2$ appears to be formed, giving the salts $B^+H_2Cl_2$ and $B^+H_2PtCl_6$, the latter forming in green needles.

Di-chloro phenazine $C_{12}H_8Cl_2N_2$ $[144^\circ]$ Made from phenazine and PCl_5 .

Nitro-phenazine $C_{12}H_8(NO_2)_2N_2$ $[210^\circ]$ Made by nitration. Yellowish green needles.

Amido-phenazine $C_6H_5 \begin{smallmatrix} \text{N} \\ \diagup \quad \diagdown \\ \text{N} \end{smallmatrix} C_6H_5(NH_2) \begin{smallmatrix} 1 \\ 2 \end{smallmatrix} \text{ s}$

$[265^\circ]$ Got by sublimation from a mixture of di amido phenazine and zinc dust (O Fischer & Hepp, B 22, 357). Long red needles with bronze lustre (from alcohol). Strong base. Its dilute solution shows orange red fluorescence— $B^+H_2PtCl_6$, 2 $\frac{1}{2}$ aq.

u-Di-amido phenazine $C_{12}H_{10}N_4$, ν s

$C_6H_5 \begin{smallmatrix} \text{N} \text{ CH } \text{N} \\ \diagup \quad \diagdown \\ \text{N} \text{ CH } \text{N} \end{smallmatrix} C_6H_5$ A product of oxidation of o phenylene-diamine by $FeCl_3$ (Fischer & Hepp, B 22, 355, 23, 841, 2788, cf Griess, B 5, 202, Rudolph, B 12, 2211, Wiesinger, A 224, 353). Formed also from o phenylene diamine and Cyl (cf Hubner, B 9, 777, 10, 1715), and by boiling o phenylene diamine hydrochloride (25 g) with amido-azo benzene (3 g) and acetic acid (30 g of 75 p c) for two hours. Long brownish-yellow needles or yellow plates (by sublimation). Conc H_2SO_4 forms a grass-green solution, turned red on dilution. Its solution in benzene or alcohol fluoresces greenish-yellow. Alcoholic solutions of the salts fluoresce dark orange red. Yields with benzil the quinoxaline $C_{16}H_{10}N_4$ — B^+HCl 3aq— $B^+H_2SO_4$ 3aq.

Di-acetyl derivative $C_{12}H_8Ac_2N_4$ $[c 270^\circ]$ Yellow needles.

Di-formyl derivative Reddish-yellow mass.

Di-amido-phenazine

$\begin{bmatrix} 1 \\ 2 \end{bmatrix} C_6H_5(NH_2) \begin{smallmatrix} \text{N} \\ \diagup \quad \diagdown \\ \text{N} \end{smallmatrix} C_6H_5(NH_2) \begin{bmatrix} 1 \\ 2 \end{bmatrix}$ $[280^\circ]$

Got by heating the tin double salt of tri amido-diphenylamine with water, $CaCO_3$, and MnO_2 (Nietzki, B 23, 1854). Long dark yellow needles, ν e sol alcohol, ether, and hot water. Yields phenazine on elimination of amidogen— B^+HNO_3 , greenish needles— $B^+C_6H_5N_3O$.

Di-acetyl derivative $C_{12}H_8N_4O_4$ $[c 280^\circ]$.

Tri amido-phenazine $C_{12}H_8N_6$, ν s.

$\begin{bmatrix} 4 \\ 2 \end{bmatrix} C_6H_5(NH_2) \begin{smallmatrix} \text{N} \\ \diagup \quad \diagdown \\ \text{N} \end{smallmatrix} C_6H_5(NH_2) \begin{bmatrix} 1 \\ 2 \end{bmatrix}$ $[16, 24 \text{ or } 6]$.

Made by passing a current of oxygen through a solution of triamidobenzene hydrochloride (10 g) and $NaOAc$ (18 g) (E Muller, Z 22, 856, cf Witt, B 10, 658). Brown needles ν sol hot alcohol and water, the solutions showing yellow fluorescence. Conc H_2SO_4 gives a yellow solution, changing violet, red, and finally yellow on dilution. It decomposes when heated to 100° — $B^+(HNO_3)_2$, 2aq needles with green lustre. Forms a crimson solution with yellow fluorescence. The solution becomes yellow on dilution.

Tri-acetyl derivative $C_{12}H_8Ac_3N_6$.

Tetra-amido-phenazine

$NH_2 \begin{smallmatrix} \text{CH} \text{ CH } \text{N} \text{ CH } \text{CH } \text{N} \end{smallmatrix} NH_2$ Formed by passing $NH_2 \begin{smallmatrix} \text{CH} \text{ CH } \text{N} \text{ CH } \text{CH } \text{N} \end{smallmatrix} NH_2$ air through a hot solution of tetra amido benzene hydrochloride (10 g) and $NaOAc$ (20 g) (Nietzki & E Muller, B 22, 447). Brown needles (from hot water or alcohol), or yellow needles containing aniline of crystallisation (from aniline). Its solutions exhibit yellowish green fluorescence. Conc H_2SO_4 forms a yellow solution, passing, on dilution, through blue, violet, and red, to yellow— $B^+(HNO_3)_2$, 2aq lustrous green needles giving a crimson solution.

References—METHYL-PHENAZINE and DI-OXY PHENAZINE.

Tolylene red $C_6H_2(NMe_2) \begin{smallmatrix} \text{N} \\ \diagup \quad \diagdown \\ \text{N} \end{smallmatrix} C_6H_2Me(NH_2)$ is also a phenazine derivative (Bernthsen, A 236, 332).

PHENTRIAZINE $C_6H_5 \begin{smallmatrix} \text{N} \text{ CH} \\ \diagup \quad \diagdown \\ \text{N} \text{ N} \end{smallmatrix}$ $[66^\circ]$

$(235^\circ-240^\circ)$ Formed from the formyl derivative of o nitro phenyl hydrazine, alcohol, $HOAc$, and sodium amalgam (Bischler, B 22, 2806), and by the action of P_2O_5 on o amido phenyl methyl hydrazine (Hempel, J pr [2] 41, 174). Yellow needles, ν sol warm water and cold alcohol, very volatile with steam.

TRIPHENAZINE DIHYDRIDE

$C_{12}H_{12}N_4$, ν s $C_6H_5 \begin{smallmatrix} \text{N} \\ \diagup \quad \diagdown \\ \text{N} \end{smallmatrix} C_6H_5 \begin{smallmatrix} \text{NH} \\ \diagup \quad \diagdown \\ \text{NH} \end{smallmatrix} C_6H_5$ or

$C_6H_5 \begin{smallmatrix} \text{NH} \\ \diagup \quad \diagdown \\ \text{N} \end{smallmatrix} C_6H_5 \begin{smallmatrix} \text{N} \\ \diagup \quad \diagdown \\ \text{NH} \end{smallmatrix} C_6H_5$ *Homofluorine*

dine Formed by heating u di amido phenazine hydrochloride (4 pts) with o phenylene diamine at 205° for 15 minutes (Fischer & Hepp, B 23, 2791). Bluish violet powder or lustrous green needles, ν sl sol alcohol, forming a violet red solution with yellowish red fluorescence. Solutions of its salts are blue, with brownish red fluorescence.

PHENAZOXINE $C_6H_5 \begin{smallmatrix} \text{NH} \\ \diagup \quad \diagdown \\ \text{O} \end{smallmatrix} C_6H_5$ *Oxy*

or *oxido-diphenyl amine* $[148^\circ]$ Obtained by heating together equal quantities of o-amidophenol and pyrocatechin at $260^\circ-280^\circ$. Silvery scales. ν sol alcohol, ether, chloroform, and benzene. Sublimable. Distills, in great part undecomposed. By nitration and reduction the leuco-base of a violet dye stuff is obtained (Bernthsen, B 20, 942).

PHENBUTYL COMPOUNDS ν . **BUTYL-PHENYL COMPOUNDS**

PHENISOBUTYL CYANIDE ν . **ISO-BUTYL BENZONITRILE**.

PHENISOBUTYL - PHENETHYL - THIOUREA *v.* **ETHYL PHENYL ISOBUTYL PHENYL THIOUREA**

PHENCAPRYLAMINE *v.* **AMIDO-PHENYLOCTANINE**

PHENENYL TRIBENZOIC ACID $C_{27}H_{18}O_6$, [α 61°] G-t by potash fusion from tri-benzoyl ene benzene (Gabriel & Michael, *B* 11, 1008) Prisms, *v* sol alcohol, ether, and HOAc Yields C_6H_5Ph , when distilled with lime— Na_2A''' — Ag_2A'''

PHENENYL TRI-METHYL TRIKETONE $C_6H_4(COCH_3)_3$, [163°] Formed by the spontaneous condensation of acetoacetic aldehyde (Claisen & Stylo *B* 21, 1144) Small needles, *v* sol HOAc, sl sol alcohol, ether, and water Oxidised by nitric acid to trimesic acid

PHENETHYLAMINE *v.* **AMIDO-PHENYLETHANINE**

PHENETHYL - PHENISOBUTYL - THIOUREA *v.* *p* **ETHYL PHENYL-*p* ISOBUTYL PHENYL THIOUREA**

PHENETIDINE *v.* *Ethyl derivative of* **AMIDO-PHENOL**

PHENETOL *v.* *Ethyl ether of* **PHENOL**

DIPHENIC ACID *v.* **DIPHENYL DICARBOXYLIC ACID**

PHENNAPHTHAZINE $C_{16}H_{10}N_2$, *αβ*

$C_{16}H_{10}N_2$ $\left\langle \begin{smallmatrix} N \\ N \end{smallmatrix} \right\rangle C_6H_5$, (*αβ*) *Naphthophenazine* [142°]

Formation—1 By mixing equal mols of *o*-phenylene diamine and (*β*) naphthoquinone in 50 p c acetic acid—2 By oxidation of equal mols of *o* phenylene diamine and (*β*) naphthol with alkaline potassium ferriyanide—3 By the decomposition of sulpho benzene azo (*β*)-naphthyl phenyl amine by treatment with boiling dilute mineral acids $C_6H_4(SO_3H)N_2C_6H_5$, NHC_6H_5 , $= C_{10}H_8N_2$, C_6H_5 , + $C_6H_5(NH_2)SO_3H$

Preparation—Fifty grms of the dyestuff are dissolved in 500 c c of boiling water, and 125 c c of conc H_2SO_4 is slowly added to the hot solution, the colour acid, which is first precipitated, redissolves, and on cooling the sulphate of the azine crystallises out in red needles, whilst sulphathionic acid remains in solution

Properties—Glistening yellow needles or prisms Sublimes at about 200° in long flat needles Distils undecomposed above 360° Sl sol alcohol, ether, and cold benzene, *v* sol hot benzene Dissolves in conc H_2SO_4 with a brownish red colour, becoming yellow on dilution

Salts—With each acid it forms two different salts— $B'HCl$ * long reddish yellow needles and warty crystals— $B'_2H_2SO_4$ * red needles and thick garnet-red prisms— $B'HNO_3$ * yellow and red needles, both sparingly soluble (Witt, *B* 20, 572)

PHENOCYANIN C_6H_5NO or $C_6H_5NO_2$ Dark blue mass with coppery lustre, got by atmospheric oxidation of a mixture of phenol and NH_3 (Phipson, *B* 6, 828) Insol water, sol alcohol Coloured red by acids

PHENOL C_6H_5O *αα* C_6H_5OH *Carbolic acid Phenyl hydrate* Mol *w* 94 [41°] (1815°) (Dale & Schorlemmer), (1829°) (Pinette, *A* 248, 82) S 66 at 17° SG ρ 1.0702 (Bruhl), ρ 1.0906 (P) SV 1018 CE (ρ 10°) 0.0083 (P). μ_p 1.6686 (B) R_{20} 45.71 HC *v* 736,500 (Berthelot, *A Ch* [6] 10, 458, 18, 829) HC *p* 787,100

H F 47,841 (Stohmann, *J pr* [2] 83, 471; 28,000 (Von Rechenberg) Occurs in castoreum (Wöhler, *A* 67, 360), and in small quantities in urine of cows, horses, and men (Stadeler, *A* 77, 18, Lieben, *A Suppl* 7, 240, Hoppe Seyler, *C J* 25, 628, Munk, *B* 9, 1596, Salkowski, *B* 9, 1595, Baumann, *B* 9, 54, 1389, 1715) Contained in considerable quantity in coal tar (Runge, *P* 31, 69, 32, 308, Laurent, *A Ch* [8] 3, 195), and in the products of the dry distillation of gum benzoin, quinic acid, wood, and bones

Formation—1 By distilling *o*-, *m*- or *p*-, oxy-benzoic acid alone or with lime (Gerhart, *Rev scient* 10, 210, Rosenthal, *Z* [2] 5, 627)—2 A product of the distillation of glycerin with $CaCl_2$ (Linnemann & Zotte, *A* 174, 87, *Suppl* 8, 254)—3 By heating anisole with conc $HIAq$ or $HClAq$ at 140° (Graebe, *A* 139, 149)—4 From aniline by the diazo reaction (Griess, *A* 137, 39)—5 From benzene sulphonic acid by potash-fusion (Wurtz, *Bl* [2] 8, 197, cf Degener, *J pr* [2] 17, 394)—6 From phenol *p* sulphonic acid by distillation with dilute H_2SO_4 and superheated steam at temperatures above 116° (Armstrong & Miller, *C J* 45, 148)—7 By heating acetylene with fuming H_2SO_4 and fusing the product with potash (Berthelot, *C R* 68, 539)—8 By shaking benzene with palladium that has absorbed hydrogen and air (Hoppe Seyler, *B* 12, 1552)—9 By the direct action of hydrogen peroxide on benzene (Leeds, *B* 14, 975)—10 In small quantity by the action of dry oxygen on boiling benzene containing $AlCl_3$ (Friedel & Crafts, *A Ch* [6] 14, 435, *C R* 86, 884)—11 A product of fermentation of proteids (Baumann, *B* 10, 685, Weyl, *H* 1, 339, Brieger, *J pr* [2] 17, 134)

Preparation—The aqueous solution got by stirring coal tar oil with $NaOHAq$ is diluted with water as long as naphthalene separates The liquid is then exposed to the air, with frequent stirring, for several days, and then fractionally ppd by acid (*e g* CO_2), the last fraction being nearly pure phenol The phenol is dried by heating to boiling in a current of air, and further purified by crystallisation (Hugo Muller, *Z* [2] 1, 270, cf Williamson & Scruggam, *C J* 7, 232) Phenol may also be dried by distilling over dried $CuSO_4$ (Bickerdike, *C N* 16, 188, cf Gladstone, *C N* 2, 98)

Properties—Long deliquescent needles, with strong smell, in sol water, miscible with alcohol and ether Attacks the skin Does not reddden litmus Nearly insol Na_2CO_3Aq , *m* sol NH_4Aq , *v* sol $KOHAq$ and $NaOHAq$ Antiseptic Not affected by distillation with P_2O_5 , lime, or BaO A solution of phenol even in 43,000 pts of water gives a pp of tri-bromophenol on addition of bromine-water (Landolt, *B* 4, 770) $FeCl_3$ gives a violet colour to an aqueous solution, the reaction is prevented by acetic acid and by alcohol (Hesse, *A* 182, 161) Conc H_2SO_4 poured beneath a solution containing phenol and nitric acid (1 p c) gives an intense red ring, with nitrous acid, red and green rings are got, with chlorates, a pale-yellow ring with blue below (Lundo, *C N* 58, 1, 15) An aqueous solution of phenol (4 vols) mixed with ammonia (1 vol) gives on warming with a few drops of bleaching powder solution a blue colour

which becomes red on addition of acids (Salkowski, *Fr* 11, 316) Boiling aqueous mercurous nitrate gives a deep-red colour (Plugge, *Fr* 11, 178) Millon's reagent gives on boiling a yellow pp which dissolves in nitric acid forming a deep red liquid, salicyloic acid behaves in like manner (Almén, *J* 1878, 1079) On adding *p*-oxybenzoic aldehyde and an equal volume of H_2SO_4 , a yellow liquid is got, turned crimson by potash (aurin) Phenol is poisonous (Wöhler, *A* 65, 844, Duplay & Carin, *C R* 112, 627)

Reactions — 1 Decomposed by passing through a red-hot tube yielding benzene, toluene, xylene, naphthalene, anthracene, and phenanthrene (Kramers, *A* 189, 129) When crude phenol is used the product deposits a crystalline hydrocarbon $C_{10}H_{12}$ [82.9°], S G 1.012 (63° at 9 mm) This hydrocarbon is volatile in the cold, smells like camphor, and is sol alcohol, ether, and petroleum-spirit It is resinified by atmospheric oxygen It absorbs bromine, yielding a liquid bromide By heating for 4 hours at 100° *in vacuo* the hydrocarbon is polymerised, yielding a solid [200°–220°] (Roscoe, *C J* 47, 669) — 2 Chlorine forms *o*- and *p*-chloro phenol (4,2,1)-di-chloro-phenol, (6,4,2,1) tri-chloro phenol, and tetra-, pent-, and hexa-chloro-phenols Exhaustive chlorination gives C_6Cl_6 , CCl_4 , C_2Cl_4 , and CO_2 (Ruoff, *B* 9, 1488) An intermediate body is C_6Cl_5O , [825°] (Hugouenq, *C R* 109, 309) $KClO_3$ and HCl yield tri-chloro-phenol and tri- and tetra- chloro-quinone — 3 Chlorine acting on phenol in alkaline solution forms the acid $C_6(OH)CCl_2 > C(OH)CO_2H$ (Hantzsch, *B* 20, 2780, 22, 1238) — 4 Bromine-water forms tri-bromo-phenol Heat (68492 units) is given out in the reaction Br in excess gives $C_6H_3Br_3O$ (Werner, *C R* 100, 799, *Bl* [2] 46, 280) — 5 Iodine and alkali at 60° form $C_6H_2I_2O$ [157°], which is violet red, insol water, and forms a red solution in alcohol and ether It is converted into tri iodo-phenol by boiling with KOH aq (Messinger & Vortmann, *B* 22, 2818) — 6 Chloride of iodine forms mono- and di-, iodo-phenol (Schützenberger, *C R* 54, 197) — 7 Oxidised by nitro *sensu* and dilute NaOH in the cold to oxalic acid and CO_2 (Stegfried, *J pr* [2] 31, 642) — 8 If a rapidly alternating electric current be passed through a solution containing phenol, magnesic sulphate, and magnesic bicarbonate, the following bodies are formed, owing to the rapidly alternating oxidation and reduction: pyrocatechin, hydroquinone, di oxy diphenyl, formic acid, succinic acid, and oxalic acid (E Drechsel, *J pr* [2] 29, 249) Another product is $C_6H_{10}O$, an oil (158°–178°) with aromatic smell which forms a phenyl hydrazide $C_{12}H_{14}N$ [108°] By continuing the alternating current this oil is converted in *n* hexoic acid — 9 Electrolysis with carbon electrodes in aqueous solution rendered slightly alkaline by KOH yields a di-oxy-benzoic acid $C_6H_4O_2$ [98°] and an amorphous acid $C_6H_6O_2$, insol water and ether, sol alcohol The amorphous acid yields picric acid with HNO_3 , and on protracted boiling with dilute HCl aq it yields amorphous infusible $C_{12}H_{10}O_2$, and amorphous $C_{12}H_{10}O$, [60°], sol water, alcohol, and ether In like manner $NaOPh$ yields on electrolysis $C_{12}H_{10}O_2$, separable by hot HCl aq into infusible $C_{12}H_{10}O_2$, insol water and ether, and

$C_{12}H_{10}O_2$, [78°], sol water (Bartoli & Papasogli, *G* 14, 90) — 10 Taken internally it is partly oxidised to hydroquinone and pyrocatechin (Nencki & Giacoso, *H* 4, 825) — 11 H_2O_2 oxidises it to pyrocatechin, hydroquinone, and quinone (Martinon, *Bl* [2] 43, 156) When treated in aqueous solution with NH_3 , H_2O_2 , Na_2CO_3 , and hydroxylamine hydrochloride it yields phenolquinonimide, which colours the liquid bright blue (Wurster, *B* 20, 2934) — 12 CrO_2Cl_2 , followed by water, gives $O(C_6H_4OH)_2$ (Etard) CrO_2Cl_2 and $HOAc$ gave tri- and tetra- chloro-quinone (Carstenjen, *J pr* [2] 2, 82) — 13 Fusion with NaOH yields resorcin, pyrocatechin, and phloroglucin Potash fusion gives *o* and *m*-oxybenzoic acids and two di oxy diphenyls (Born & Schröder, *B* 11, 1332, 12, 417) — 14 PCl_5 forms $PCl_2(OPh)$, $PCl(OPh)_2$, and $P(OPh)_3$, which may be separated by fractional distillation *in vacuo* (Noack, *A* 218, 85; Anschütz & Emery, *A* 239, 810, *A* 253, 110) The compound $PCl_2(OPh)$, (90° at 11 mm), (216° at 760), S G 1.854, is converted by chlorine into $PCl_3(OPh)$, whence SO_2 produces $POCl_2(OPh)$ (122° at 11 mm) Bromine converts $PCl_3(OPh)$ into $PCl_2Br(OPh)$, which is crystalline but very unstable Sulphur at 190° converts $PCl_3(OPh)$ into $PSCl_2(OPh)$, a colourless liquid (120° at 11 mm), S G 1.4059 The compound $PCl(OPh)_2$ (172° at 11 mm) (295° at 760 mm) is converted by chlorine into $PCl_3(OPh)_2$, which is crystalline, and insol ether Bromine converts $PCl(OPh)_2$ in ether into orange yellow crystals of $PClBr_2(OPh)_2$ Sulphur and $PCl(OPh)_2$, at 190° yield $PSCl(OPh)_2$, crystallising in colourless needles [64°], (194° at 11 mm) Tri-phenyl phosphite $P(OPh)_3$ combines with chlorine, forming $PCl_2(OPh)_3$, whence water produces tri phenyl-phosphate $PO(OPh)_3$, [45°] (245° at 11 mm) $P(OPh)_3$ with sulphur at 190° yields $PS(OPh)_3$, [50°] (245° at 11 mm) S G 1.2341 — 15 PCl_5 forms hardly any chloro-benzene (Otto, *A* 145, 817, cf Glutz, *A* 143, 181) — 16 P_2S_5 forms, on heating, phenyl mercaptan, di phenylenedisulphide $C_{12}H_{10}S_2$, and some Ph_3S (Kekulé, *C R* 64, 752, Graebe, *B* 7, 51, 397, Geuther, *A* 221, 57) P_2S_5 forms benzene, Ph_3PO , and H_2S — 17 Aqueous $KMnO_4$ oxidises it to oxalic acid and CO_2 (Tollens, *Z* [2] 4, 715) — 18 Distillation over heated zinc dust yields benzene — 19 H_2SO_4 (1 pt) forms *o* and *p* sulphonic acids (Kekulé, *Z* [2] 3, 197) H_2SO_4 (1½ pts) at 160° forms di oxy di phenyl sulphone (Glutz, *A* 147, 52) $K_2S_2O_8$, heated with a solution of KOC_2H_3 at 65°–70°, forms $C_6H_5O SO_3OK$, which crystallises in tables, S 14 at 15°, sol hot alcohol This salt occurs in urine It is decomposed by boiling with water and dilute acids into phenol and H_2SO_4 At 150° the dry salt changes to the isomeric potassium phenol *p*-sulphonate Phenyl sulphuric acid is also a product of the passage of a rapidly-alternating electric current through a solution of phenol, $MgSO_4$, and magnesic bicarbonate The free acid is very unstable (Baumann, *B* 11, 1907; Brieger, *H* 8, 311, Drechsel, *J pr* [2] 29, 240) — 20 SO_2Cl_2 at 150° forms chloro-phenol (Dubois, *Z* [2] 2, 706) SO_2Cl_2 forms *o*- and *p*-chlorophenols and their sulphonic acids (Armstrong & Pike, *C N* 29, 288) — 21 Nitric acid to *ms* *o*- and *p*-nitro-, di-nitro, and tri-nitro-phenols.—

92 Potassium and sodium dissolve, giving off hydrogen and forming phenylates. These absorb CO_2 , forming $\text{PhO CO}_2\text{K}$ and $\text{PhO CO}_2\text{Na}$, which yield oxy-benzoates when strongly heated (v o oxy benzoate actm) (Kolbe, *J pr* [2] 10, 89) — 23 CrO_3 forms phenoquinone $\text{C}_6\text{H}_4\text{O}_2$ (Wichelhaus, *B* 5, 248, 846) — 24 Nitrous acid forms nitroso phenol, the mono-oxim of quinone (Baeyer, *B* 7, 967) H_2SO_4 , to which 6 p c of KNO_3 has been added, gives when shaken with a mixture of phenol (1 vol) and H_2SO_4 (1 vol) a brown colour, changing to green, and finally to a magnificent blue. On pouring into water brown flakes are deposited (Liebermann, *B* 7, 248 1098). In the action of H_2SO_4 and nitrous acid on phenol there is formed (a) phenol dichroin $\text{C}_6\text{H}_3\text{NO}_2$ or $\text{C}_6\text{H}_4(\text{OH})\text{N}(\text{OPh})_2$, a brown powder, sol ether, and also phenoxyls chrom $\text{C}_6\text{H}_3\text{NO}_2$, which is black and insol ether (Kramer, *B* 17, 1877, Brunner & Chuit, *B* 21, 250) (a) Phenol dichroin is also formed from quinone mono oxim and H_2SO_4 (Baeyer & Caro, *B* 7, 966) Each body yields an amorphous acetyl derivative. Phenol-dichroin forms a blue solution in alkalis and H_2SO_4 . Phenol oxychromin forms a green solution in H_2SO_4 , and a brown solution in alkalis — 25 NOCl forms chlorinated quinones (Tilden, *C J* 27, 851) — 26 Distillation with PbO yields diphenylene oxide $\text{C}_{12}\text{H}_8\text{O}$ and $\text{C}_{12}\text{H}_6\text{O}_2$, crystallising in needles [174°] (Graebe, *B* 7, 396, Behr & Van Dorp, *B* 7, 398) — 27 COCl_2 at 150° forms $\text{CO}(\text{OPh})_2$ and $\text{COCl}(\text{OPh})$ (Kempt, *J pr* [2] 1, 402) — 28 *Ammoniacalising chloride* at 290° forms aniline, diphenylamine, and Ph_2O (Merz & Weith, *B* 13, 1299) 29 *Hydrazine* solution in excess forms a white unstable substance [57°], possibly $\text{C}_6\text{H}_3\text{O}(\text{N}_2\text{H}_4)_2$ (Curtius & Thun, *J pr* [2] 44, 190) — 30 TiCl_4 acting on a benzene solution of phenol forms dark red crystals of $\text{Ti}(\text{OPh})_3\text{HCl}$, decomposed by water into phenol, titanic acid, and HCl (Schumann, *B* 21, 1079) — 31 AlCl_3 gives the solid $\text{AlCl}_2(\text{OPh})_2$, v sol hot CS_2 , insol ligroin, decomposed at once by water into phenol, alumina, and HCl (Claus & Merklin, *B* 18, 2933) On heating phenol (2 pts) with AlCl_3 (1 pt), benzene, Ph_2O , and diphenylene methane oxide are formed (Merz & Weith, *B* 14, 191) Phenol (5g) added to AlBr_3 (10g) forms amorphous $\text{Al}_2\text{Br}_3(\text{OPh})_2$, which is quickly decomposed by water (Gustavson, *J R* 16, 242) — 32 AlCl_3 and CCl_4NO_2 followed by water give aurin — 33 Heating with oxalic acid and H_2SO_4 gives rosolic acid 34 *Cyanic acid vapour* is absorbed by dry phenol forming phenyl allophanate, which crystallises from hot alcohol in unctuous crystals (Tuttle, *J* 1857, 451) — 35 *Benzyl chloride* and zinc form $\text{PhCH}_2\text{C}_6\text{H}_4\text{OH}$ on heating (Paterno, *G* 2, 2) — 36 Phenol (10g) boiled with Ac_2O (30g) and ZnCl_2 (20g) forms phenacetin, a red dye $\text{C}_6\text{H}_4\text{H}_2\text{O}_2$, which is insol benzene, sol alcohol, ether, and HOAc . Its solution in alkalis is raspberry-red (Rasnfski, *J pr* [2] 26, 54) — 37 $\text{C}_6\text{H}_5\text{CCl}_3$ forms benzaurin — 38 *Phthalic anhydride* and H_2SO_4 form phenol phthalein $\text{C}_{20}\text{H}_{14}\text{O}_4$ on heating (Baeyer, *B* 4, 658) — 39 — *Acetamide* and *benzamide* on heating form respectively PhOAc and PhOBz , while NH_3 is given off (Guarechi, *A* 171, 140) — 40 Heated in a alcoholic solution with CCl_4 and KOH or NaOH it yields o- and p- oxy benzoic acids —

41 *Paraldehyde* and stannic chloride form $\text{CH}_3\text{CH}(\text{C}_6\text{H}_4\text{OH})_2$ — 42 *Benzoyl aldehyde*, alcohol, and a few drops of HCl form white resinous $\text{C}_{20}\text{H}_{14}\text{O}_2$, whence amorphous $\text{C}_{20}\text{H}_{14}\text{Ac}_2\text{O}_2$ may be got (Michael & Ryder, *Am* 9, 130) — 43 *Benzene sulphochloride* added to a slightly alkaline solution of phenol forms $\text{C}_6\text{H}_5\text{SO}_2\text{OPh}$ as very stable colourless crystals [36°], sol alcohol, sl sol ether, saponified by alcoholic potash (Georgesen, *B* 24, 417) — 44 *Acetoacetic ether* and H_2SO_4 form (8) methyl coumarin — 45 *Chloro-acetal* and alcoholic NaOPh at 160° form $\text{C}_6\text{H}_5\text{OCH}_2\text{CH}(\text{OEt})_2$ (255°) (Autenrieth, *B* 24, 162) — 46 CCl_4 , COCl_2 and AlCl_3 give diphenyl carbonate (Heutschel, *J pr* [2], 36, 815) 47 o *Oxy-benzoyl aldehyde*, HOAc , and H_2SO_4 form, on warming, oxyaurin $\text{C}_{10}\text{H}_6\text{O}_4$, which greatly resembles aurin (Liebermann, *B* 9, 801, 11, 1436) Zulkowsky (*M* 5, 111) obtained a dyestuff $\text{C}_{20}\text{H}_{14}\text{O}_4$ soluble in aqueous NaHSO_4 , and another dyestuff insol NaHSO_4 — 48 CH_2Cl_2 and dilute caustic soda form $\text{CH}_2\text{C}(\text{OPh})_2$ [98 5°] crystallising in plates, v sl sol water (Heiber, *B* 24, 3678)

Estimation — 1 Titrated by adding bromine water till the solution is yellow, or by adding excess of bromine, KI , and starch, then titrating with $\text{Na}_2\text{S}_2\text{O}_3$. The bromine solution contains 40 grms Br and 20 grms KBr per litre (Degener, *J pr* [2] 17, 880, cf Landolt, *B* 4, 770, Koppeschaar, *Fr* 15, 233, Weinreb & Bondi, *M* 6, 506, Giacosa, *Fr* 6, 45) — 2 14 to 15 grms of pure KOH are dissolved in 1 litre of water and 10 grms of bromine gradually added, the solution is then diluted till 50 c c corresponds to 0.05 gm of pure phenol. To ascertain the strength of any phenol solution 50 grms of the above solution are taken and the phenol added till a drop of the solution ceases to give a blue colouration with KI and starch (Chandelon, *Bl* [2] 38, 69) — 3 2 or 3 grms phenol are dissolved in three times the molecular proportion of caustic soda. The solution is made up to 500 c c, and 10 c c are put into a small flask warmed to 60°, and one tenth normal iodine solution is allowed to flow in until it is coloured strongly yellow by excess of iodine, by shaking, a red pp is formed. The excess of iodine is estimated by sodium thio sulphate. The quantity of iodine taken up by the phenol multiplied by 0.123518 gives the amount of pure phenol (Messinger & Vortmann, *B* 23, 2753) — 4 To separate phenol from mixtures in poisoning cases the substance is triturated with dilute H_2SO_4 and extracted with alcohol, the alcohol evaporated, and the residue extracted with benzene (Dragendorff & Jacobson, *C* 1886, 828), or the alcoholic extract may be mixed with NaOH aq, evaporated to a small bulk, filtered, and the phenol ppd by HCl aq in a graduated tube (Muter & De Koningh, *An* 12, 191, cf Staveley, *Chem Zeit* 13, 1126) — 5 Phenol may be titrated by adding standard NaOH to a solution containing s-tri-nitro benzene as indicator until a red colour appears (Bader, *Fr* 31, 58)

Salts — Phenol behaves as a very weak acid. It dissolves in KOH aq, but does not expel CO_2 from sodium carbonate in the cold. — $\text{C}_6\text{H}_5\text{OK}$. White hygroscopic needles, v sol alcohol, ether, and water. Oxidises rapidly in air — $\text{C}_6\text{H}_5\text{ONa}^+$. Yields Ph_2O and $\text{C}_{12}\text{H}_{10}\text{O}$ when distilled with NaPO_3 (Niederhäusern, *B* 16,

1123) Sulphur at 200° gives $S_2(C_6H_5OH)_2$.— $Ba(OC_6H_5)_2$ 8aq Crystalline crusts, got by boiling phenol with baryta water and evaporating *in vacuo*— $Ca(OC_6H_5)_2$ * Yields diphenylene oxide and a little benzene on distillation— $Pb(OH)OC_6H_5$ Made by boiling phenol with iitharge (Calvert, *C J* 18, 68)— $TiOOC_6H_5$ Crystals, al sol cold water (Kuhlmann, *J* 1864, 254)— $Hg(OC_6H_5)_2$ 4sq Pp got by adding mercurous chloride to a solution of NaOPh (Pouchet, *C R* 106, 276)— $Al(OC_6H_5)_3$ Made by heating phenol with aluminium and AlI_3 and pouring off the liquid product (Gladstone & Tribe, *C J* 89, 9, 41, 5, Hodgkinson, *C N* 1877, 237) Split up on distillation into Al_2O_3 and Ph_2O , other products being phenol and a ketone $C_{12}H_{10}O$ [97°] (c 280°), V.D. 182.2—Aniline salt $C_6H_5ONH_2Ph$ [87°] (181°) Formed by boiling aniline with phenol (Dale & Schorlemmer, *A* 217, 388, Dyson, *C J* 43, 466, Mylins, *B* 19, 1002) Tables (from alcohol)—*p*-Toluidine salt $C_6H_5ONH_2C_6H_4Me$ [81°] Needles (from ligroin) (Dyson)

Combinations—1 With SO_2 phenol forms an unstable compound which may be distilled at 140° (best in a current of SO_2), and which crystallises in rectangular tablets. If exposed to air it absorbs water and gives off SO_2 . The compound melts between 25° and 30° (A. Hölzer, *J pr* [2] 25, 463)—2 With CO_2 If salicylic acid is heated in a sealed tube for two hours at 260° it is resolved into phenol and CO_2 , but on cooling crystals resembling common salt with sides like staircases are formed. These melt at 37° They are decomposed by heat, and also by water, alcohol, ether, and chloroform, into phenol and CO_2 (A. Kleip, *J pr* [2] 25, 464). The same compound is got from phenol and liquid CO_2 (Barth, *A* 148, 49)

Formyl derivative C_6H_5OCHO Liquid, boiling with decomposition at 180° (Seifert, *J pr* [2] 31, 467)

Acetyl derivative C_6H_5OAc *Phenyl acetate* (195°) at 733 mm (Orndorff, *Am* 10, 368) Formed by boiling an alcoholic solution of phenyl phosphate with KOAc (Williamson & Scrugham, *A* 92, 317, Kreyser, *B* 18, 1716) and by the action of $AcCl$ on phenol (Cahours, *A* 92, 816) or of $POCl_3$ (1 mol) on phenol (3 mols) mixed with $HOAc$ (3 mols) (Nencki, *J pr* [2] 25, 282, Seifert, *J pr* [2] 31, 467) Obtained also by boiling phenol with acetamide (Guareschi, *A* 171, 142) Heavy oil, saponified by KOH *Aq* **Reactions**—1 Sodium acts violently, giving off and forming $EtOAc$, phenol, salicylic acid, $C_{10}H_{12}O_2$ [48°] crystallising from alcohol in needles, and $C_{10}H_{14}O$ [188°] nearly insol in alcohol, but crystallising therefrom in yellow needles (Hodgkinson & W. H. Perkin, jun, *C J* 37, 487, 721)—2 Benzyl chloride after heating for 14 days leaves an oil with strong blue fluorescence which when saponified by alcoholic potash forms $C_{10}H_{10}O$ [39°] (c 295°) and benzylphenol $C_{12}H_{14}O$ [81°] (321°) (H. A. P.)—3 Benzyl chloride and $AlCl_3$ give HCl , toluene, anthracene, Ac_2O , and $OH_2PhC_6H_4OAc$ —4 $NaSEt$ forms $PhONa$ and $EtSAc$ (Seifert)—5 PCl_5 at 100° gives phenyl phosphate and $C_6H_5OCCl_2CCl_2$ [26°] (Michael, *Am* 9, 207)—6 Chlorine in the cold forms $AcCl$, *p*-chloro-phenol, and $C_6H_4Cl(OAc)$ Chlorine at 160° yields $AcCl$,

$C_6H_4Cl_2(OAc)$, and mono- and di- chloro phenols (Seelig, *J pr* [2] 89, 175)—7 Bromine forms $AcBr$ and C_6H_4BrOAc Excess of Br gives $C_6H_4Br_2OH$ and $C_6H_4Br_2OAc$ (Seelig)—8 $BzCl$ and a little $ZnCl_2$ give $PhOBz$ (Doebner, *A* 210, 255)

Propionyl derivative C_6H_5OCOEt [20°] (211°) $SG \frac{1}{2}$ 1 0643, $\frac{1}{2}$ 1 0542 Formed by distilling phenol with excess of propionyl chloride (Perkin, *C J* 55, 546) Large transparent prisms PCl_5 at 100° followed by water gives only $C_6H_4Cl_2O$ (116°) (Michael, *Am* 9, 212)

Butyryl derivative $C_6H_5OCO C_2H_5$ (228°) $SG \frac{1}{2}$ 1 0364, $\frac{1}{2}$ 1 0269 Formed from phenol and butyryl chloride (Pérkin)

Benzoyl derivative C_6H_5OBz *Phenyl benzoate* [89°] (314° cor) HF 61,804 (Stohmann, *J pr* [2] 86, 7) Made by melting benzoic acid (11 g) with phenol (10 g) and gradually adding $POCl_3$ (13 g) The product is washed with dilute KOH and crystallised from dilute alcohol, the yield being fair (12 g) (Rasinski, *J pr* [2] 26, 62, cf Ettinger, *A* 53, 87, Stenhouse, *A* 58, 91, Laurent & Gerhardt, *A* 75, 75, List & Limpriht, *A* 90, 190) Formed also by boiling phenol with benzamide (Guareschi, *A* 171, 141) and from phenyl phosphate and NaOBz (Kreyser, *B* 18, 1719) Monoclinic prisms (from ether alcohol) Not saponified by boiling aqueous KOH, but saponified by alcoholic potash

Salicyl derivative *o* $Oxybenzoic acid$

Methyl derivative C_6H_5OMe *Anisole* Mol w 108 (155°) VD 3.79 (calc 3.73) $SG \frac{1}{2}$ 1 0110 (Pinette, *A* 248, 34), $\frac{1}{2}$ 997 (Vincent, *Bl* [2] 40, 106) CE (0°–10°) 00083 SV 125.2 (R. Schiff, *A* 220, 105) SH 405 at 0° (Schiff, *A* 234, 300) R_{∞} 55.33 (Nasini & Bernheimer, *G* 15, 93) HfP 15,860 HfV 13,830 (Thomsen, *Th*), 32,718 $[C_6O_2=94,000, H_2O=69,000]$ (Stohmann, *J pr* [2] 35, 23) Formed by distilling anisole acid or the methyl derivative of salicylic acid with baryta (Cahours, *A Ch* [3] 2, 274, 10, 353, 27, 439) Made also by heating $PhOK$ with MeI at 110° (Cahours, *A* 78, 225) Prepared by heating NaOPh at 200° in a rapid current of MeCl (Vincent, *Bl* [2] 40, 106) Oil with pleasant odour, v sol alcohol and ether Conc $HfAq$ at 140° splits it up into phenol and MeI (Graebe, *A* 139, 149) At 400° it decomposes into phenol and ethylene (Bamberger, *B* 19, 1820) Chlorine in the cold gives rise to $[1\ 2\ C_6H_4ClOMe$ (c 200°), $[1\ 3\ 6\ C_6H_4Cl_2OMe$ [28°], (233°), and $[1\ 3\ 5\ 6\ C_6H_3Cl_3OMe$ [60°] (249°) In presence of iodine, chlorine forms $[1\ 2\ 3\ 5\ 6\ C_6H_3Cl_4OMe$ [100°] (279°) and also C_6Cl_5OMe [107°] (c 289°) Chlorine with I at 60° forms C_6Cl_4 [220°], C_6HCl [87°] and $C_6H_2Cl_3$ [136°] Chlorine with I acting on boiling anisole forms C_6Cl_4 , $COCl$, HCl , and some CCl_4 Chlorine with $SbCl_3$ forms C_6Cl_4OH [187°] and CCl_4 Chlorine with $SbCl_3$ at 100° gives C_6Cl_4O [107°] (Hugouenq, *A Ch* [6] 20, 504) $AcCl$ in CS_2 in presence of $AlCl_3$ forms CH_3Cl , C_6H_5OMe (Gattermann, *B* 22, 1129), $BzCl$ acts in like manner

Ethyl derivative C_6H_5OEt *Phenetole*. (172°) $SG \frac{1}{2}$ 9822 (Pinette, *A* 248, 35), VD 4.27 (calc. 4.21) CE (0°–10°) 00087 SV 148.5 (R. Schiff, *A* 220, 105) SH 429 (Schiff,

A 284, 300 H.F.p 39,775 (Stohmann) Formed by distilling barium ethyl salicylate and by the action of EtI on C_6H_5OK (Cahours, *A Ch.* [3] 27, 468, *Baly*, *A* 70, 269) Formed also by the action of alcohol and $ZnCl_2$ on phenol (Kastrop, *B* 10, 1855) and by the action of dry alcohol on diazobenzene nitrate or sulphate (Remsen & Orndorff, *Am.* 9, 392)

Preparation—Equal volumes of H_2SO_4 and alcohol are mixed and allowed to cool. The liquid is then neutralised with soda and evaporated until Na_2SO_4 begins to crystallise out. The liquid is poured off from the crystals, mixed with a strong solution of sodium phenylate, and heated in an autoclave at 150° under 7 atmospheres' pressure. The sodic phenylate is made by dissolving phenol in caustic soda solution (S.G. 1.33) In calculating the amount of phenol it is assumed that half the alcohol forms sodic ethyl sulphate (Kolbe, *J pr* [2] 27, 425)

Properties—Oil Yields with fuming HNO_3 a di nitro derivative [87°] At 400° it yields phenol and ethylene $BzCl$ in CS_2 in presence of $AlCl_3$ forms $C_6H_5CO_2C_2H_5OEt$, $AcCl$ acts in like manner (Gattermann, *B* 22, 1129)

Propyl derivative C_6H_5OPr (191°) S.G. $\frac{2}{3}$ 9639 C.E. ($0^\circ-10^\circ$) 00086 S.V. 172 (Pinette, *A* 243, 35) H.F.p 46,574 (Stohmann)

Isopropyl derivative $PhOPr$ (176°) S.G. $\frac{2}{3}$ 958 (Silva, *Z* 1870, 249)

Butyl derivative $C_6H_5OC_4H_9$ (210°) S.G. $\frac{2}{3}$ 95 C.E. ($0^\circ-10^\circ$) 00089 S.V. 195.3

Isobutyl derivative $C_6H_5OC_4H_9Pr$ (198°) S.G. $\frac{2}{3}$ 939 (Riess, *B* 3, 780)

Isoamyl derivative $C_6H_5OC_5H_{11}$ (225°) Formed from $PhOK$ and isoamyl iodide (Cahours, *C R* 32, 61) Oil, lighter than water

Heptyl derivative $C_6H_5OC_7H_{15}$ (267°) S.G. $\frac{2}{3}$ 9319 C.E. ($0^\circ-10^\circ$) 00091 S.V. 270.8

Octyl derivative $C_6H_5OC_8H_{17}$ (283°) S.G. $\frac{2}{3}$ 9221 C.E. 00084 S.V. 296.1 (Pinette, *A* 243, 36)

Vinyl derivative $C_6H_5OC_2H_3$ (c 155°) S.G. $\frac{2}{3}$ 992 From the bromo ethyl derivative and alcoholic potash (Sabaneff, *Bl* [2] 41, 253)

Bromo vinyl derivative $C_6H_5OC_2H_3Br$ Formed from CH_2Br_2 , phenol, and alcoholic potash (Sabaneff, *A* 216, 277) Oil, decomposed by distillation

Di bromo vinyl derivative $C_6H_5OC_2H_3Br_2$ [38°] Got from phenol, C_2H_5Br , and alcoholic potash Gives $PhOC_2H_3Br$ [59°]

Tri-chloro vinyl derivative $C_6H_5OCCl_2CCl_2$ [26.5°] (106° at 12 mm) Formed by the action of PCl_5 upon phenyl acetate Also obtained by the action of potassium phenol upon per chloro ethylene Crystalline solid Volatile with steam (Michael, *B* 19, 840, *Am* 9, 307)

Bromo-ethyl derivative $PhOC_2H_4Br$ [39°] ($240^\circ-250^\circ$) Made from ethylene bromide and sodium phenol in alcoholic solution by digesting for 3 hours at 100° Separated from the ethylene-ether, $(PhO)_2C_2H_4$, by distilling with steam, in which it is much more volatile Colourless crystals with characteristic odour, partly decomposes on boiling V sol alcohol and ether (Weddige, *J pr* [2] 24, 242) **Reactions**—1 Heated at 110° with alcoholic NH_3 it gives crystals of $HN(C_6H_5OPh)_2HBr$, [216°], sl sol alcohol Alkalies liberate from

this salt the free base imido di-ethylene di phenyl ether as an alkaline oil. With HCl it forms a salt, $HN(C_6H_5OPh)_2HCl$ [218°] It also forms a nitrate [197°]—2 With *o* nitro phenol potassium it forms $PhOC_2H_4OC_6H_4NO_2$, crystals in prisms, [86°]—3 Heated with sodium *p* oxybenzoic ether in alcoholic solution it forms $C_6H_5OC_2H_4OC_6H_4CO_2Et$ [81°], which, when saponified by alcoholic potash, yields the acid, $C_6H_5OC_2H_4OC_6H_4CO_2H$ melting at $[196^\circ]$ (Wagner, *J pr* [2] 27, 227)

Chloro-ethyl derivative $PhOC_2H_4Cl$ [25°] (221°) Made from C_2H_5ClBr and $KOPh$ Reacts with $KOEt$ forming $PhOC_2H_4OEt$ (230°), S.G. $\frac{2}{3}$ 1.018 (Henry, *C R* 96, 1233, or (252°), S.G. $\frac{2}{3}$ 1.037 (Sabaneff, *Bl* [2] 41, 253)

Penta-bromo ethyl derivative $PhOC_2H_4Br_5$ [103°-106°] From the di bromo vinyl derivative by successive treatment with alcoholic potash and Br (Sabaneff, *A* 216, 282) **Methylene derivative** $CH_2(OPh)_2$ (294°) (Henry, (299°) (Arnhold, *A* 240, 201) S.G. $\frac{2}{3}$ 1.114 (H), $\frac{2}{3}$ 1.092 (A) Formed from CH_2Br_2 and $KOPh$ (Henry, *A Ch* [5] 86, 269)

Ethylene derivative $C_6H_5(OPh)_2$ [98.5°] (Burr, *Z* [2] 5, 165), [95°] (Lippmann, *C C* 1870, 45) Made from C_2H_5Br and $KOPh$ at 140° Crystals, sl sol alcohol Br yields $C_6H_5Br_2O_2$ [c 100°] H_2SO_4 yields a disulphonic acid, whence $C_6H_5(O_2C_6H_4SO_3)_2Pb$ may be got, crystallising from hot water in laminae

Bromo propyl derivative $C_6H_5OC_3H_7$ ($246^\circ-256^\circ$) Formed from phenol, trimethylene bromide, and $NaOEt$ (Lohmann, *B* 24, 2632) Oil Alcoholic NH_3 at 100° forms $(PhOC_3H_7)_2NH$ which distils above 300° and gives B^+HCl [206°], and the nitros amine $(PhOC_3H_7)_2NNO$ [61°] Aniline forms $PhOC_3H_7NHPH$ [32°] which gives B^+HCl [170°] With $NaOMe$ in $MeOH$ it yields $PhOC_3H_7OMe$ (231°), while alcoholic potash solution forms $PhOC_3H_7OEt$ (329° , or probably 239°) The compound $PhOC_3H_7OH$ (250°) is formed by the action of nitrous acid on $PhOC_3H_7NH_2$ (242°) which is itself obtained by the hydrolysis of $PhOC_3H_7NHCO_2C_6H_5CO_2H$

Trimethylene derivative $CH_2(CH_2OPh)_2$ [61°] A by product in the preparation of the bromo propyl derivative

Allyl derivative $PhOC_3H_5$ (194°) From C_2H_5Br and $NaOPh$ (Henry, *B* 5, 455)

Di chloro allyl derivative $PhOC_3H_5Cl_2$ ($114^\circ-118^\circ$ in vacuo) Got from $PhOCOEt$ and PCl_5 (Michael)

Bromo allyl derivative $PhOC_3H_5Br$ (240°) Got from $CH_2CHBrCH_2Br$ and $KOPh$ (Henry, *Bl* [2] 40, 324) Alcoholic potash forms the propargyl derivative (210°)

Benzyll derivative *v* Phenyl ether of BENZYL ALCOHOL

Glucoside $C_6H_5OC_6H_4(OH)_4CHO$ [172°] Formed by adding acetochlorhydrase to an alcoholic solution of $PhOK$ (Michael, *C R* 89, 855, *Am* 1, 306) Long needles (from hot water), v sol water, sol alcohol and $HOAc$ H_2SO_4 forms a yellow solution Acetic anhydride and $NaOAc$ at 100° yield the tetra acetyl derivative $C_6H_5OC_6H_4(OAc)_4CHO$, which crystallises from alcohol in long white needles

References—AMIDO, AMIDO DI-IMIDO-, BROMO-, BROMO AMIDO-, BROMO IODO-NITRO-, 3 H 2

BROMO-NITRO-, CHLORO-, TRI-CHLORO-IODO-, CHLORO-NITRO-, IODO-, DI-iodo AMIDO-, IODO-NITRO-, and NITRO-phenols

Diphenol *v* Di-oxy-DIPHENYL

Tetraphenol *v* FURFURAL

PHENOL-AZO-COMPOUNDS *v* Oxy-benzene-AZO-COMPOUNDS

Phenol-bisazo- compounds *v* Disazo compounds

PHENOL-BLUE *v* Di-methyl amido phenyl-imide of QUINONE

PHENOL CARBOXYLIC ACID *v* OXY-BENZOIC ACID.

Phenol dicarboxylic acid *v* OXY-PHTHALIC, OXY-ISOPHTHALIC, and OXY-TEREPHTHALIC ACIDS

Phenol tricarboxylic acid *v* OXY-TRIMESIC ACID

PHENOLISATIN *v* Di-oxy DIPHENYLOXINDOLE

PHENOL-PHTHALEIN $C_{20}H_{14}O_4$, *tc*

$C_6H_5 \begin{smallmatrix} \diagup C(C_6H_5OH)_2 \\ \diagdown CO \end{smallmatrix}$ [c 258°] Formed by

heating phenol with phthalic anhydride and conc H_2SO_4 . Obtained also from di-amido-di-phenyl-phthalide by the diazo-reaction (Baeyer, *A* 202, 36, *B* 9, 1230, 12, 642) Small crystals (from alcohol), *v* sol hot alcohol, sl sol water. Its solutions in alkalis and alkaline carbonates are red, but become colourless on neutralisation, and also on addition of excess of KOHAq but not by NH_3 . The red colour of a solution in very dilute ammonia disappears on standing (Long, *Am* 11, 84) Aniline does not give any colour with phenol-phthalain $AgNO_3$ gives a violet pp

Reactions—1 H_2SO_4 at 100° forms a sulphonic acid, but at 200° it yields oxy-anthraquinone—2 PCl_5 yields di-chloro di phenyl phthalide [156°]—3 Zinc-dust and NaOHAq yield phenol-phthalin $CO_2H \cdot C_6H_4 \cdot CH(C_6H_5OH)_2$ —4 Potash fusion gives dioxybenzophenone and benzoic acid—5 NH_3 at 170° forms $C_{20}H_{14}N_2O_4$ [266°]

Di-acetyl derivative $C_{20}H_{12}Ac_2O_4$ [143°] Colourless tables (from wood spirit)

Anhydride $C_{20}H_{12}O_4$ [175°] A by product in the preparation of phenol-phthalain (Baeyer, *A* 212, 347) Needles, insol NaOHAq. Conc H_2SO_4 forms a solution with green fluorescence. Bromine yields $C_{20}H_{12}Br_2O_4$ [258°] Alcoholic potash and zinc dust reduce it to phthalin $C_{20}H_{14}O_2$

Tetra bromo-phenol phthalain $C_{20}H_8Br_4O_4$, *tc* $C_6H_4 \cdot C_2O_2(C_6H_2Br_2OH)_2$ [220°-280°] Formed by adding Br in HOAc to a solution of phenol phthalain in alcohol. Colourless needles (from ether), sl sol alcohol and HOAc. Alkalis form violet solutions, decolourised by excess H_2SO_4 , produces di bromo-oxy-anthraquinone on heating to 150°. Ac₂O yields a di-acetyl derivative [134°] H_2SO_4 containing KNO_3 forms bromosoquinone $C_{12}H_6Br_2O_8$, separating from alcohol chloroform in red crystals with steel-blue lustre, insol water, forming a violet solution in H_2SO_4 , and converted by boiling alcoholic potash into $C_{12}H_6Br_2O_8$ [264°] NH_3 at 170° forms $C_{20}H_{12}Br_2N_2O_4$ (above 280°), whence nitrous acid produces $C_{20}H_{12}Br_2N_2O_8$, and Ac₂O yields $C_{20}H_{10}Ac_2Br_2N_2O_4$ [241°]

PHENOL-PHTHALIDEIN $C_{20}H_{14}O_4$, *tc*

$C_6H_5 \begin{smallmatrix} \diagup C(OH)(C_6H_5OH) \\ \diagdown CO \end{smallmatrix}$ *Dr-oxy-*

phenyl-oxanthranol [212°]. Formed by oxidation of phenol-phthalidin by dilute alkaline $KMnO_4$ (Baeyer, *A* 202, 100) Monoclinic tables (from HOAc), $a \cdot b \cdot c = 46 \cdot 1 \cdot 49$, $\beta = 69^\circ 5'$, *v* sol alcohol and acetone. It forms a pale-yellow solution in KOHAq, and is reprecipitated by HCl. Conc H_2SO_4 forms a violet solution, and on heating produces oxyanthraquinone and pthalic acid. Reduced by zinc-dust and NaOHAq to phenol-phthalidin. PCl_5 yields di-chloro phenyl-oxanthranol $C_{20}H_{12}Cl_2O_4$ [156°]. Bromine gives $C_{20}H_{12}Br_2O_4$ (above 280°), whence $C_{20}H_{12}Ac_2Br_2O_4$ [182°] may be obtained. On adding phenol to a solution of phenol-phthalidein in H_2SO_4 , the violet colour changes to blood-red, and on addition of water a red amorphous pp is thrown down. This pp forms a deep-violet solution in alkalis, and yields a crystalline bromo-derivative. A mixture of NH_3 and alcohol at 160° converts the red pp into $C_{20}H_{14}N_2O_4$ [260°], crystallising in yellow needles

Acetyl derivative $C_{20}H_{12}Ac_2O_4$ [109°] Monoclinic prisms, $a \cdot b \cdot c = 2 \cdot 78 \cdot 1 \cdot 44$, $\beta = 77^\circ 2'$

PHENOL-PHTHALIDEIN CHLORIDE *v*

Di-CHLORO PHENYL-OXANTHRANOL

PHENOL-PHTHALIDIN *v* Di-OXY-PHENYL

ANTHRANOL

PHENOL-PHTHALIN $C_{20}H_{14}O_4$, *tc*

$CO_2H \cdot C_6H_4 \cdot CH(C_6H_5OH)_2$ *Dr-oxy-tri-phenyl-methane carboxylic acid* [225°] Formed by reduction of phenol-phthalain with zinc-dust and NaOHAq (Baeyer, *A* 202, 80, 212, 350) Small needles. Forms a colourless solution in potash, coloured red by K_2FeCy , through formation of phenol-phthalain. Yields a di-acetyl derivative [146°] Sodium-amalgam yields phenol-phthalol $CH(OH) \cdot C_6H_4 \cdot CH(C_6H_5OH)_2$ [190°], which gives a tri-acetyl derivative [40°]

Anhydride $C_{20}H_{12}O_4$ [217°] Formed by reducing the anhydride of phenol-phthalain. Small needles (from dilute alcohol)

Di-chloro-phenol-phthalin *Anhydride*

$O \begin{smallmatrix} \diagup C_6H_4Cl \\ \diagdown C_6H_4Cl \end{smallmatrix} OH$ $C_6H_5 \cdot CO_2H$ [226°-230°] Got

by the action of zinc and NaOHAq on the compound $C_{20}H_{12}Cl_2O_4$, derived from fluorescein (*v* vol II p 558) Small needles (from dil alcohol)

Tetra-bromo-phenol-phthalin

$CO_2H \cdot C_6H_4 \cdot CH(C_6H_2Br_2OH)_2$ [205°] Formed by bromination. Crystals (from benzene) Yields a di-acetyl derivative [166°] (Baeyer, *A* 202, 80) Conc H_2SO_4 converts it into tetra bromo phenol-phthalidin $C_{20}H_8Br_4O_4$, which crystallises from alcohol in golden needles, and is converted by oxidising agents into tetra bromo-phthalidein $C_{20}H_8Br_4O_4$. Tetra-bromo-phenol-phthalidin is tetra-bromo di-oxy phenyl-anthranol, and its di-acetyl derivative crystallises in needles [256°]

PHENOL-QUINOLINE *v* OXY-PHENYL-QUINOLINE

PHENOLS Compounds containing hydroxyl united to carbon, which forms part of a benzene nucleus (*v* BENZENE and ALCOHOLS) They much resemble the tertiary fatty alcohols, being more acid in character than primary alcohols. Thus phenols and tertiary alcohols form hardly any acetate on heating with HOAc. Phenols in which hydrogen in the *o*- or *p*-position has been displaced by NO_2 , or by a halogen are even more strongly acid in character (*v* *g* picric acid)

Formation.—1. By potash-fusion from sul-

phonic acids —2 By the diazo-reaction from amido-compounds (*v* DIAZO-COMPOUNDS) —3 By distillation of oxyacids either alone or with lime —4. Halogens in a benzene nucleus may sometimes be displaced by hydroxyl by heating with KOHAq in sealed tubes, thus *o*- and *p*-chloro-nitro-benzenes give *o*- and *p*-nitro phenols by this treatment

Properties of Phenol and its homologues — Dissolve in NaOHAq but not in Na₂CO₃Aq Phenols do not give rise on oxidation to an acid or aldehyde containing the same number of atoms of carbon in the molecule They do not exchange H⁺ or K⁺ on treatment with KSH, nor do they form ammonium salts They do not react (like alcohol) with HCl FeCl₃ colours aqueous solutions of phenols, and products of oxidation are often produced. H₂SO₄ to which 6 p.c. of KNO₃ has been added gives with a solution of a phenol in H₂SO₄ a brown colour, changing on warming to green and blue (Liebermann, *B* 7, 248, 806, 1098)

Reactions —1 Readily undergo bromination, chlorination, and nitration —2 H₂SO₄ forms sulphonic acids, the isomeric sulphuric acids being very unstable —3 Sodium and CO₂ form oxyacids on heating (Kolbe) —4 Chloroform and NaOHAq form, on boiling, oxy-aldehydes (Tiemann a Reimer, *B* 9, 824) —5 CCl₄ and alcoholic potash at 100° form oxyacids —6 Ammoniacal ZnCl₂ at 200°–300° converts phenols into amines (Merz a Weith, *B* 13, 1298, 20, 544) —7 Solutions of NaHCO₃ and (NH₄)HCO₃ yield oxyacids on heating with dioxyphe-nols (Kostanecki, *B* 18, 3203) —8 Distillation over zinc-dust reduces the hydroxyl to hydrogen, forming the corresponding hydrocarbons —9 Alkyl chlorides and anhydrides yield alkyl derivatives of phenols, especially if a little powdered zinc be added (Schisparelli, *G* 11, 69) When an alkyl chloride is used oxy ketones are also often formed through the alkyl entering the benzene nucleus —10 Organic acids mixed with phenols readily yield alkyl derivatives of the phenols on adding POCl₃ or on heating in presence of ZnCl₂ (Rasinski, *J pr* [2] 26, 62, Nencki, *M* 10, 906) Alcohols in presence of ZnCl₂ form alkyl ether and higher homologous phenols —11 Alkyl derivatives of phenols split up on distillation into phenols and olefines (Bamberger, *B* 19, 1819) —12 The acid sulphuric ethers of the form C₆H₄XO.SO₃H derived from *o* and *p*-alkyl-phenols are oxidised by alkaline KMnO₄ to *o*- and *p*-oxy benzoic acids (Heymann, *B* 19, 704) —13 Alkaline solutions of phenols absorb atmospheric oxygen, forming dark coloured products —14 Homologues of phenol frequently yield oxyacids on fusion with potash, the alkyl side-chain being oxidised to carboxyl 15 Aldehydes in presence of dehydrating agents form compounds of the form XOCH(OY), which change to XOCH(Y' OH). Thus (β) naphthol and benzoic aldehyde give PhCH(OC₆H₅)₂, changing to PhCH(C₆H₅OH)₂, which is at once converted into the anhydride PhCH<C₆H₅OH>O (Baeyer, *B* 5, 26, Clausen, *B* 19, 5316). —16 Phthalyl chlorides usually forms phthalic ethers (R Meyer, *B* 24, 2600). —17 Iodine and potash give di- and tri- iodo- derivatives (Messinger a-Vortmann, *B* 22, 2315) —18 Nitrous

acid displaces hydrogen in the *o*- or *p*- position by nitrosyl The products are identical with the mono-oxims formed by warming quinones with hydroxylamine hydrochloride, but yield nitro-phenols on oxidation with alkaline K₂FeO₄. —19 Diazo salts readily form azo-compounds (*q v*) 20 Diazo-amido-compounds form oxy azo-compounds Thus C₆H₄Cl.N₂.NH.C₆H₄Cl heated with phenol on a water bath yields C₆H₄Cl.N₂.C₆H₄OH and C₆H₄Cl.NH₂ (Heumann a Oeconomidis, *B* 20, 904) —21 Benzotrichloride acting on phenols in which there is an atom of hydrogen in the *p* position to hydroxyl yields colouring matters related to aurin (Doebner, *A* 217, 228, 257, 56)

PHENOL *o*-SULPHONIC ACID C₆H₄SO₃H C₆H₄(OH)SO₃H Formed, together with the *p* acid, by the action of H₂SO₄ on phenol, especially in the cold (Kekulé, *Z* 1867, 199, *B* 2, 330, Faust, *Z* 1871, 339, Armstrong, *C J* 24, 1112, 25, 12, 857) Prepared by dropping H₂SO₄ on dry phenol at –10°, allowing the product to stand for a week, and then distilling, treating with BaCO₃ and KHCO₃ successively, and crystallising the K salt (Allan le Cannu, *C R* 109, 225) Crystals (containing 1½ aq), melting at 50° and decomposing at a higher temperature, with liberation of H₂SO₄ Sol ether When heated with water it changes to phenol *p*-sulphonic acid (Post, *A* 205, 64) Possesses powerful antiseptic properties (Serrant, *C R* 103, 1079)

Reactions —1 Potash-fusion gives some pyrocatechin The action begins about 280° and attains a maximum (20 p.c) at 320° Soda-fusion also gives pyrocatechin The action begins at 310° At 330° the yield is 6 p.c Fusion with a mixture of potash and soda gives a still smaller yield (1 p.c) of pyrocatechin (Degener, *J pr* [2] 20, 308) —2 Bromine-vapour acting at –8° on the K salt yields bromo and di bromo derivatives —3 BzCl forms PhOBz on heating with the K salt (Solommanoff, *Z* [2] 5, 296) —4 Heated with dilute HCl at 140° it is split up into phenol and H₂SO₄ (Armstrong)

Salts —KA' 2aq [235°–240°] Colourless trimetric prisms, sol water —NaA' 1½aq —BaA', aq needles *S* 25 at 100° —BaA', 2aq —PhA', aq tables, sl sol water —The aniline salt yields phenol and amido-benzene *p*-sulphonic acid on distillation (Kopp, *B* 4, 978)

Methyl derivative C₆H₄(OMe)SO₃H Formed from the acid, KOH, MeI, and MeOH (Kekulé) —KA' aq needles, more soluble than its *p*-isomeride (Hartinger, *M* 4, 173) Yields a chloride [55°]

Ethyl derivative Forms a K salt crystallising in plates

Phenol m-sulphonic acid C₆H₄(OH)SO₃H Formed by potash-fusion from benzene *m*- or *p*-disulphonic acid (Barth a Senhofer, *B* 9, 969) Needles (containing 2aq) FeCl₃ gives a violet colour Yields resorcin on fusion with potash —NaA' aq —KA' aq [200°–210°] Efflorescent needles —K₂C₆H₄SO₃ aq —PhA', 8aq. tables, v sol water —BaA', ½aq —CaA', 6aq green tables

Ethyl derivative C₆H₄(OEt)SO₃H Got by the action of KOH and EtI on the acid (Dehlis a Lagan, *B* 23, 3393) Deliquescent needles Yields a chloride [38°] and an amide [131°], and is reduced by zinc-dust and dilute H₂SO₄ to C₆H₄(OEt)SH (289°) —Salts KA' aq BaA', ½aq needles (from water) —CaA', 8aq

Phenol *p*-sulphonic acid $C_6H_4(OH)SO_3H$ Formed by warming phenol with H_2SO_4 or $ClSO_3H$. Formed also from amido benzene *p*-sulphonic acid by the diazo-reaction. Highly deliquescent hydrated needles (Le Canu, *C R* 103, 388).

Reactions—1 Bromine acting on a cold solution forms mono- and di-bromo-derivatives and finally tri-bromo-phenol (Le Canu)—2 Boiling $HIAq$ gives phenol (Benedikt & Bamberger, *M* 12, 4)—3 Potash fusion forms a little resorcin (Luncke, *J pr* [2] 8, 43)—4 PCl_5 gives *p*-di-chloro benzene and $C_6H_4ClOPOCl_2$ (Kekulé, *B* 5, 875, 6, 943)—5 MnO_2 and H_2SO_4 yield quinone (Schrader, *B* 8, 759)—6 Dilute HCl at 150° forms phenol and H_2SO_4 .

Salts— KA' [400°] Six sided trimetric lamellae $abc = 879 \times 11002$ — NaA' 2aq monoclinic prisms (Shadwell, *J* 1881, 874)— BaA' 3aq S 50 at 100° — $BaC_2H_5SO_3$ 2aq— PbA' 2aq— CoA' 8aq (Freund, *A* 120, 85)— NiA' 8aq— CuA' 10aq— AgA' —Aniline salt Plates [170°] Decomposed at 190° into phenol and amido benzene *p*-sulphonic acid—*o*-Toluidine salt [c 192°] S 107 at 14° (Lecco, *J* 1874, 747)—*p*-Toluidine salt [202°] S 52 at 17° .

Benzoyl derivative $C_6H_4(OBz)SO_3H$ Formed from $PhOBz$ and SO_3 (Engelhardt & Latschinoff, *Z* 1868, 76)— KA' long needles (from water)— CaA' — BaA' — PbA' 2aq— CuA' 6aq— AgA' .

Methyl derivative $C_6H_4(OMe)SO_3H$ Got by methylation. The K salt forms needles.

Ethyl derivative $C_6H_4(OEt)SO_3H$ Formed from the acid, KOH , and EtH (Kekulé, *Z* 1867, 200), and also, together with the *o*-iso meride, by sulphonating phenetole (Lippmann, *C R* 68, 1332)— BaA' 4aq crystals, v sl sol hot water— KA' aq needles (from alcohol).

Phenyl ether PhA' Formed from phenol (2 mols) and $ClSO_3H$ (Engelhardt & Latschinoff, *Z* 1869, 298). Syrup, split up by water into phenol and phenol sulphonic acid.

Anhydride $C_{12}H_8S_2O_6$ Formed from the acid and $POCl_3$ (Schiff, *A* 178, 171). Powder, v sol water, v sl sol $HClAq$. Coloured violet-red by $FeCl_3$. Ppts gelatin and alkaloids. Yields an acetyl derivative $C_{12}H_8AcS_2O_6$.

Phenol disulphonic acid $C_6H_3(OH)(SO_3H)_2$ [1 2 4] Formed by warming phenol with H_2SO_4 and SO_3 , and by allowing separate vessels of phenol and fuming H_2SO_4 (S G 1 85) to remain for six months *in vacuo*, the acid absorbing the vapour of phenol (Kekulé, *Z* 1866, 693, Le Canu, *C R* 109, 442). Formed also from diazo-benzene sulphate and fuming H_2SO_4 (Griess, *A* 137, 69, Armstrong & Prevost, *B* 6, 664). Nodular groups of deliquescent needles, v e sol water and alcohol. HNO_3 yields picric acid. Potash fusion gives pyrocatechin and its sulphonic acid, soda fusion forms also proto-catechuic acid (Barth & Schmidt, *B* 12, 1260). $FeCl_3$ gives a red colour (Stadeler, *A* 144, 299).

Salts— KA' aq. Needles, decomposing about 270° — S 38 at 100° — BaA' 4aq S 19 6 at 15° — $Ba_2(C_2H_5S_2O_6)_2$ 6aq— $Ba_3(C_2H_5S_2O_6)_3$ 3aq Sl sol water— $Pb_2(C_2H_5S_2O_6)_2$ 6aq Sl sol water— AgA' plates, v sol water.

Methyl derivative $C_6H_3(OMe)(SO_3H)_2$ Formed by sulphonation of anisic acid (Zervas,

A 103, 342), and also from $SO_3(C_2H_5OMe)_2$ and conc H_2SO_4 at 170° (Annaheim, *A* 172, 47)— BaA' 4aq monoclinic crystals.

Ethyl derivative $C_6H_3(OEt)(SO_3H)_2$ [1 2 4] Formed from amido benzene disulphonic acid by the diazo-reaction (Zander, *A* 198, 25). Very deliquescent needles. Yields a chloride [108°] and an amide [233°]— KHA' aq— BaA' 2aq— BaA' 3aq v sol water.

Ethylene derivative $C_6H_3(O C_2H_4 SO_3H)_2$ Formed from $C_2H_4(OPh)_2$ and H_2SO_4 at 120° (Lippmann, *Z* 1869, 447)— BaA' 7aq— PbA' 7aq plates, $insol$ cold water.

Phenol disulphonic acid An unstable acid is formed by heating the trisulphonic acid with KOH (3 pts) and some water at 150° (Senhofer, *J* 1879, 749). Yields K_2A' 3aq— BaA' 4aq— PbA' 4aq long prisms, v sol water.

Phenol trisulphonic acid $C_6H_3(OH)(SO_3H)_3$ [1 2 4 6] Formed by heating phenol (6 pts) with H_2SO_4 (30 pts) and P_2O_5 (15 pts) at 180° (Senhofer, *A* 170, 110). Formed also by heating $SO_3(C_2H_5OH)_2$ with fuming H_2SO_4 at 190° (Annaheim, *A* 172, 28). Needles or short prisms (containing 3 aq). Coloured red by $FeCl_3$ — Na_3A' 3aq— K_3A' 4aq— $K_2C_2H_5S_3O_6$ 2aq— Ba_3A' 4aq scales— Ba_4A' 10aq plates— Cd_4A' 7aq— $Pb_4C_2H_5S_3O_6(OH)_4$ 4aq crystal line powder— Ag_4A' 14aq slender needles.

References—AMIDO-, BROMO-, BROMO NITRO-, DI IODO-, IODO NITRO- PHENOL SULPHONIC ACIDS.

PHENOQUINONE $C_{12}H_6O_4$ [71°] Formed from phenol (1 mol) and quinone (2 mols), or by boiling phenol with aqueous CrO_3 (Wichelhaus, *B* 5, 248, 846, Nietzki, *A* 215, 134, Hesse, *A* 200, 232, Erhart, *Ar Ph* [3] 8, 481). Red needles with green lustre, sol cold water, alcohol, ether, and ligroin. Reduced by SO_2 to hydroquinone. Bromine forms di-bromo phenol.

PHENOSAFRANINE *v* SAFRANINE.

PHENOSE $C_{12}H_{10}O_6$ Formed by the action of aqueous ClO_H on benzene in the dark, the resulting $C_6H_4Cl_2O_3$ [10°] being extracted with ether and heated with very dilute Na_2CO_3 (Carius, *A* 136, 323). Formed also by the electrolysis of toluene mixed with alcohol and dilute H_2SO_4 (Renard, *C R* 92, 965). Deliquescent amorphous mass, v sol water and alcohol, $insol$ ether. Has a sweet taste. Decomposes above 100° . Turned brown by acids and alkalis. Prevents ppn of cupric sulphate by potash, and reduces Fehling's solution. Does not undergo alcoholic fermentation. Reduced by HI to hexyl iodide. HNO_3 yields oxalic acid. Ammoniacal lead acetate ppts $C_6H_4PbO_6$.

PHENOXYACETIC ACID *v* Phenyl derivative of GLYCOLLIC ACID, vol II p 638, and CHLORO-PHENOXYACETIC ACID.

PHENOXYACETIC ACRYLIC ACID *v* Carbonyl methyl derivative of COUMARIC ACID.

PHENOXYACETIC-CARBOXYLIC ACID *v* CARBOXYPHENOXYACETIC ACID.

γ -PHENOXY BUTYRIC ACID $C_{10}H_{12}O_4$ v $C_6H_5OCH_2CH_2CH_2CO_2H$ [60°] Made from the nitrile and $HClAq$ at 100° . Silvery plates, $insol$ water, v sol alcohol— AgA' white crystalline powder, decomposing at 200° .

Nitrile $PhO C_3H_6CN$ [46°] (288°). Formed from $PhO C_3H_6Br$, alcohol, and aqueous $KOCy$ (Lohmann, *B* 24, 2640), and from γ -chlorobutyronitrile and $NaOPh$ (Gabriel, *B* 24 8231).

White needles Conc H_2SO_4 converts it into $\text{C}_6\text{H}_5(\text{SO}_3\text{H})\text{O}\text{C}_6\text{H}_5\text{CONH}_2$ [211°], a crystalline powder which gives BaA^+ , as white crystals insol Aq Yields $\text{PhO}\text{C}_6\text{H}_5\text{NH}_2$ (256°) on reduction

PHENPROPYL-COMPOUNDS v. PROPYL-

PHENYL-COMPOUNDS

PHENOPYLAMINE v. AMIDO PHENYL-

PROPANE

PHENUVIC ACID v. PHENYL METHYL FUR-FURANE CARBOXYLIC ACID

PHENYL The monovalent radicle C_6H_5 . It is more chlorous in character than methyl and ethyl (V Meyer, *B* 20, 534)

Diphenyl $\text{C}_{12}\text{H}_{10}$ vs C_6H_5 , C_6H_5 Mol w 154 [70°] (254°) (Schultz, *A* 174, 205), (253° at 716 mm) (Bamberger a Lodter, *B* 20, 3077) H C v 1,492,800 (Stohmann, *J pr* [2] 40, 86), 1,508,700 (Berthelot, *Bl* [2] 47, 865, *A Ch* [6] 10, 448) H C p 1,494,300 (S), 1,510,100 (B) H F -19,800 (S), -37,100 (B) SVS 154.25 (Schiff). Occurs in the coal tar oils boiling at 200°-300° (Buchner, *B* 8, 22, Schulze, *B* 17, 1203)

Formation—1 By adding slices of sodium to bromo benzene and distilling after 24 hours (Fittig, *A* 121, 361, 132, 201, Schultz, *A* 174, 201, Engelhardt a Latschinoff, *Z* [2] 7, 259)—2 From chloro benzene and sodium amalgam (Cl.urch, *C J* 16, 76)—3 With other hydrocarbons by passing benzene vapour through a red hot tube (Berthelot, *Z* 1866, 707), and formed consequently in many reactions in which benzene is produced at a high temperature—4 By passing benzene vapour mixed with SbCl_5 or SnCl_4 through a red hot tube (Watson Smith, *C J* 30, 30, Aronheim, *B* 9, 1898)—5 A product of the passage of azobenzene through red hot tubes (Claus, *B* 8, 37)—6 By the action of red hot zinc dust on di phenylene ketone-oxide 7 By heating diphenylene ketone dicarboxylic acid with lime (Bamberger a Hooker, *A* 229, 155)—8 From benzidine by the diazo-reaction (Griess, *Tr* 1864 [8] 692)—9 By heating phenol with K at 240° and ppg with water—10 By subjecting bromo benzene to an electric current, using zinc for the positive electrode (Christomanos, *G* 5, 402)—11 In small quantity in the decomposition of diazobenzene salts by alcohol (Griess) and by SnCl_4 (Culman a Gasiorowski, *J pr* [2] 40, 97)

Preparation—1 By slowly passing benzene through a red hot iron tube By successive passage of the unchanged benzene through the tube the yield may be raised to 97 p c of the theoretical (Adam, *A Ch* [6] 15, 224, *C Luddens*, *B* 8, 870)—2 31 g aniline are dissolved in 40 g conc. H_2SO_4 and 150 g water, and diazotized with 31 g NaNO_2 100 g 90 p c alcohol is then added, followed gradually by 50 g finely-divided copper After stirring for one hour it is finally distilled with steam The yield is 6-7 grms Iron and zinc-dust may also be used The temperature must not rise above 80°-40° (L Gattermann, *B* 23, 1226)

Properties—Indescent nacrous scales (from alcohol), v sol hot alcohol and ether

Reactions—1 Readily yields di-bromo- and di-nitro-derivatives Chlorine, in presence of SbCl_5 , yields $\text{C}_{12}\text{H}_8\text{Cl}_2$ and $\text{C}_{12}\text{H}_6\text{Cl}_4$ (Kramers, *A* 189, 142)—2 Heated in a stream of Cl in

presence of I the product is $\text{C}_{12}\text{Cl}_{10}$ (Ruoff, *B* 9, 1048, of Merz a Weith, *B* 18, 2881)—3 A mixture of diphenyl and ethylene passed through a red hot tube yields benzene, styrene, anthracene, and phenanthrene (Barbier, *A Ch* [5] 7, 582)—4 CrO_3Cl_2 in HOAc oxidises it to benzoic acid (Carstanjen, *J pr* [2] 2, 79)—5 AlCl_3 yields, on heating, benzene and a pitchy mass (Friedel a Crafts, *C R* 100, 692) Diphenyl (15 pts) heated with AlCl_3 (1 pt) and CH_2Cl_2 (10 pts) yields diphenylene-methane (fluorene) [118°] and $(\text{C}_6\text{H}_5\text{O}_2\text{H})_2\text{CH}_2$ (810°-820°) (Adam, *Bl* [2] 47, 686)—6 AlCl_3 and COCl_2 give $(\text{C}_6\text{H}_5\text{O}_2\text{H})_2\text{CO}$ [229°] *Acetyl chloride* and AlCl_3 give $\text{C}_6\text{H}_5\text{C}_6\text{H}_5\text{COCH}_3$ [121°]— EtCl and AlCl_3 give $\text{C}_6\text{H}_5\text{C}_6\text{H}_5\text{C}_2\text{H}_5$ (285°) and $\text{C}_6\text{H}_5\text{C}_6\text{H}_5\text{Et}$ (a. 307°) In all these cases the compounds are m-derivatives (Adam, *A Ch* [6] 15, 224)

Dihydrates $\text{C}_{12}\text{H}_{12}$ (249°). Got by the action of boiling alcoholic potash on $\text{C}_6\text{H}_5\text{Br}$, which is formed from $\text{C}_6\text{H}_5\text{I}$ and Br (Bamberger a Lodter, *B* 21, 843) Oil, volatile with steam, sol ether Yields $\text{C}_{12}\text{H}_{12}\text{Br}_2$

Tetra hydrate $\text{C}_{12}\text{H}_{14}$ (245°) at 716 mm Got by reducing diphenyl dissolved in amyl alcohol with sodium (Bamberger a Lodter, *B* 20, 3077) Liquid Yields $\text{C}_{12}\text{H}_{14}\text{Br}_2$ and $\text{C}_{12}\text{H}_{12}\text{Br}_2$ [184°]

References—AMIDO, BROMO, BROMO AMIDO, BROMO NITRO, CHLORO, DI CHLORO DI NITRO, DI IODO, NITRO, and OXY-, DIPHENYL

PHENYL-ACETAMIDE v. Acetyl derivative of ANILINE and Amide of PHENYL-ACETIC ACID

PHENYL-ACETAMIDINE $\text{C}_6\text{H}_5\text{N}_2$ vs $\text{C}_6\text{H}_5\text{CH}_2\text{C}(\text{NH})\text{NH}_2$ Formed from the thio amide $\text{CH}_2\text{Ph CS NH}_2$ by the action of ammonia and atmospheric oxygen, or of ammonia and HgCl_2 (Bernthsen, *B* 8, 1319, 9, 429, *A* 184, 321) Formed also from $\text{CH}_2\text{Ph C}(\text{NH}_2)\text{OEt}$ and ammonia (Luckenbach, *B* 17, 1423) Crystals, decomposed by heat, v sol alcohol, ether, and warm water Absorbs CO from the air Decomposed by heating with water and alcohol into NH_3 and $\text{CH}_2\text{Ph CONH}_2$

Salts— $\text{B}^+\text{HCl aq}$ — $\text{B}^+\text{H}_2\text{PtCl}_4$ — $\text{B}^+\text{H}_2\text{SO}_4$ tables, v sol water and alcohol— $\text{B}^+\text{H}_2\text{S}_2\text{O}_8$ [198°] Needles (from alcohol)— B^+HNO_3 Monoclinic crystals (Lossen, *A* 265, 165)— B^+HOAc [193°] Stellate groups of needles (from alcohol)— $\text{B}^+\text{H}_2\text{C}_2\text{O}_4$ prisms

Di-acetyl derivative

$\text{C}_6\text{H}_5\text{CH}_2\text{C}(\text{OAc})\text{NHAc}$ [178°] Four sided tables (from water), al sol alcohol and ether

Phenyl-acetamidine $\text{CH}_2\text{C}(\text{NPh})\text{NH}_2$ Formed from acetonitrile and aniline hydrochloride at 170° (Bernthsen, *A* 184, 358) Oil $\text{B}^+\text{H}_2\text{C}_2\text{O}_4$ v e sol water

s-Di phenyl-acetamidine $\text{C}_{12}\text{H}_{12}\text{N}_2$ vs $\text{CH}_2\text{C}(\text{NPh})\text{NHPh}$ *Ethyl-di-phenyl-amidine* [132°] **Formation**—1 From aniline (8 pts), HOAc (2 pts) and PCl_5 (2 pts) (Hofmann, *C R* 62, 729)—2 From acetanilide (2 mols) and PCl_5 (1 mol) (Lippmann, *B* 7, 541, Wallach, *B* 8, 1567)—3 From acetonitrile and aniline hydrochloride at 240° (Bernthsen).—4 By passing HCl into heated acetanilide (Wallach, *B* 15, 210)—5 By-product in the preparation of this acetanilide from acetanilide and P_2S_5 .—6 By distillation of thioacetanilide (Jacobsen, *B* 19, 1073)

Preparation—By heating a mixture of equal weights of acetanilide and aniline hydrochloride.

Properties—Slender needles or thin prisms, sl sol. alcohol, forming a neutral solution, nearly insol water, v. sol ether. Conc H_2SO_4 forms amido-benzene *p* sulphonic acid and acetic acid. Tin and HCl reduce it to aniline and HOAc. Fuming HNO_3 forms the insoluble nitrate of a di nitro-derivative $\text{C}_{12}\text{H}_9\text{N}_2\text{O}_6\text{HNO}_3$, [182°] (Biedermann, *B* 7, 540) COCl_2 at 60° forms $\text{C}_{12}\text{H}_9\text{N}_2\text{O}_6\text{Cl}_2$, whence NaOEt yields $\text{C}_{12}\text{H}_9\text{N}_2\text{O}_6$, [91°] crystallising from ether (Loeb, *B* 18, 2427, 19, 2340). Excess of COCl_2 in benzene yields $\text{C}_{12}\text{H}_9\text{N}_2\text{O}_6$ [118°], which is decomposed by dilute HCl into aniline and phenyl cyanate. Cyanogen passed into an ethereal solution forms $\text{C}_{12}\text{H}_9\text{N}_2\text{O}_6$, a white crystalline powder [165°]. EtI yields only $\text{CH}_3\text{C}(\text{NPh})\text{NPhEt}$, whence MeI followed by moist Ag_2O produces strongly alkaline $\text{CH}_3\text{C}(\text{NPh})\text{NPhEtMe}(\text{OH})$ — B^+HCl tables, m sol water— $\text{B}^+\text{H}_2\text{PtCl}_4$ — B^+HNO_3 .

u-Di phenyl-acetamide $\text{CH}_3\text{C}(\text{NH})\text{NPh}_2$, [68°] Formed from diphenylamine hydrochloride and acetonitrile by heating for a week at 150° (Bernthsen, *A* 192, 25). Monoclinic tablets or thick prisms, a 0.6142 b 1.1222 c 1.222 V sol alcohol, forming an alkaline solution. CS, at 100° gives di phenyl-thioacetamide. The hydrochloride is v e sol water— $\text{B}^+\text{H}_2\text{PtCl}_4$ — B^+HNO_3 . Colourless tablets.

Di phenyl-acetamide

$\text{C}_{12}\text{H}_{11}\text{CH}_3\text{C}(\text{NPh})\text{NH}_2$ *Phenyl phenyl acetamide* [c 128°] Formed, with H_2S , by heating $\text{C}_6\text{H}_5\text{CH}_2\text{CH}_2\text{CSNH}_2$ with aniline hydrochloride (B). Formed also by heating phenyl-acetonitrile (benzyl cyanide) with aniline hydrochloride at 230° and by the action of iodine on a mixture of aniline and phenyl thioacetamide. Small needles or laminae, v sol water, v sol alcohol and ether. May be sublimed. Boiling dilute alcohol decomposes it into aniline and $\text{CH}_3\text{PhCONH}_2$.

Tri-phenyl-acetamide

$\text{C}_6\text{H}_5\text{CH}_2\text{C}(\text{NPh})\text{NPh}_2$ [108°] Formed from $\text{C}_6\text{H}_5\text{CH}_2\text{C}(\text{NH}_2\text{Cl})\text{OEt}$ (1 mol) and alcoholic aniline (3 mols) (Luckenbach, *B* 17, 1427). Plates (from alcohol)— $\text{B}^+\text{H}_2\text{PtCl}_4$, plates.

Reference—DI-BROMO DI PHENYL ACETAMIDE.

PHENYL ACETATE *v* *Acetyl derivative of PHENOL*.

PHENYL-ACETCHLORAMIDE *v* *Chloracetamide in the article ANILINE*.

PHENYL-ACETIC ACID

$\text{C}_6\text{H}_5\text{CH}_2\text{CO}_2\text{H}$ *Alphatoluylic acid* Mol w 136 [77°] (265.5° cor) SG $\frac{1}{4}$ 1.0778, $\frac{1}{2}$ 1.0834 H C p 933,200 (Stohmann, *J pr* [2] 40, 134). H C v 932,600 HF 94,800 (Stohmann), 59,000 (von Rechenberg).

Formation—1 By saponification of its nitrile (Cannizzaro, *A* 96, 246, *C R* 52, 966, 54, 1225)—2 By boiling vulpic acid with baryta water (Möller a Strecker, *A* 118, 64)—3 By reducing mandelic acid with conc HIAq and P (Orum Brown, *Proc Roy Soc Edinb* 5, 409)—4 By the putrefactive fermentation of proteids (Salkowski, *B* 13, 649, *H* 2, 420, 9, 507).

Preparation—By converting benzyl chloride into the nitrile (benzyl cyanide), mixing 100 grms of the cyanide with 800 grms of a mixture of 8 vols of H_2SO_4 and 2 vols Aq , and heating till

gas bubbles begin to rise. A violent reaction occurs, heat being again applied when this subsides, until no further reaction takes place. The acid partly crystallises out, and is partly extracted with ether (W Slædel, *B* 19, 1949).

Properties—Thin laminae, sl sol cold water, v. sol hot water, alcohol, and ether.

Reactions—1 Yields benzoic aldehyde, formic acid, and CO_2 when oxidised by chromic acid mixture—2 HNO_3 yields *o*- and *p*-nitro acids (Pirogoff, *B* 5, 332)—3 *Electrolysis* of a dilute solution gives benzoic aldehyde and benzoic acid and two neutral substances [93°] and [115°] (Slawik, *B* 7, 1051)—4 *Ozonised oxygen* yields $\text{PhCH}_2\text{COOCH}_2\text{Ph}$ (318°)—5 When administered to animals it appears in the urine as phenyl aceturic acid, and increases the amount of urea (Salkowski, *B* 12, 653, *H* 12, 222)—6 *Phthalic anhydride* and HOAc form benzylidene phthalide on heating. Tetra chlorophthalic anhydride (40 pts), phenyl acetic acid (20 pts), and NaOAc (1 pt) form, in like manner, $\text{CHPhC}_2\text{O}_2\text{C}_6\text{H}_5\text{Cl}_4$, melting above 360°, whence NaOH forms $\text{CH}_2\text{PhCO}_2\text{C}_6\text{H}_5\text{Cl}_4$ [175°] (Gabriel, *B* 20, 2869). Di-chloro phthalic anhydride forms the corresponding compound $\text{CHPhC}_2\text{O}_2\text{C}_6\text{H}_5\text{Cl}_2$ [210°], whence alkalis yield $\text{C}_6\text{H}_5\text{CH}_2\text{Cl}_2\text{O}$ [117°]—7 HIAq and P at 200° form $\text{C}_6\text{H}_5\text{PO}$, [136°], which crystallises from water in needles, and gives CaA'' 2aq, BaA'' 2aq, and AgA'' (Guye, *J* 1884, 468).

Salts— CaA , 3aq— CaA' , 2aq— BaA' , 3aq v sol water— PbA' , aq— AgA' , small laminae. *Methyl ether* MeA' [220°] SG $\frac{1}{2}$ 1.044 Liquid (Radziszewski, *Z* [2] 5, 358).

Ethyl ether EtA' (229° cor) SG $\frac{1}{2}$ 1.086 Converted by heating with sodium into acetic ether, sodium phenyl acetate, a solid $\text{C}_{12}\text{H}_{11}\text{O}_2$, [175°] (? tri phenyl-phloroglucin), and an oil (? $\text{CH}_2\text{PhCOCHPhCO}_2\text{Et}$) (Hodgkinson, *C J* 37, 481, *C J Proc* 2, 189).

Amido ethyl ether $\text{CH}_3(\text{NH}_2)\text{CH}_2\text{A}'$ Formed from $\text{C}_6\text{H}_5\text{BrNHCOCH}_2\text{Ph}$ and hot water (Elfeldt, *B* 24, 3222)— $\text{B}^+\text{C}_6\text{H}_5\text{N}_2\text{O}$, [138°] *n* *Propyl ether* PrA' (238° cor) SG $\frac{1}{2}$ 1.0142 Made by heating alcoholic potassium phenyl acetate with PrI for two days. Sodium acts on it, forming propyl acetate, sodium phenyl acetate, a yellow oil $\text{C}_{12}\text{H}_{13}\text{O}_2$, (335° at 50 mm), and a small quantity of a solid, $\text{C}_{12}\text{H}_{13}\text{O}_2$, forming white needles (from petroleum), [170°], SG $\frac{1}{2}$ 1.039 This solid is also formed by the action of sodium on the oil $\text{C}_{12}\text{H}_{13}\text{O}_2$, as well as on the corresponding oil $\text{C}_{12}\text{H}_{13}\text{O}$, formed from ethyl phenyl acetate. The solid forms an acetyl derivative [100°] (Hodgkinson, *C J* 37, 483).

Iso-butyl ether $(\text{CH}_3)_2\text{CHCH}_2\text{A}'$ (147° cor) Sodium gives isobutyl acetate and A oil (Hodgkinson, *C J* 37, 485).

Benzyl ether $\text{PhCH}_2\text{A}'$ (318°) (S), (270° at 160 mm) SG $\frac{1}{2}$ 1.094 Got by heating benzyl chloride with alcoholic potassium phenylacetate for a week (H). Heated with sodium it forms sodium phenyl acetate, the salt of an acid $\text{C}_{12}\text{H}_{11}\text{O}_2$, [120°], possibly α - β -di-phenyl-propionic acid, and an oil, $\text{C}_{12}\text{H}_{13}\text{O}_2$, (820° at 60 mm), possibly the benzyl ether of that acid.

Chloride CH_2PhCOCl (108° at 17 mm.). SG $\frac{1}{2}$ 1.1692 (Anschütz a Berns, *B* 20, 1889; Hinsberg, *B* 28, 2952).

Anhydride $(\text{CH}_3\text{Ph CO})_2\text{O}$ [72 5°] Formed from the chloride and the Ag salt (A a B)

Amide $\text{CH}_3\text{Ph CO NH}_2$ [168°] (283°) Formed from the nitrile by heating with alcoholic KSH (Weddige, *J pr* [2] 7, 99) Formed also by heating phenylacetamide hydrochloride with oxalacetic ether and dilute (10 p c) NaOH aq (Pinner, *B* 22, 1627), and by the action of yellow ammonium sulphide on acetophenone (Willgerodt, *B* 21, 534) Plates, v sol hot water Forms with HgO a compound crystallising in needles [208°] (Reimer, *B* 13, 741) K_2CrO_4 and HOAc give $\text{CH}_3\text{Ph CO NHBr}$ [12°], crystallising from benzene in needles, and yielding benzylamine on warming with potash (Hoogewerf a Van Dorp, *R T C* 6, 385) Aldehyde and a few drops conc HCl aq give $(\text{C}_6\text{H}_5\text{CH}_2\text{CO NH})_2\text{CHMe}$ [228°] Chloral gives an analogous body

Diethylamide $\text{CH}_3\text{Ph CO NEt}_2$ [86°] (296° uncor) Formed from the chloride and diethylamine in ether (Hausknecht, *B* 22, 324)

Bromo-ethyl amide
 $\text{CH}_3\text{Ph CO NH C}_2\text{H}_4\text{Br}$ [85°] Plates

β Bromo propyl amide
 $\text{CH}_3\text{Ph CO NH CH}_2\text{CH}_2\text{CH}_2\text{Br}$ [46°] Formed from the chloride and $\text{C}_2\text{H}_5\text{Br}(\text{NH}_4)$ Needles

γ Bromo propyl-amine
 $\text{CH}_3\text{Ph CO NH CH}_2\text{CH}_2\text{CH}_2\text{Br}$ [44°] Needles, converted by KOH into $\text{CH}_2\text{CH}_2\text{CH}_2\text{O} \rightarrow \text{C CH}_2\text{Ph}$ (Eilfeldt, *B* 24, 3224)

Amide $\text{CH}_3\text{Ph CO NHPh}$ [117°] (Hofmann, *B* 13, 1225) Got, by intramolecular change, by the action of PCl_5 on the oxim of deoxybenzoin (Günther, *A* 252, 71)

Di-phenyl amide $\text{CH}_3\text{Ph CO NPh}_2$ [72°] Satiny needles (from ether) (Hausknecht)

Phenyl hydrazide $\text{C}_6\text{H}_5\text{N}_2\text{O}$ [167°] White flakes (Bulow, *A* 236, 196)

Nitrile $\text{C}_6\text{H}_5\text{CH}_2\text{CN}$ **Benzyl cyanide** (233° cor) $\text{S G } 131015 \text{ H C } 1,023,000 \text{ H F } -27,900$ (Berthelot a Petit, *C R* 108, 1219) The chief constituent of the oils of garden cress (*Lepidium sativum*) and of nasturtium (*Tropaeolum majus*) (Hofmann, *B* 7, 1293) Formed by boiling benzyl chloride with alcoholic KCy (Cannizzaro, *A Ch* [3] 45, 468) **Preparation**—By digesting benzyl chloride (1 kilo), crude KCy (1 kilo), and dilute alcohol for a week Tribenzylamine remains in the retort after distilling off the benzyl cyanide below 300° **Reactions**—1 A specimen (200°–230°) treated with ZnEt_2 , the temperature being kept below 130°, formed a product which, when decomposed by alcohol, gave cyanobenzene ($\text{C}_6\text{H}_5\text{N}$), and benzoin, $\text{C}_{14}\text{H}_{11}\text{N}_2\text{O}$, an indifferent body [160°]—2 Reduced by zinc and HCl aq to $\text{C}_6\text{H}_5\text{CH}_2\text{CH}_2\text{NH}_2$ (Berthsen, *B* 8, 691)—3 Alcoholic ammonium sulphide forms $\text{C}_6\text{H}_5\text{CH}_2\text{CS NH}_2$ [98°]—4 Water at 250° forms the amide, ammonium acetate acts in like manner (Berthsen, *B* 9, 429)—5 Bromine forms $\text{C}_6\text{H}_5\text{CHBr CN}$ and very sparingly soluble $\text{C}_6\text{H}_5\text{CHBr CBr}(\text{NH})$ [c 200°], which is crystalline, is decomposed by hot water and alcohol, and yields mandelic acid when heated with dilute HCl at 150° (Reimer, *B* 14, 1797)—6 Pyruvic acid and H_2SO_4 in the cold form $\text{C}_6\text{H}_5\text{N}_2\text{O}$ [145°] (Böttger, *B* 14, 1600).

Paranitrile $(\text{C}_6\text{H}_5\text{N})_n$, $n=8?$ **Cyanobenzene**, [171°] (F a T), [221°] (P.) Obtained by ex-

tracting with alcohol the product of the action of ZnEt_2 on the isomeric nitrile (Frankland a. Tompkins, *C J* 37, 568) Occurs in small quantity in the product of the action of alcoholic KCy on benzyl chloride (Pinner, *B* 17, 2010) Silky needles Hardly sol alcohol, sol. CS_2 , and glacial acetic acid Forms a gummy mass with warm dilute HCl, possibly a salt

Di-phenyl-acetic acid $\text{C}_{14}\text{H}_{12}\text{O}_2$, etc
CHPh₂CO₂H **Deoxybenzic acid** Mol w 212 [148°] Formed by reducing benzilic acid with HIAq (Jena, *A* 155, 84) and by heating $\text{CHPhBr CO}_2\text{H}$ (1 mol.) with benzene (1 mol.) and zinc dust (Symons a Zincke, *B* 6, 1188, *A* 171, 122) Obtained also by saponifying its nitrile or amide Needles (from water), sl sol cold water, v sol alcohol and ether Yields benzophenone on oxidation, and di phenyl methane on distillation with soda lime Bromine forms $\text{CPh}_2\text{Br CO}_2\text{H}$

Salts— $\text{BaA}'_2\text{2aq}$ Efflorescent monoclinic crystals— $\text{CaA}'_2\text{2aq}$ — AgA' amorphous pp

Methyl ether MeA' [60°] Plates (from alcohol) (Rattnier, *B* 21, 1316)

Ethylether EtA' [58°] Prisms

Amide $\text{CHPh}_2\text{CONH}_2$ [166°] Got by action of KOH (6 g) in alcohol on the nitrile (2 g) (Anschutz a Romig, *A* 233, 347), and by heating the ammonium salt under pressure at 230° (Neure, *A* 250, 141) Plates (from alcohol)

Nitrile CHPh_2CN [76°] (N), [73°] (F a S) (183° at 12 mm) Formed by heating CHPh_2Br with HgCy_2 for 18 hours at 165°, and extracting with benzene (Friedel a Balsahn, *Bl* [2] 33, 589) and by heating the acid with lead sulphocyanide at 170° (Freund a Immerwahr, *B* 23, 2845) Got also, together with a polymeride [168°], by nitrating diphenyl ethane and reducing the resulting $\text{C}_{14}\text{H}_{10}\text{N}_2\text{O}$ [148°] with SnCl_4 and alcoholic HCl (A a R), and by treating the amide with PCl_5 in POCl₃ (N) White needles (from ligroin and ether) Alcoholic NaOEt and benzyl chloride form $\text{CPh}_2(\text{CH}_2\text{Ph})\text{CN}$ [126°] An ethereal solution of iodine (1 mol) gradually added to an alcoholic solution of the nitrile (2 mols) and NaOEt (1 mol) forms $\text{CN CPh}_2\text{CPh}_2\text{CN}$ [c 230°]

Tri-phenyl acetic acid $\text{CPh}_3\text{CO}_2\text{H}$ [264°] Formed by the action of fuming HCl aq and HOAc on the nitrile at 200° (E a O Fischer, *B* 11, 1598, *A* 194, 242) Prepared by heating CCl_4 , CO_2H (250 g), benzene (340 g), and AlCl_3 (250 g), mixing the product with water and distilling with steam, the residue of aluminum triphenylacetate is decomposed by NH_4Aq and the filtrate ppd by HCl, the yield being 5 p c (Elbs a Tölle, *J pr* [3] 32, 624) In this preparation too by products $(\text{C}_6\text{H}_5\text{O})_n$ [825°] and $(\text{C}_6\text{H}_5\text{O})_n$ [75°] crystallising in yellow needles are also formed Monoclinic prisms (from alcohol), sl sol HOAc Very feeble acid The NH_4 salt gives off NH_3 spontaneously The K salt, ppd as needles by conc KOH aq, is decomposed by water Fuming H_2SO_4 at 100° forms $\text{SO}_3\text{H C}_6\text{H}_5\text{CPh}_2\text{CO}_2\text{H}$, which yields $\text{BaA}'_2\text{aq}$, an amorphous salt—Salt AgA' Powder, not affected by light

Nitrile CPh_3CN [127 5°] Formed from HgCy_2 and CPh_3Cl at 160° and from KCy and CPh_3Br (Elbs, *B* 17, 700) Monoclinic prisms

(from ligroin or HOAc) Converted by alcoholic potash into a polymeride [210°] crystallising in colourless needles

References—AMIDO-, BROMO-, BROMO AMIDO, BROMO NITRO-, CHLORO, IODO-, and OXY PHENYL-ACETIC ACID

PHENYL-ACETIC ALDEHYDE C_6H_5O & $C_6H_5CH_2CHO$. (206°) (Radziszewsky, *B* 9, 872), (194°) (Etard)

Formation—1 By distilling calcium phenylacetate with calcium formate (Cannizzaro, *A* 119, 254)—2 From a bromo- (or chloro) β oxyphenyl-propionic acid $CHPh(OH)CHBrCO_2H$ by distilling with dilute Na_2CO_3 , the yield being 75 p.c. (Lupp *B* 16, 1286, Erlenmeyer, *A* 219, 179)—3 By boiling $PhCH(OH)CH_2OH$ (1 pt) for an hour with H_2SO_4 (3 pts) and water (12 pts) (Zincke, *A* 216, 301) Stronger H_2SO_4 yields $C_6H_5H_{12}$ [101°]—4 From ethyl benzene by successive treatment with CrO_2Cl_2 and water (Etard, *A* *Ch* [5] 22, 248)—5 By distilling phenyl glycidic acid with dilute H_2SO_4 (Erlenmeyer, *B* 13, 304).

Properties—Oil. Forms a crystalline compound with $NaHSO_4$.

Reactions—1 With acetoacetic ether and NH_3 , it yields benzyl-di-methyl pyridine di-carboxylic ether dihydride $C_6H_5N(C_2H_5)_2Me_2(CO_2Et)_2$ [1 8 5 2 6] [115°] (Jeaurenaud, *B* 21, 1784)—2 HCl forms $CH_2PhCH(OH)CN$ [58°], *S* 1 at 15°, crystallising in stellate groups of colourless needles, *v* sol alcohol.—3 HNO_3 (*S* G 1 49) at -12° gave *o* and *p*-nitro benzoic aldehyde

Phenyl-hydrasid CH_2PhCHN_2HPh [58°] Prisms, *v* sol alcohol. Converted into phenyl indole by heating with $ZnCl_2$ at 180° (Fischer a Schmitt, *B* 21, 1072)

ψ **Ethyl derivative** $C_6H_5CHCH_2OEt$ (217°) *S* G 3 981 Formed from ω chlorostyrene and alcoholic potash (Erlenmeyer, *B* 14, 1868) Oil, easily decomposed by heating with water into alcohol and the aldehyde

Di-phenyl-acetic aldehyde $CHPh_2CHO$ (315°) Formed by heating hydrobenzoin (1 pt) or isohydrobenzoin with (6 pts of) dilute (20 p.c) H_2SO_4 for 8 hours at 210° (Zincke a Breuer, *B* 9, 1769, *A* 198, 182, Weise, *A* 248, 38) Oil, *v* sol alcohol and ether. Yields benzophenone and CO_2 on oxidation. Forms a crystalline compound with $KHSO_4$. Alcoholic potash yields di-phenyl-carbinol and di-phenyl methane. Nascent HCl yields $CHPh_2CH(OH)CN$, upon which alcoholic hydrogen chloride reacts with formation of $CHPh_2CH(OH)C(NH_2)ClOEt$ [185°]. The aldehyde appears to form condensation products [168°] and [214°] on standing

Phenyl hydrazide $CHPh_2CHN_2HPh$ Flat stellate needles (from hot alcohol), *v* sol ether (Rudolph, *A* 248, 101)

Oxam $CHPh_2CHNOH$ [120°] White needles, formed together with another body [145°] by the action of hydroxylamine in dilute alcohol (Auwers, *B* 24, 1780)

PHENYL-ACETIC CARBOXYLIC ACID *v*. CARBOXY-PHENYL-ACETIC ACID

PHENYL-ACET-IMIDO-ACETATE

$C_6H_5CH_2C(NH)OAc$ [129°] White needles *v* sol alcohol, al sol cold water. Formed by boiling phenyl acet-imido ethyl ether with Ac_2O (Luckenbach, *B* 17, 1493)

PHENYL-ACETIMIDAMIDE *v* **PHENYL-**

ACETAMIDINE

PHENYL-ACET-IMIDO-ETHYL ETHER

$C_6H_5CH_2C(NH)OEt$ The hydrochloride B^+HCl [c 85°] is formed by passing dry HCl into a solution of phenyl-acetomitrile in absolute alcohol (Luckenbach, *B* 17, 1421) The free base is a colourless liquid of aromatic odour, decomposed on distillation into alcohol and phenyl acetomitrile. The hydrochloride is resolved on fusion into phenyl acetamide and $EtCl$

PHENYL-ACETONITRILE *v* **Nitrile** of **PHENYL ACETIC ACID**

PHENYL ACETONYL SULPHIDE C_6H_5SO & $PhSCH_2COCH_3$ [35°] (144° at 15 mm) Formed from chloro acetone and $NaSPh$ (Delisle, *B* 22, 308) Large tables (from ether), turned dark violet by hot H_2SO_4 . Yields a phenyl-hydrazide [83°]

The corresponding sulphone $PhSO_2CH_2Ac$ [58°] is formed from chloro acetone and sodium benzene sulphinate (R. Otto, *B* 19, 1642)

PHENYL ACETO-PHENONE

[1 8] $C_6H_5C_6H_5COCH_3$ [121°] (*c* 326°) Formed from diphenyl, $AcCl$, and $AlCl_3$ (Adam, *A* *Ch* [6] 15, 255) Prisms, reduced by sodium amalgam to the carbinol [86°]

DI PHENYL ACETOXIM (so called) *v* **Oxim** of **BENZOPHENONE**

PHENYL-ACETURIC ACID

$CH_2PhCOONHCH_2CO_2H$ [142°] *S* 73 at 12° Occurs in horses' urine, and is also found in urine after a dose of phenyl acetic acid (Sal kowski, *B* 17, 3010, *H* 7, 162) Formed by the action of the chloride or anhydride of phenyl acetic acid on glycocholi (Hotter, *B* 20, 81, *J* pr [2] 38, 98) Small trimetric crystals (from alcohol), *a* *b* *c* = 890 12374 *V* sl sol ether, *v* sol hot water and alcohol— CaA_2 , 2aq *S* 3 16 at 11 2°— CuA_2 , aq— PbA_2 , aq prisms— ZnA_2 , plates, *v* sol hot water— AgA , amorphous

Methyl ether MeA [86°] Needles

Ethyl ether EtA [79°] Prisms

Propyl ether PrA [31°] Plates

Amide $CH_2PhCOONHCH_2CONH_2$ [174°] Pearly hexagonal plates, *v* sol hot water. With HgO it yields $Hg(C_6H_5N_2O_2)_2$

PHENYL-ACETYL ACETONE $C_6H_5H_3O_2$ & $C_6H_5CH_2COCH_2COCH_3$ *Benzyl methyl methylene di-ketone* (266°–269° at 748 mm) Has weak acid properties

Formation—Phenyl acetyl acetoacetic ether, formed by the action of phenyl acetyl chloride upon sodio aceto-acetic ether, is boiled with water for 6 hours

Properties—Colourless oil. Sol alcohol, ether, benzene, hot water, strong acids, and dilute alkalis, al sol cold water. With phenyl-hydrazine it condenses to phenyl benzyl methyl pyrazole

Salts— A^+Ag white pp— A^+Na^+ crystalline solid (Fischer a Bulow, *B* 18, 2136)

PHENYL-ACETYLENE C_6H_5 & $C_6H_5C\equiv CH$ *Acetylenic benzene* (141 6° cor) *S* G 3 9295 μ = 1 542 (Brühl, *A* 285, 13) *S* V 125 8

Formation—1 By distilling phenyl propiolic acid with dry $Ba(OH)_2$ (Weger, *A* 241, 70)—2 By boiling styrene dibromide with alcoholic potash, and heating the resulting bromo styrene with alcoholic potash (Glaser, *Z* [2] 5, 97, *A* 154, 155, Hollemann, *B* 20, 3080).—3 From

acetophenone by treatment with PCl_5 , the resulting $\text{C}_6\text{H}_5\text{COCl}$, being heated with conc alcoholic potash at 120° (Friedel, *Z* [2] 5, 123)
4 By the passage through a red hot tube of a mixture of styrene and hydrogen, or of benzene and ethylene (Berthelot, *C R* 87, 952)

Properties—Liquid with peculiar odour. Its alcoholic solution forms a yellow pp $\text{Cu}_2(\text{C}_6\text{H}_5)_2\text{O}$ with ammoniacal cuprous chloride and a white pp $\text{Ag}_2(\text{C}_6\text{H}_5)_2\text{O}$ with ammoniacal AgNO_3 . These pps are explosive. Sodium added to its ethereal solution ppts CPh CNa as a white powder, which takes fire in the air, and is reconverted by water into phenyl-acetylene.

Reactions—1 Unites with bromine—2 Sodium and CO_2 give sodium phenyl propiolate, (Paterno, *G* 2, 553)—3 Sodium and EtI give PhCCEt (202°) (Morgan, *C J* 29, 162)—4 H_2SO_4 , followed by water, yields acetophenone (Friedel a Balsoun, *BI* [2] 35, 54)—5 Boiling HOAc and zinc dust reduce it to styrene (Aronstein a Hollemann, *B* 22, 1181)

Di phenyl acetylene C_{10}H_8 , i.e. CPh CPh. **Tolane** [71°] (Béhal) Formed by boiling CPhHBr or the corresponding di chloro di phenyl ethane, with alcoholic potash (Lamprecht a Schwanert, *A* 145, 347, Fittig, *A* 168, 74) Formed also by the action of sodium amalgam on an alcoholic solution of CPhCl_2 , CPhCl , (Liebermann a Homeyer, *B* 12, 1974) Large crystals (from ether) May be distilled. Yields di phenyl ethylene (stilbene) when heated with HIAq and P at 150° (Barber, *J* 1876, 366) Yields benzoic acid on oxidation. Conc H_2SO_4 gives a green colour, and on warming to 60° , diluting with water and distilling with steam, phenyl benzyl ketone (deoxybenzoin) [55°] is produced (Béhal, *BI* [2] 49, 337)

Chlorides—1 Di chloro di phenyl ethylene and tetra chloro di phenyl-ethane

Dibromide v Di bromo di phenyl ethylene

Di phenyl di acetylene C_{10}H_8 , i.e. CPh C C CPh [97°] (G), [88°] (H) Formed by shaking the cuprous compound of phenyl-acetylene with alcoholic NH_3 and air, or better, with alkaline K_2FeCy_4 (Glaser, *A* 154, 159, Baeyer a Landsberg, *B* 15, 57) Long needles (from dilute alcohol), v sol ether. Hot H_2SO_4 carbonises it. In ethereal solution Br forms $\text{C}_{10}\text{H}_8\text{Br}_2$ [178°] and $\text{C}_{12}\text{H}_{10}\text{Br}_2$ [149° – 153°] (Holleman, *B* 20, 3081)

Picric acid compound $\text{C}_{11}\text{H}_9\text{O}_7\text{N}_3$, [108°] Yellow crystals (from alcohol)

References—Amido, Iodo, and Nitro-phenyl-acetylene

TRI PHENYL ACETYLENE-TRIAMINE

$\text{C}_{18}\text{H}_{12}\text{N}_6$, i.e. $\text{N}_3\text{H}(\text{C}_6\text{H}_5)_3$, [190°] Formed from acetylene tetrabromide, aniline, and alcoholic potash (Sabanejeff, *A* 178, 125) Needles, v sl sol cold alcohol—Salts $\text{B}'_2\text{H}_2\text{PtCl}_4$ — $\text{B}'_2\text{H}_2\text{HgCl}_4$ amorphous pp

DI-PHENYL ACETYLENE-DI-THIO-DI-

UREA $\text{CS} \begin{smallmatrix} \text{NH CPh NH} \\ \text{NH CPh NH} \end{smallmatrix} \text{CS}$ Formed from benzil and thio urea at 145° (Anschütz a Geldermann, *A* 261, 134) Crystals, decomposing at 300° , v sl sol alcohol

DI-PHENYL-ACETYLENE-DI-UREA

$\text{CO} \begin{smallmatrix} \text{NH CPh NH} \\ \text{NH CPh NH} \end{smallmatrix} \text{CO}$ Tolane-urea Formed

by heating benzil with urea at 175° (Anschütz a Geldermann, *A* 261, 133, Angeli, *B* 24, 606, *G* 19, 568) Crystals, not decomposed at 315° , v sl sol alcohol and ether. Reduces ammoniacal AgNO_3 .

Di acetyl derivative $\text{C}_{18}\text{H}_{12}\text{Ac}_2\text{N}_6\text{O}_2$ [266°]

PHENYL-ACETYL-SUCCINIC ACID v ACETYL-PHENYL SUCCINIC ACID The mono ethyl ether of this acid yields on treatment with alcoholic NH_3 , a compound $\text{C}_{12}\text{H}_{12}\text{N}_2\text{O}_2$ [264°], which forms, on saponification, a compound $\text{C}_{12}\text{H}_{11}\text{NO}_2$ [148°], crystallising in small needles (Weltner, *B* 18, 793)

PHENYL ACRIDINE $\text{C}_{15}\text{H}_{13}\text{N}$ i.e.

$\text{C}_6\text{H}_5 \begin{smallmatrix} \text{CPh} \\ \text{N} \end{smallmatrix} \text{C}_6\text{H}_5$, [181°] (c 404°) VD 129.7 [$\text{H} = 1$]

Formation—(Bernthsen, *B* 15, 3011, 16, 1809, 19, 425, 20, 1552, *A* 192, 19, 224, 12) 1 By heating benzoyl diphenylamine (30 g) with ZnCl_2 (30 g) at 210° – 280° —2 From benzotrichloride, diphenylamine, and ZnCl_2 —3 By heating benzonitrile with diphenylamine hydrochloride at 240° —4 By heating di phenyl benzamidine hydrochloride to 250° —5 In small quantity by heating di phenyl amine with cinnamic acid and ZnCl_2 —6 By diazotising chrysaniline (di amido phenyl acridine) and boiling the product with alcohol (Fischer a Körner, *B* 17, 206, *A* 226, 184)

Preparation—By heating benzoic acid (50 g) diphenylamine (70 g) and ZnCl_2 (150 g) at 260° for 10 hours. The yield of crude base (50 g) is good

Properties—Colourless thin prisms (from alcohol), thick yellowish prisms (containing C_6H_5) or tablets (from benzene) V sol benzene m sol ether, sl sol alcohol, insol water. Solutions of its salts exhibit green fluorescence. Not affected by boiling dilute HNO_3 , by fuming HCl at 260° , by AcCl , by AcO at 190° , by Bz_2O at 160° , by potash fusion, by heating with H_2SO_4 (3 vols) diluted with water (1 vol), or by distillation with soda lime

Reactions—1 CrO_3 in HOAc gives benzoic acid and CO_2 —2 On oxidation with KMnO_4 it gives phenyl quinoline carboxylic acid. On the other hand, its alkyl halogen addition products give rise to phenyl amido benzoic acid $\text{C}_{16}\text{H}_{11}(\text{NHPH})\text{CO}_2\text{H}$ on oxidation with KMnO_4 (Claus a Nicolaysen, *B* 18, 2706)—3 Yields on nitration with H_2SO_4 and fuming HNO_3 di nitro phenyl-acridine $\text{C}_{15}\text{H}_{11}(\text{NO}_2)_2\text{N}$, which is sol aqueous or alcoholic HCl but sl sol alcohol, and tri nitro phenyl acridine, which crystallises from toluene and alcohol in minute yellow needles, hardly sol alcoholic HCl , and converted by hot SnCl_4 into a red dye (? tri-amido-phenyl acridine), which becomes colourless on further reduction

Salts—Partially decomposed by water— $\text{B}'\text{HCl}$ orange needles or prisms, sl sol cold water, sol alcohol. Its solution exhibits dark-green fluorescence— $\text{B}'\text{HCl}$ 3aq red crystals— $\text{B}'_2\text{H}_2\text{PtCl}_4$ —Nitrate [188°] Yellow needles, sl sol HNO_3 —Sulphate yellowish-red trimetric crystals, v sl hot water—Picrate Hair-like crystals (from alcohol) Phenyl-acridine does not appear to form an acetate

Methylo-chloride $B'MeCl$ soluble needles— $B'Me.PtCl$, decomposes below 70°

Methylo-nitrate $B'MeNO_3$. Sparingly sol yellow needles

Methylo-iodide $B'MeI$. Black monoclinic prisms (from alcohol), insol ether. Converted by $KOH.Aq$, by $NH_3.Aq$, or by moist Ag_2O , into $B'MeOH$ [108°], which is also got by oxidising $C_6H_5 \begin{smallmatrix} CHPh \\ \diagup \\ NMe \end{smallmatrix} > C_6H_5$, a body to which it is reduced by tin and $HCl.Aq$

Hydrate $C_6H_5 \begin{smallmatrix} CHPh \\ \diagup \\ NH \end{smallmatrix} > C_6H_5$, [164°]

Formed by reducing phenyl-acridine hydrochloride in aqueous solution with zinc-dust or by adding sodium amalgam to an alcoholic solution. Colourless needles, with no basic properties, sol alcohol and ether. Alcoholic $AgNO_3$ oxidises it to phenyl-acridine, a change which is slowly effected even by air. It yields an acetyl derivative $C_{10}H_7.N(Ac)N$ [128°], and is converted by MeI at 140° into $C_{10}H_7.MeN$ [104°], whence HCl and $NaNO_2$ produce the methylo-hydroxide of phenyl-acridine (*v supra*)

Di-sulphonic acid. Got by heating phenyl-acridine (10 g), H_2SO_4 (15 g), and $H_2S_2O_8$ (15 g) for 12 hours at 150° . Its dilute solutions exhibit green fluorescence. The salt $C_{10}H_7.N(SO_3Na)_2$ is insol absolute alcohol, and in very dilute solutions exhibits blue fluorescence. The Ba and mercurous salts are nearly insol. Aq.

References — AMIDO- and OXY-PHENYL-

ACRIDINE

PHENYL-ACRIDINE CARBOXYLIC ACID

$C_6H_5 \begin{smallmatrix} C(C_6H_5, CO_2H) \\ \diagup \\ N \end{smallmatrix} > C_6H_5$, *Acridyl-benzonic acid*

Formed by heating phthalic anhydride (30 g) with diphenylamine (45 g) and $ZnCl_2$ (75 g) for 12 hours at 190° (Bernthsen & Traube, *B* 17, 1510, *A* 224, 45). Yellow crystalline powder, sol acids and alkalis, insol water, sl sol boiling alcohol. Its solutions in dilute acids have a green fluorescence, in alkalis a blue fluorescence. Split up by heat into CO_2 and phenyl-acridine. Yields a hydride [160° – 165°]

Salts— NaA . aq. colourless pearly plates or long needles— $HA.HCl$ [163°]. Yellow crystals (from hot dilute HCl), sl sol hot water

An isomeric acid $C_6H_5 \begin{smallmatrix} CPh \\ \diagup \\ N \end{smallmatrix} > C_6H_5, CO_2H$, [252° – 255°], formed by oxidation of phenyl-methyl-acridine with chromic acid mixture, is sl sol alcohol, and gives sparingly soluble BaA , (Bonna, *A* 239, 62)

PHENYL-ACRYLIC ACID *v* CINNAMIC ACID and ATROPIC ACID

Di-phenyl-acrylic acid *v* PHENYL-CINNAMIC ACID

PHENYL-ALANINE *v* α -AMIDO-PHENYL-PROPIONIC ACID and PHENYL-AMIDO-PROPIONIC ACID

PHENYL ALDEHYDE *v* PHENYL-ACETIC ALDEHYDE

PHENYL ALDEHYDO-BENZYL KETONE $Ph.CO.CHPh.CHO$ [110°]. Formed from de-oxybenzoin, formic ether, and $NaOEt$ in ether (Claisen & Meyerowitz, *B* 22, 3279). Yellowish-white crystals. $FeCl_3$ gives a dark violet colour

PHENYL ALDEHYDO-ETHYL KETONE $Ph.CO.CHMe.CHO$ [119°]. Formed from

phenyl ethyl ketone, formic ether, and $NaOEt$ in Et_2O (Claisen & Meyerowitz, *B* 22, 3276). White needles, v sol alcohol and water. $FeCl_3$ colours its alcoholic solution dark violet. Gives an anilide [132°] crystallising in needles

PHENYL ALDEHYDO-PROPYL KETONOL $Ph.CO.CHEt.CHO$ [87°]. Formed from phenyl *n*-propyl ketone, formic ether, and $NaOEt$ in ether (Claisen & Meyerowitz, *B* 22, 3278). White plates. $FeCl_3$ gives a dark-violet colour. The anilide [120°] crystallises in white needles

TETRA-PHENYL-ALDINE *v* TETRA-PHENYL-PYRAZINE

PHENYL-ALLENYL-AMIDO-AMINE *v* α -AMIDO-AMINOXIM

PHENYL-ALLOPHANIC ACID *Ethyl ether* $C_6H_5.NH.CO.NH.CO$ [120°]. Formed from phenyl-urea and chloro-glyoxylic ether $COCl.CO.Et$ (Stojentini, *J pr* [2] 32, 18). Needles, v sol alkalis and conc $HCl.Aq$

Phenyl ether $NHPh.CO.NH.CO.Ph$. Formed from phenol and cyanic acid vapour (Tuttle, *J* 1857, 451). Crystals (from alcohol), insol cold water. Decomposes at 150° into phenol and cyanuric acid

Benzyl ether $NHPh.CO.NH.CO.C_6H_5$, [158°]. Formed by heating phenylcyanate with benzyl carbonate at 150° under pressure (Traube, *B* 22, 1574). V sl sol water, m sol alcohol

Di-phenyl-allophanic acid

$NHPh.CO.NPh.CO.H$

Methyl ether MeA [231°]

Ethyl ether EtA [188°]. Formed from di-phenyl diisocyanate $Ph.N=C=O$, and alcohol (Hofmann, *B* 4, 246), and by the action of HgO and alcoholic NH_3 on $NHPh.CS.NPh.CO.Et$ (Seidel, *J pr* [2] 32, 266). Needles, sl sol ether. Yields $(C_6H_5.N_2O_3).3HgO$ [129°], crystallising in prisms (from alcohol)

Isoamyl ether $C_6H_5.A$ [58°]

Reference — DI-BROMO DI-PHENYL ALLOPHANIC ACID

PHENYL ALLYL-CYANAMIDE KETONE

$C_6H_5.CO.CHPh.C_6H_5$, (337°). Formed from de-oxybenzoin, $NaOEt$, and allyl iodide (Buddeberg, *B* 23, 2067). Oil, v sol alcohol

PHENYL ALLYL-CYANAMIDE $C_6H_5.N_2.C_6H_5.N.CO.NH.C_6H_5$, *Carboallylphenylamide* [105°]. Formed from phenyl allyl thio-urea and $Pb(OH)_2$ (Bizio, *J* 1861, 497). Silky needles (from dilute alcohol)— $B.HgCl_2$, amorphous— $B.H.PtCl_5$

PHENYL-ALLYLENE $C_6H_5.C=CH_2$, (185°). Formed by the action of alcoholic potash on the bromo-propenyl-benzene, which is got by boiling $CHPhBr.CMe.CO.H$ with water (Körner, *B* 21, 276). Yellowish liquid, with unpleasant odour. Yields a liquid dibromide (c 258°) and a solid tetrabromide [75°]. Aqueous $HgCl_2$ forms white amorphous $(C_6H_5)_2Hg_2Cl_2O_2$, which on heating with HCl yields phenyl ethyl ketone

v PHENYL-ALLYL-HYDRAZINE

$C_6H_5.NPh.NH_2$, (198° at 184 mm). Formed by reducing the nitrosamine derived from allyl-aniline (Michaelis & Claessen, *B* 22, 2239). Got also from allyl-bromide and phenyl hydrazine. Oil, v sol dilute HCl . Reduces Fehling's solution on warming. Oxidised by aqueous $FeCl_3$ to the tetrazene $C_6H_5.NPh.N.NPh.C_6H_5$, (86°). Benzoic aldehyde yields $C_6H_5.NPh.N.C_6H_5$ [52°].— $B.HCl$ [187°]. Silky needles

Benzoyl derivative $C_6H_5.AcN$, [139°]

s-Phenyl allyl-hydrazine $C_9H_{11}N_2$ \pm e
Ph NH NH C_3H_5 (172° at 60 mm) Got from phenyl-hydrazine and allyl bromide (Fischer a. Knoevenagel, A 239, 203) Oil, sol dilute acids Reduces cold Fehling's solution Oxidises by HgO to benzene-azo propylene $Ph.N_2.C_3H_5$, an oil, (95°–100° at 27 mm)

DI-PHENYL-DI-ALLYLIDENE-DIAMINE
 $C_{18}H_{19}N_2$ \pm e $N_2Ph_2(C_3H_5)_2$ Formed from aniline and acrolein below 0° (Schiff, A Suppl 3, 358) Yellow resin — $B'H_2PtCl_6$

DI-PHENYL-DI-ALLYLIDENE-ETHYLENE DIAMINE $C_{16}H_{17}N_2$ (N CH CH CH C_2H_5)₂ [110°] Formed by mixing ethylene-diamine (1 mol) with cinnamic aldehyde (2 mols) (Mason, B 20, 271) Large colourless tables (from ether) V sol alcohol and benzene, m sol ether Decomposed into its constituents by acids

PHENYL ALLYL OXIDE C_8H_7O C_6H_5 (194°) Formed from allyl bromide and NaOPh (Henry, B 5, 455) Liquid

PHENYL ALLYL SULPHIDE C_8H_7S C_6H_5 (c 208°) Formed by distilling the acid $CH_3C(SPh)CH_2CO_2H$ (Escales a. Baumann, B 19, 1792) Liquid, forming a blue solution in H_2SO_4

PHENYL-ALLYL-THIOSEMICARBAZIDE
 $C_9H_9NH NH CS NHC_6H_5$ [119°] Formed from phenyl hydrazine and allyl thiocarbimide dissolved in benzene (A E Dixon, C J 57, 263, Avenarius, B 24, 268) Silvery needles, insol water, v sol alcohol and ether Gives a mahogany colour with $FeCl_3$

Di phenyl-allyl thiosemicarbazide
 $C_{12}H_{13}N(C_6H_5)_2NH CS NPh$ [103°] Formed from phenyl allyl-hydrazine and phenyl thio carbimide (Michaelis a. Claessen, B 22, 2237) Needles (from hot alcohol)

PHENYL ALLYL THIO UREA $C_{10}H_{11}N_2S$ \pm e
NPh CS NHC_6H_5 *Phenyl thiosumamine* [98°] S (alcohol) 70 at 16° Formed by the action of aniline on oil of mustard (Zinn, A 84, 348) and of allylamine on phenyl thiocarbimide (Weith, B 8, 1529) Monoclinic crystals, insol water Cyanogen passed into its alcoholic solution forms $C_{12}H_{12}N_2S$, which crystallises from alcohol in stellate forms, and is converted by dilute H_2SO_4 into $CS \begin{smallmatrix} N(C_6H_5) \\ N(C_6H_5) \end{smallmatrix} CO$ [161°], whence alcoholic silver nitrate forms $CO \begin{smallmatrix} N(C_6H_5) \\ N(C_6H_5) \end{smallmatrix} CO$ [108°], which is resolved by baryta-water into oxalic acid and phenyl allyl-urea (Maly, Z [2] 5, 258)

PHENYL ALLYL THIO-UREA CARB OXYLIC ACID v **ALLYL-THIO URAMIDO-BENZOIC ACID**

PHENYL-ALLYL UREA NPh CO NHC_6H_5 [97°] Formed as described under phenyl allyl-thio-urea Shining white needles

PHENYL-AMENYL-AMIDINE $C_{11}H_{11}N_2$ \pm e
Me, C(NH₂)(NPh) The oxalate $B'H_2C_2O_4$ [192°] s formed by heating the corresponding valerionitrile with aniline hydrochloride at 170° Freund a. Lenz, B 24, 2155)

PHENYL-AMIDO-ACETIC ACID $C_8H_7NO_2$ \pm s
 $Ph(NHPh)CO_2H$ *Phenyl glycocoll Amidido-acetic acid* [127°] Formed by the action of aniline on bromo acetic acid and on chloro-acetic acid (Michaelson a. Lippmann, Z 1866,

15, Schwebel, B 10, 2046) Prepared by boiling aniline (45 g) dissolved in a little ether with chloro-acetic acid (25 g) and water (1500 cc) for half an hour, and rapidly evaporating to crystallisation (Rebuffat, G 17, 233) By-products are $C_8H_7NO_4$ [99°] (Meyer, B 14, 1325) and $C_8H_7N_2O_2$ [218°], which is perhaps $CH_2(NHPh)CO NPh CH_2CO_2H$, though this formula may also be assigned to an isomeric acid [180°] formed by the action of alcoholic potash on di-oxy-di-phenyl-pyrazine (Hausdörfer, B 22, 1803, Abenius, J pr [2] 40, 432)

Properties—Small crystals, m sol water, insol ether By fusion with potash at 260° it yields a mass which forms indigo on oxidation (Heumann, B 23, 3043, cf Lederer, J pr [2] 42, 383, Biedermann a. Lepetit, B 23, 3289)

Salt— CaA_2 2aq Needles (from dilute alcohol), m sol hot water, insol alcohol On distillation with calcium formate it yields indole (Mauthner a. Suida, M 10, 251)

Methyl ether MeA' [48°] Formed by heating aniline with methyl chloro-acetate (Meyer, B 8, 1157) and by the action of aniline on methyl diazo acetate (Curtius, J pr [2] 38, 437) Rectangular plates (from ether) or needles (from alcohol)

Ethyl ether EtA' [59°] Formed in like manner Colourless plates (from ether)

Formyl derivative
NPh(CHO) CH_2CO_2H [124°] Formed by saponification of its ether (290°–295°), which is the product of the action of chloro-acetic ether on sodium formanilide (Paal a. Otten, B 23, 2593) Needles, v sol ether and hot water Yields $C_8H_7NO_2Na$, a hygroscopic mass

Acetyl derivative NPhAc CH_2CO_2H [191°] Formed from phenyl amido acetic acid, Ac₂O, and benzene (R) Formed also by heating aniline with chloro acetic acid and NaOAc at 110°–150° (Hausdörfer, B 22, 1797), and by the action of boiling alcoholic potash on its ether (300°), which is got from sodium acetanilide and chloro-acetic ether (P a O) Pearly plates (from water) or needles (from HOAc-hgroin), v sol alcohol and hot water Yields the salts NaA' , BaA' , 3aq, and CuA'_2

Chloro-acetyl derivative
 $CH_2Cl CO NPh CH_2CO_2H$ [133°] Formed from phenyl-amido acetic acid and chloro acetyl chloride (Abenius, J pr [2] 40, 429, 499) Plates or prisms, v sol alcohol On boiling with aqueous sodium carbonate it yields $CH_2(OH) CO NPh CH_2CO_2H$ [128°], which crystallises in plates, forms the salts CaA'_2 6aq and BaA'_2 , the anhydride $PhN \begin{smallmatrix} < CO CH_2 > \\ < CH_2 CO > \end{smallmatrix} O$ [169°], and the amide $CH_2(OH) CO NPh CH_2 CONH_2$ [129°]

Bromo-acetyl derivative Plates, decomposing at 153° (Hausdörfer, B 22, 1803)

Benzoyl derivative $CH_2(NPhBz) CO_2H$ [63°] Amorphous pp (Rebuffat)

Nitrosamine NPh(NO) CH_2CO_2H [105°] Yellow needles (Schwebel, B 11, 1132), v sol warm water

Anhydride C_8H_7NO [263°] Got by heating the acid at 150° (Meyer, B 10, 1967) Needles (from alcohol), insol water.

Amide $CH_2(NHPh) CONH_2$ [133°] Formed by heating chloro acetamide with aniline and

NaOAc at 180° (Bischoff, *B* 22, 1809, cf Meyer, *B* 8, 1154) Minute needles (from water)

Amide $\text{CH}_2(\text{NHPh})\text{CO NHPh}$ [112°]

Formed by boiling chloro-acetic ether or $\text{CH}_2\text{Cl CO NHPh}$ with aniline (Wilm a Wischin, *Z* 1868, 74, Meyer, *B* 8, 1156, Hausdörfer, *B* 22, 1796) Formed also by heating aniline with alcohol and the compound of glyoxal with NaHSO_3 (Hinsberg, *B* 21, 110) Needles (from alcohol), sl sol cold water

p-Toluide $\text{CH}_2(\text{NHPh})\text{CO NHC}_6\text{H}_4\text{Me}$ [165°] Formed by heating aniline with fused NaOAc and $\text{CH}_2\text{Cl CO NHC}_6\text{H}_4\text{Me}$ at 150° (Bischoff, *B* 23, 2000, cf Meyer, *B* 8, 1156)

Nitrile $\text{CH}_2(\text{NHPh})\text{CN}$ Formed from chloro acetonitrile and aniline in ether at 90° (Engler, *B* 6, 1004) Thick oil, forming a crystalline hydrochloride, decomposed by water

Reference — Bromo PHENYL AMIDO-ACETIC ACID

Isomeride — AMIDO PHENYL ACETIC ACID

Di-phenyl-di-amido-acetic acid $\text{C}_{16}\text{H}_{15}\text{N}_2\text{O}_2$, $\text{e CH}(\text{NHPh})_2\text{CO}_2\text{H}$ Formed as a yellow crystalline powder by adding aniline to an aqueous solution of glyoxylic acid (Böttinger, *B* 11, 1559) Decomposed by hot water

ω -PHENYL AMIDO ACETOPHENONE

$\text{C}_{16}\text{H}_{15}\text{ON}$ $\text{e C}_6\text{H}_5\text{CO CH}_2\text{NHPh}$ *Aceto phenone-anilide* [93°] Prepared by the action of aniline on ω bromo acetophenone (Möhlau, *B* 14, 171, 15, 2467) Prisms or fine needles Sol alcohol, ether, chloroform, benzene, and CS_2 On boiling with aniline it gives phenyl-indole Nitrous acid forms a nitrosamine $\text{C}_6\text{H}_5\text{CO CH}_2\text{NPh(NO)}$ [73°] — B'HCl prisms, decomposed by water — B'HB'r

Acetyl derivative $\text{C}_{16}\text{H}_{13}\text{NO}_2$ [127°]

Benzoyl derivative [145°] Prisms

PHENYL AMIDO-ACRIDINE

$\text{C}_{16}\text{H}_{13}\text{N}$ $\text{N} \begin{array}{c} \text{—} \text{C CH C NHPh} \\ \text{CH C CH CH} \end{array}$ [176°] Formed by heating di-phenyl *m* phenylene diamine (5 g) with ZnCl_2 (10 g) and formic acid (5 g of 90 p.c.) for 5 hours at 250° (Besthorn a Curtman, *B* 24, 2042) Flat brownish red needles, *m* sol alcohol, sl sol ether

PHENYL-AMIDO-ACRIDYL BENZOIC ACID

$\text{C}_{16}\text{H}_{13}\text{N}$ $\text{N} \begin{array}{c} \text{—} \text{C CH C NHPh} \\ \text{C(CH}_2\text{CO}_2\text{H) C CH CH} \end{array}$ [above 300°] Formed by heating di-phenyl *m* phenylene-diamine (5 g) with phthalic anhydride (10 g) at 250° (Besthorn a Curtman, *B* 24, 2047) Small red crystals, sol HOAc

PHENYL AMIDO-ACRYLIC ACID

$\text{CH}(\text{NHPh})\text{CH CO}_2\text{H}$ [194°] Formed from the product of the action of aniline on di bromo-succinic acid by boiling with potash (Reissert, *B* 20, 5105) Insol water, *v* sol alcohol — $\text{NaA}^2\frac{1}{2}\text{aq}$ silky plates

Ethyl ether EtA' [144°]

PHENYL-AMIDO-AZOBENZENE *v* Benzene-diphenylamine

DI PHENYL-AMIDO-BENZAMIDINE

$\text{C}_{16}\text{H}_{15}\text{N}_3$, $\text{e C}_6\text{H}_5(\text{NH})_2\text{C(NPh)NHPh}$ *Carbo triphenylamine* [198°] Prepared by heating *p*-nitro benzoic acid with aniline and PCl_5 for two hours at 180° and reducing the product with tin and HCl (Weith, *B* 12, 101) Formed also by the action of COCl_2 , of CBr_4 , and of CCl_4 , SO_2Cl_2 on aniline (Hofmann, *Pr* 3, 284, Bolas a Groves, *A*, 180, 173, Michler a Walder, *B* 14,

2174) Four-sided tables Split up by heating with HCl at 180° into aniline and *p* amido-benzoic acid Decomposed on distillation — B'HCl [282°] — $\text{B}'_2\text{H}_2\text{PtCl}_4$

PHENYL-DIAMIDOBENZENE *v* AMIDO-DIPHENYLAMINE

TRI-PHENYL-TRI-AMIDO-BENZENE

$\text{C}_{18}\text{H}_{15}(\text{NHPh})_3$ [198°] Made by heating phloroglucin with aniline at 210° (Minunni, *B* 21, 1984) Needles (from alcohol) — B'HCl yellow powder, melting below 100°, decomposed by warm water — $\text{B}'_2\text{H}_2\text{PtCl}_4$ [c 250°]

Tri acetyl derivative [173°] Needles.

Tri benzoyl derivative [above 304°]

PHENYL AMIDO BENZOIC ACID

$\text{C}_6\text{H}_5(\text{NHPh})\text{CO}_2\text{H}$ [222° uncor] Formed by oxidation of the methyl-o-iodide of phenyl-acridine with MMnO_2 (Claus a Nicolaysen, *B* 18, 2709) Small colourless needles *V* sol ether and chloroform, insol water

Salts — $\text{A}^+\text{Na}^4\text{aq}$ easily soluble colourless plates — $\text{A}^+\text{Ba}^5\text{aq}$ small sparingly-soluble colourless plates — A^+Ag^* very sparingly-soluble white curdy pp

PHENYL AMIDO BENZYL KETONE

$\text{C}_{16}\text{H}_{15}\text{CO CH}(\text{NH}_2)\text{C}_6\text{H}_5$ [60°-70°] Formed by reducing the mono oxim of benzil with HCl and SnCl_2 (Braun, *B* 22, 556) and by heating desyl-phthalamic acid with HCl (Neumann, *B* 23, 996) White crystalline mass, becoming brown in air — B'HCl [210°] White needles — $\text{B}'_2\text{H}_2\text{PtCl}_4$ [193°] — *Picroste* $\text{B'O}_2\text{C}_6\text{H}_5\text{N}_2\text{O}$

Carboxy-benzoyl derivative

$\text{C}_{16}\text{H}_{15}\text{CO CHPh NH CO C}_6\text{H}_5\text{CO}_2\text{H}$ [168°] Got by warming desyl-phthalimide with NaOHAq — $\text{AgC}_6\text{H}_5\text{NO}_2$, crystalline

Phenyl *p* amido benzyl ketone

$\text{C}_6\text{H}_5\text{CO CH}_2\text{C}_6\text{H}_4\text{NH}_2$ [95°] *S* 33 at 100° Formed by reducing nitro benzil or nitro deoxybenzoin with tin and HCl (Golubeff, *J R* 6, 114, 11, 101, *Bn* 3, 109) Thin needles (from dilute alcohol) — B'HCl Tables (from alcohol) *S* 25 in the cold — $\text{B}'_2\text{H}_2\text{PtCl}_4$ — $\text{B}'_2\text{H}_2\text{SO}_4$

Oxim $\text{C}_6\text{H}_5\text{C(NO)}\text{CH}_2\text{C}_6\text{H}_4\text{NH}_2$ [141°]

Crystallises from alcohol (Ney, *B* 21, 2449)

PHENYL- α -AMIDO BUTYRIC ACID

$\text{C}_{16}\text{H}_{15}\text{NO}_2$, $\text{e CH}_2\text{CH}_2\text{CH}(\text{NHPh})\text{CO}_2\text{H}$ [140°] Prepared by heating α bromo butyric acid with an ethereal solution of aniline at 100° (Duvillier, *A Ch* [5] 20, 205, Nastvogel, *B* 22, 1792) Feebly acid — B'HCl — EtA' (278°)

Phenyl β amido-butyric acid

$\text{CH}_2\text{CH}(\text{NHPh})\text{CH}_2\text{CO}_2\text{H}$ [128°] Formed by the action of baryts on the neutral isomeride (? betaine) which is obtained, together with the anilide, by boiling β -chloro butyric ether with aniline (Balbiano, *G* 10, 144, *B* 13, 312) Tufts of needles, sl sol water — BaA' (dried at 100°) Scales — B'HCl — $\text{B}'_2\text{H}_2\text{C}_2\text{O}_4$ [189°]

Anilide $\text{CHMe}(\text{NHPh})\text{CH}_2\text{CO NHPh}$ The hydrochloride B'HCl [207°] crystallises in colourless plates, sl sol hot water

Phenyl- α -amido-isobutyric acid

$\text{OMe}(\text{NHPh})\text{CO}_2\text{H}$ [185°] Formed by saponification of the nitrile or the ether (Tiemann, *B* 15, 2042, Bischoff, *B* 24, 1044) Concentric needles, *m* sol water May be sublimed

Ethyl ether EtA' [80°] (271°) Formed by heating aniline (2 mols.) with bromo isobutyric acid (1 mol.). Thick prisms.

Amide $\text{C}_6\text{H}_5(\text{NHPH})\text{CONH}_2$ [137°]
Needles, sol alcohol and HClAq

Nitrile $\text{C}_6\text{H}_5(\text{NHPH})\text{CN}$ [94°] Made by heating acetone cyanhydrin with aniline Long white prisms, sl sol hot water

Di phenyl di γ amido-butyric acid *Nitrile* $\text{CH}(\text{NHPH})\text{CH}_2\text{CH}_2\text{CN}$ [103°] Formed by heating $\text{CH}_3\text{CH}_2\text{CH}_2\text{CN}$ with aniline at 350° (Chautard, *A Ch* [6] 16, 185) Scales, sol water and alcohol

PHENYL AMIDO CHLORO v CHLORO PHENYL AMIDO

PHENYL β AMIDO CINNAMIC ACID

$\text{C}_6\text{H}_5\text{C}(\text{NHPH})\text{CHCO}_2\text{H}$
Methyl ether MeA' [93°] Formed from methyl benzoyl acetate and aniline (Knorr, *A* 245, 372) Prisms

Ethyl ether EtA' Formed in like manner (Conrad a Limpach, *B* 21, 521) Oil, v sol alcohol At 250° it yields (*Py* 1,3) oxy phenyl-quinoline and alcohol

Anilide $\text{C}_6\text{H}_5\text{C}(\text{NHPH})\text{CHCO}_2\text{NHPH}$ [133°] Needles (from alcohol), almost insol Aq

PHENYL AMIDO CITRACONIC PHENYL IMIDE? $\text{C}(\text{NHPH})\text{CO} \begin{smallmatrix} \diagup \\ \diagdown \end{smallmatrix} \text{NPh}$ [160°] Formed by heating methyl oxalacetic ether with aniline at 180° (Wislicenus a Spiro, *B* 22, 3351) Golden plates, v sol hot alcohol

PHENYL AMIDO GRESOL $\text{C}_6\text{H}_5\text{NO}$ *s.e.* $\text{NHPH C}_6\text{H}_5\text{MeOH}$ [135] [79°] (345°) Formed by heating orcin (1 pt), aniline (2 pts), and CaCl_2 (1 pt) at 260°-270° (Zega a Buch, *J pr* [2] 33, 538) Thick needles (from alcohol) Reduced by distilling over zinc dust to phenyl-m tolyl amine

Salt B'HCl crystalline powder, decomposed by water

Isomeride Oxy BENZYL ANILINE

PHENYL AMIDO CROTONIC ACID *Methyl ether* $\text{CH}_3\text{C}(\text{NHPH})\text{CHCO}_2\text{Me}$ [51°] Formed by mixing aniline with methyl acetoacetate (Conrad a Limpach, *B* 21, 1965) Prisms (from MeOH) Yields (*Py* 1,3) oxy methyl-quinoline when heated to 240°

Ethyl ether $\text{C}_6\text{H}_5\text{NO}$ *s.e.* $\text{CH}_3\text{C}(\text{NHPH})\text{CHCO}_2\text{Et}$ Formed by allowing a mixture of aniline (1 mol) and acetoacetic ether (1 mol) to stand for some days in the cold, drying with K_2CO_3 , and filtering (Knorr, *B* 20, 1397, cf Conrad a Limpach, *B* 20, 944, 22, 83) Strongly refracting oil, split up by acids and alkalis into its components At 200° it forms (*Py* 1,3) oxy methyl quinoline and a smaller quantity of oxy phenyl di methyl pyridine carboxylic ether

The anilide of acetoacetic acid (vol 1 p 19) $\text{CH}_3\text{C}(\text{CH}_3)\text{CONHPH}$ [85°], which is isomeric with phenyl-amido crotonic acid, is completely decomposed by heat, but yields the same oxy methyl quinoline on treatment with conc HClAq

PHENYL AMIDO ψ CUMYL KETONE

$\text{C}_6\text{H}_5\text{CO C}_6\text{HMe}(\text{NH}_2)$ [51346] *Benzocumide* [130°] (c 360°) Formed by saponification of its phthalyl derivative (Froehlich, *B* 17, 1804, 2674) Long yellow needles or plates, v sol. alcohol and ether, not volatile with steam B'H.PtCl_4 orange needles MeI forms $\text{C}_6\text{H}_5\text{CMe}(\text{Bz})\text{NMMe}_2$ [187°], crystallising from water in prisms (containing aq) ClCO_2Et gives

$\text{C}_6\text{HMe}(\text{Bz})\text{NHCO}_2\text{Et}$ [105°], crystallising in needles

Acetyl derivative [170°] Needles
Benzoyl derivative [227°] Needles.
Phthalyl derivative

$\text{C}_6\text{HMe}(\text{Bz})\text{N} \begin{smallmatrix} \diagup \text{CO} \\ \diagdown \end{smallmatrix} \text{C}_6\text{H}_5$ [181°] Formed by heating the phthalyl derivative of ψ cumidine with BzCl and a little ZnCl_2 at 180° Small rhombohedra, v sol hot water On saponification by alcoholic potash it yields the acid $\text{C}_6\text{HMe}(\text{Bz})\text{NHCO}_2\text{C}_6\text{H}_5\text{CO}_2\text{H}$ [195°], which crystallises in minute needles (containing aq)

PHENYL AMIDO-CYANURIC ACID v Cy anuric acid in the article CYANIC ACID

PHENYL-AMIDO-ETHANE SULPHONIC ACID $\text{C}_6\text{H}_5\text{NHCH}_2\text{CH}_2\text{SO}_3\text{H}$ *Phenyl taurine* [c 280°]

Formation —1 By heating the anilide with HCl —2 By heating chloro isethionic acid with an ethereal solution of aniline at 130° (Leymann, *B* 18, 871, James, *C J* 47, 369, *J pr* [2] 31, 415) —3 By the oxidation of phenyl thiohydantoin with KClO_4 and HCl there is formed diphenyltaurocarbamic anhydride $\text{C}_6\text{H}_5\text{N SO}_2$ [186°], which is split up by boiling baryta water into aniline, CO_2 , and phenyl taurine (Andreassch, *M* 4, 137)

Properties —Colourless leaflets, insol alcohol and ether, sol water forming an acid solution Aqueous cyanamide at 110° forms phenyltaurocyanine $\text{NH}_2\text{C}(\text{NH})\text{NPh C}_6\text{H}_5\text{SO}_3\text{H}$, which crystallises from water in plates melting above 300° — BaA' , 3aq leaflets, m sol water

Anilide $\text{NHPH C}_6\text{H}_5\text{SO}_2\text{NHPH}$ [74°] Formed, together with the anhydride $\text{C}_6\text{H}_5\text{N SO}_2$ [69°], by the action of an ethereal solution of aniline (3 mols) on $\text{CH}_2\text{Cl CH}_2\text{SO}_2\text{Cl}$ (1 mol) — B'HCl [169°] Colourless crystals

DI-PHENYL-AMIDO-ETHYL AMIDO ACETIC ACID $\text{NHPH CH}_2\text{CH}_2\text{NPh CHCO}_2\text{H}$ [116°] Formed from oxy di phenyl pyrazine hexahydrate and alcoholic potash (Bischoff, *B* 23, 2026) Prisms, sol ether Converted into the parent substance by boiling with water or alcohol

PHENYL AMIDO-ETHYL KETONE

$\text{C}_6\text{H}_5\text{CO C}_6\text{H}_5\text{NH}_2$ Formed by the action of boiling HClAq on $\text{C}_6\text{H}_5\text{Bz N C}_6\text{O}_2\text{C}_6\text{H}_5$ [85°], which is got from $\text{C}_6\text{H}_5\text{CO C}_6\text{H}_5\text{Br}$ and potassium phthalimide (Schmidt, *B* 22, 3251) — B'HCl crystalline, v *s.e.* sol water and alcohol — B'H.PtCl_4 — $\text{B'C}_6\text{H}_5\text{N}_2\text{O}$, [160°] Needles

PHENYL-AMIDO-ETHYL-PHTHALIMIDE

$\text{C}_6\text{H}_5\text{C}_6\text{O}_2\text{NCH}_2\text{CH}_2\text{NHPH}$ [100°] Formed by heating bromo ethyl phthalimide with aniline at 150° (Gabriel, *B* 23, 2224) Lemon yellow needles, v sol dilute HClAq Converted by NaOHAq into $\text{C}_6\text{H}_5(\text{CO}_2\text{H})\text{CO NH C}_6\text{H}_5\text{NHPH}$ [120°-130°], a crystalline powder

PHENYL-AMIDO-FUMARIC IMIDE

$\text{C}(\text{NHPH}) \begin{smallmatrix} \diagup \text{CO} \\ \diagdown \end{smallmatrix} \text{CHCO} \begin{smallmatrix} \diagup \text{CO} \\ \diagdown \end{smallmatrix} \text{NH}$ [202°] Formed by heating bromo-fumarimide with aniline (Löschner, *B* 21, 2718) Golden plates, sl sol hot water

PHENYL-AMIDO-GLUTACONIC PHENYL-IMIDE $\text{C}_6\text{H}_5\text{N}_2\text{O}$ *s.e.*

$\text{C}(\text{NHPH}) \begin{smallmatrix} \diagup \text{CH}_2\text{CO} \\ \diagdown \end{smallmatrix} \text{CHCO} \begin{smallmatrix} \diagup \text{CH}_2\text{CO} \\ \diagdown \end{smallmatrix} \text{NPh}$ [275°] Formed from acetone dicarboxylic ether and aniline (Emery, *B* 23, 3764) Yellowish plates, v sl sol alcohol

PHENYL-AMIDIMIDO-ETHENYL-*o*-AMIDO-PHENYL MERCAPTAN $C_6H_5N_3S$ probably

$C_6H_5\langle\overset{N}{S}\rangle C(NHPh)NH$ [118°] Formed together with the di-phenyl derivative by heating amido-imido-ethenyl-*o*-amido-phenyl-mercaptan with aniline, ammonia being evolved (Hofmann, *B* 20, 2254) Plates Sol. alcohol and ether

Salts $-B'H_2Cl.PtCl_3$, small needles, m sol water $-B'HClAuCl_2$, soluble yellow plates

Di-phenyl-amidimido-ethenyl-*o*-amido-phenyl-mercaptan $C_{22}H_{15}N_3S$ which is probably

$C_6H_5\langle\overset{N}{S}\rangle C(NHPh)NPh$ [129°] Obtained as above White silvery plates V sol alcohol and ether

Salts $-B'H_2Cl.PtCl_3$, dark-yellow sparingly-soluble plates $-B'H_2Cl_2AuCl_2$, yellow needles

PHENYL-AMIDO-JUGLONE v *Juglone***PHENYL-AMIDO MALEIC ACID**

The phenylimide $NHPh.C_6H_4.C_4O_2.NPh$ [232°] is formed, together with the mono-anilide $NHPh.C_6H_4(CO_2H)CONHPh$ [176°], by boiling dibromo succinic acid or chloro- or bromo-maleic or fumaric acids with aniline and water (Reissert a Tiemann, *B* 19, 626, Michael, *B* 19, 1377) The phenylimide is also formed by heating aniline with oxalacetic ether at 140° (Wishlencus a. Spiro, *B* 22, 3350)

PHENYL-AMIDO MALONIC ACID *Anilide* $CH(NHPh)(CO NHPh)_2$ [162°] Formed by boiling chloro-malonic acid with aniline (Conrad a Bischoff, *A* 209, 231) Prisms

PHENYL-AMIDO-METHENYL-AMIDO-

NAPHTHOL $C_{10}H_7\langle\overset{O}{N}\rangle C(NHPh)$ [168°].

Formed from benzene-azo (β) naphthol by heating with CS_2 at 250° (Jacobson, *B* 21, 419) Small needles, v e sol cold alcohol With $HClAq$ at 190° it yields amido (β) naphthol, aniline, and CO_2 —Picrate [210°] Acetate [120°-130°]

PHENYL-AMIDO-METHENYL-AMIDO-

PHENYL MERCAPTAN $C_6H_5\langle\overset{N}{S}\rangle C(NHPh)$

[159°] Formed from $C_6H_5NS.CCl$ and aniline (Hofmann, *B* 12, 1130) Formed also by heating phenyl thiocarbimide with azobenzene at 270° (Jacobson a Frankenbacher, *B* 24, 1410) Needles, sol alcohol, insol water Decomposed by alcoholic potash at 200° into aniline and amido-phenyl mercaptan $-B'HCl -B'HAuCl_3 -B'_2H_2PtCl_6 -B'_2C_2H_2N_2O_7$ [222°]

Acetyl derivative [167°] Needles

PHENYL-AMIDO-METHENYL-AMIDO-

PHENOL $C_6H_5\langle\overset{N}{O}\rangle C(NHPh)$ [178°] Formed by boiling $C_6H_5\langle\overset{N}{O}\rangle CSH$ with aniline (Kalckhoff, *B* 16, 1826) Needles, sol alcohol, ether, and $HOAc$ May be distilled. Weak base $-B'H_2PtCl_6$

PHENYL-AMIDO-METHENYL-AMIDO-

PHENYL-MERCAPTAN $C_6H_5N_3S$ *see*

$C_6H_5\langle\overset{N}{S}\rangle C(NHPh)$ [159°] Formed by heating chloro-phenyl-thiocarbimide with aniline (Hofmann, *B* 12, 1129, 11, 12). Needles

PHENYL-AMIDO-METHENYL-BENZAMIDESULPHIM

$C_6H_5\langle\overset{NS}{N}\rangle C(NHPh)$ [174°] Formed by boiling benzamidoxim (1 mol) with phenyl thiocarbimide (2 mols) for 10 hours (Koch, *B* 24, 394) White plates, insol water, sol alcohol, ether, and conc $HClAq$, insol water and alkalis Conc $HClAq$ at 150° decomposes into S, H_2S , aniline, $HOBz$, and NH_3 Bromine forms $CPh.N_3S.CNHC_6H_4Br$ [14°] Yields a nitrosamine [119°] and an acetyl derivative [196°], both being crystalline

PHENYL-AMIDO METHYL-ACRIDINE

$C_6H_5\langle\overset{N}{N}\rangle CCH.CNHPh$ [216°] Fc med

by heating the diacetyl derivative of *m*-phenylene-di phenyl diamine (1 pt) with $ZnCl_2$ (2 pts) for 6 hours at 250° (Besthorn a Curtman, *B* 24, 2044) Brownish red needles, v sol alcohol sl sol ether Split up by $HClAq$ at 250° into aniline and oxy methyl-acridine

DI-PHENYL-DI-AMIDO METHYLENE-*o*-PHENYLENE-DIAMINE

$C_6H_5\langle\overset{NH}{NH}\rangle C(NHPh)_2$ [160°] Formed from di-phenyl cyanamide $C(NPh)_2$ and *o*-phenylene diamine at 135° (Moore, *B* 22, 1635, 3190) Needles (from benzene ligroin), insol ligroin, v sol alcohol Yields with nitrous acid a nitrosamine $C_6H_5N_3(NO)_2$, $C(NPhNO)_2$ [138°] Salts $-B'_2H_2Cl_2$, needles, v sol water $-B'_2H_2SO_4$, needles, sl sol hot water $-B'_2H_2PtCl_6$, *see* [165°]

Tetra-benzoyl derivative [148°]

*Tetra-phenyl-tetra-amido-di-methylene-*o*-phenylene-diamine* $C_6H_5N_4C(NHPh)_4$ [139°] Formed by heating $C(NPh)_2$ with *o*-phenylene diamine at 200° (Moore) Colourless prisms v sol alcohol Split up by heat into aniline and the preceding body

Salts $-B'_2H_2Cl_2$, needles, v sol hot water $-B'_2H_2PtCl_6 -B'_2H_2SO_4$, prisms, v sol hot water

Tetra-acetyl derivative

$C_6H_5N_4C(NPhAc)_4$ [126°] White prisms

Tetra-benzoyl derivative [182°]

DI-PHENYL-DI-AMIDO-METHYLENE-*o*-PHENYLENE-PHENYL GUANIDINE

$C_6H_5\langle\overset{N}{N}\rangle C(NPh)\langle\overset{N}{N}\rangle C(NHPh)_2$ [188°]. Formed from phenylene phenyl-guanidine and $C(NPh)_2$ at 210° (Keller, *B* 24, 2506) The homologous compound from *o*-tolylene phenyl-guanidine melts at 200° while the corresponding body from *o*-tolylene-*p*-tolyl guanidine melts at 176° All three compounds crystallise from alcohol in white needles, sol benzene

PHENYL-AMIDO-METHYL-DI-ETHYL-

PYRIMIDINE $N\langle\overset{Cet}{Cet}\rangle CMe_2\langle\overset{N}{N}\rangle C(NHPh)$

[99°] Formed from chloro-methyl-di-ethyl pyrimidine (derived from cyanethine) by heating with aniline and alcohol at 220° (Von Meyer *J. pr* [2] 89, 274) Needles, v sol. alcohol. $-B'_2H_2PtCl_6$, needles

PHENYL-AMIDO-METHYL-MALONIC ACID *Amic ether*

$CO_2Et.OMe(NHPh)CONH_2$ [86°] Formed by dissolving the nitrile in cold conc. H_2SO_4 and

pouring into water (Gersen, *B* 19, 2965) The nitrile $\text{CO}_2\text{Et CMe}(\text{NHPh})\text{CN}$ [102°] is formed by heating $\text{CO}_2\text{Et CMe}(\text{OH})\text{CN}$ with an alcoholic solution of aniline for 24 hours at 80°. The amic ether crystallises in white needles, sol hot water, and is converted by boiling aqueous NaOH into $\text{CH}_2\text{CH}(\text{NHPh})\text{CO}_2\text{H}$

PHENYL-L-(a)-AMIDO METHYL-NAPHTHO-

PHENAZINE $\text{C}_6\text{H}_5\text{Me} \begin{smallmatrix} \text{N} \\ \text{N} \end{smallmatrix} \text{C}_6\text{H}_4\text{NHPh}$

[214°]. Formed by heating (1,3,4)-tolylene-diamine with benzene azo phenyl (a)-naphthylamine hydrochloride and alcohol at 140° (Eicker, *B* 23, 306) Brass-yellow needles, forming yellowish-red solutions with yellowish-green fluorescence. — $\text{B}'\text{H}_2\text{PtCl}_6$. minute dark-red needles.

PHENYL-AMIDO-DI-METHYL-PYRIDINE

$\text{N} \begin{smallmatrix} \text{CMe CH} \\ \text{CMe CH} \end{smallmatrix} \text{CNHPh}$. *Phenyl-amido lutidine* [150°] (C a E); [144°] (G a M). (235°-238°) Obtained by heating chloro-di-methyl-pyridine with aniline at 185° (Conrad a Epstein, *B* 20, 165) Formed also by heating the corresponding oxy-di-methyl-pyridine (1 mol) with phenyl cyanate (1 mol.) and benzene at 100°, CO_2 being evolved (Goldschmidt a Meissler, *B* 23, 274) White crystalline solid. V sol. alcohol, ether, and dilute acids With H_2SO_4 and a trace of HNO_3 it gives a blood-red colouration — $\text{B}'\text{H}_2\text{PtCl}_6$ [209°] (C a E), [204°] (G a M), microscopic yellow needles

PHENYL-AMIDO-DI-METHYL-PYRROLE

$\text{NHPh N} \begin{smallmatrix} \text{CMe CH} \\ \text{CMe CH} \end{smallmatrix}$ [92°] (c 270°) Formed by heating its dicarboxylic acid, and also by the action of phenyl hydrazine on acetyl acetone (Knorr, *B* 18, 1568, 22, 170) Crystalline mass, volatile with steam Insol water and alkalis, sol conc HClAq Reddens pine wood moistened with HClAq

PHENYL-AMIDO-DI-METHYL-PYRROLE-

DICARBOXYLIC ACID $\text{NHPh N} \begin{smallmatrix} \text{CMe C CO}_2\text{H} \\ \text{CMe C CO}_2\text{H} \end{smallmatrix}$ Needles. Got from its ether $\text{Et}_2\text{A}''$ [127°], which is formed by mixing solutions of di acetyl-succinic ether and phenyl-hydrazine in HOAc (Knorr, *B* 18, 804, 1568).

PHENYL (Py 1)-AMIDO-(Py 3)-METHYL

QUINOLINE $\text{C}_6\text{H}_5 \begin{smallmatrix} \text{C}(\text{NHPh})\text{CH} \\ \text{N} \text{CMe} \end{smallmatrix}$ [151°]. Formed by heating chloro-methyl-quinoline with aniline at 190° (Conrad a Limpach, *B* 20, 953) Prisms, al sol ether, v sol alcohol.

Phenyl-(Py 3)-amido-(Py 1)-methyl-quinoline $\text{C}_6\text{H}_5\text{N}_2$ [130°] Formed by heating (Py 3,1)-chloro-methyl quinoline with aniline (Knorr, *A* 236, 102) Shining plates. — $\text{B}'\text{H}_2\text{PtCl}_6$ [235°] Yellow needles, insol Aq. *Phenyl-(Py 1)-amido-tri-methyl-quinoline*

$\text{C}_6\text{H}_5\text{Me} \begin{smallmatrix} \text{C}(\text{NHPh})\text{CH} \\ \text{N} \text{CMe} \end{smallmatrix}$ [150°]. Formed by heating the corresponding chloro-tri methyl-quinoline with aniline at 185° (Conrad a Limpach, *B* 21, 538). Prisms (from dilute alcohol).

PHENYL-AMIDO-METHYL-THIAZOLE

$\text{CMe.N} \begin{smallmatrix} \text{CH} \\ \text{S} \end{smallmatrix} \text{CNHPh}$ [117°]. Formed by heating oxy-methyl-thiazole with aniline (Hantzsch a Weber, *B* 20, 8130), and by the action of chloroacetone on phenyl-thio-urea (Traumann, *A* 249, *Yac. III*.

47) Small white needles (from alcohol) Yields aniline when heated with HClAq at 240°

PHENYL-AMIDO-NAPHTHALENE v.

PHENYL-NAPHTHYL-AMINE

Di-phenyl-di-amido naphthalene v. *Di*

PHENYL-NAPHTHYLENE-DIAMINE.

Tri-phenyl tri-amido naphthalene

$\text{C}_6\text{H}_5 \begin{smallmatrix} \text{C}(\text{NHPh})\text{CH} \\ \text{C}(\text{NHPh})\text{CNHPh} \end{smallmatrix}$ [148°] One of the products formed by heating benzene azo (a)-naphthylamine with aniline at 160° (Fischer a Hepp, *A* 256, 251) Needles, v sol alcohol. Yields phenyl-rosinduline and a yellow substance [140°] on oxidation

Tetra-phenyl-tetra-amido-naphthalene

$\text{C}_6\text{H}_5(\text{NHPh})_4$ [191°]. Accompanies the preceding body On oxidation it yields phenyl-amido phenyl-rosinduline, di-phenyl-di-amido-naphthoquinone, and a body melting at 169°.

PHENYL-AMIDO (a)-NAPHTHOQUINONE

$\text{C}_6\text{H}_5\text{NO}_2$, i.e. $\text{C}_6\text{H}_5 \begin{smallmatrix} \text{CO CNHPh} \\ \text{CO CH} \end{smallmatrix}$ [191°].

Formed by heating (a)-naphthoquinone with an alcoholic solution of aniline (Zincke, *B* 12, 1645), by boiling phenyl-amido (β) naphthoquinone with HCl , and by the action of aniline and HOAc on oxy-(a)-naphthoquinone (Liebermann, *B* 14, 1665) Long red needles, v sol. hot alcohol and ether, insol cold alkalis Alcoholic potash forms a purple solution Boiling dilute NaOH splits it up into aniline and oxy-(a)-naphthoquinone.

Anilide $\text{C}_6\text{H}_5 \begin{smallmatrix} \text{CO} \\ \text{C}(\text{NHPh})\text{CH} \end{smallmatrix}$ *Di*.

phenyl di-imidonaphthol [180°]

Formation — 1 By heating each of the naphthoquinone oxims with aniline and HOAc at 100° (Fuchs, *B* 8, 1023, Brömme, *B* 21, 893) — 2 By heating di-imido-(a)-naphthol hydrochloride with aniline at 120° (Göbe, *B* 13, 124) — 3 By boiling (β)-naphthoquinone with an alcoholic solution of aniline (Zincke, *B* 15, 481) — 4 By heating di-bromo-(a)-naphthol with aniline (Meldola, *C J* 45, 157) — 5 By the action of aniline and aniline hydrochloride at 100° on benzene-azo-(a) naphthol, on benzene-azo-(a)-naphthylamine, and on similar bodies (Fischer a Hepp, *B* 21, 679)

Properties — Orange red needles (from alcohol), forming orange solutions in benzene, acetone, and CHCl_3 . Weak base Its solution in HOAc is orange when hot, but crimson when cold Hot HClAq forms a red solution Insol alkalis Not attacked by boiling with alkalis, or by AqO , AcCl , and MeI H_2SO_4 at 120° yields oxy-naphthoquinone and aniline. Zinc dust and HOAc reduce it to aniline and naphthalene. Alcoholic HCl or EtBr at 150° yields aniline and phenyl-amido-(a)-naphthoquinone HNO_3 forms a di-nitro- derivative [148°].

Salts. — $\text{B}'\text{HCl}$: golden-green plates, forming a violet solution in alcohol. Decomposed by water — $\text{B}'\text{H}_2\text{PtCl}_6$: bronze needles. — $\text{B}'\text{H}_2\text{ZnCl}_6$: bronze-green plates. — $\text{B}'\text{HI}$: black needles. — $\text{B}'\text{H}_2\text{SO}_4$ — $\text{B}'\text{HNO}_3$: plates

Di-anilide $\text{C}_6\text{H}_5(\text{NHPh})_2$ [189°].

A product of the action of benzene-azo-(a)-naphthylamine on aniline (Fischer a Hepp, *A* 252, 246). Orange-yellow prisms.

Phenyl-amido-(β)-naphthoquinone

$C_{16}H_9O_2(NHPh)$ or $C_{16}H_8O_2 \begin{matrix} \text{CO} \\ \text{C}(NHPh) \end{matrix} \begin{matrix} \text{OH} \\ \text{OH} \end{matrix}$ *Oxy-*

(α)-naphthoquinone amide. (β)-Naphthoquinone amide' [245°-250°] Prepared by adding aniline to an alcoholic solution of (β)-naphthoquinone (Zincke, *B* 14, 1494, 15, 279, Liebermann, *B* 14, 1664). Red needles, sl sol alcohol. Weak acid. Readily converted by boiling with HOAc into the isomeric phenyl-amido-(α)-naphthoquinone. Boiling HClAq splits it up into aniline and oxy-(α) naphthoquinone. Nitrous acid passed into a solution in alcohol and HOAc forms a nitroso derivative $C_{16}H_9N_2O_2$, crystallising in red needles, and converted by NaOHAlq into a yellow body [217°].

Salts—BaA', red needles, sl sol water—PbA', brown pp—ZnA',—HgA',—AgA' red pp. *Methyl ether MeA'* [161°] Formed from the Ag salt and MeI. Yellow needles, sol alcohol and ether.

Ethyl ether EtA' [104°] Prisms

n-Propyl ether PrA' [104°] Prisms

Isopropyl ether IPA' [100°]

Di-phenyl-amido (α)-naphthoquinone

$C_{16}H_8O_2(NPh)_2$ [164°] Formed by boiling (α) naphthoquinone (5 g) with diphenylamine (6 g), alcohol, and conc HClAq (10 cc) for twenty minutes, and ppg with water (Plimpton, *O J* 37, 644). Needles (from alcohol).

Acetyl derivative $C_{16}H_8AcNO_2$ [173°]

PHENYL-AMIDO NAPHTHYL-UREA

$NHPh \cdot CO \cdot NH \cdot C_6H_4 \cdot NH_2$. Formed from (1,2)-naphthylene diamine and phenylcyanate (Schiefelin, *B* 22, 1377, cf Goldschmidt, *B* 23, 502). Granules (from alcohol), not fused at 335°.

DI-PHENYL-DI-AMIDO-OIAZATHIOLE

$S \begin{matrix} \text{C}(NHPh)N \\ \text{C}(NHPh)N \end{matrix}$ [181°] Formed by adding hydrogen peroxide solution (8 p.c.) to phenylthio urea dissolved in dilute alcohol, acidulated with HCl (Hector, *B* 22, 1176). Needles, insol water, v sl sol cold alcohol. Nitrous acid forms $C_{16}H_{11}(NO)_2S$, a green insoluble powder, exploding at 179°. Cyanogen passed into the warm alcoholic solution ppts $C_{16}H_{12}N_2S$, crystallising in needles—B'HNO₂—B'H₂PtCl₄—B'AgNO₃, aq granules, v sl sol hot alcohol.

Acetyl derivative $C_{16}H_{11}AcN_2S$ [238°].

Benzoyl derivative $C_{16}H_{11}BzN_2S$ [238°].

PHENYL-AMIDO-PALMITIC ACID

$C_{16}H_{21}(NHPh)O_2$ [142°] Formed by boiling α -bromo palmitic acid with aniline (Hell a Jordanoff, *B* 24, 942). Sol ether and benzene.

PHENYL-AMIDO-PHENOL v **Oxy-di-**

PHENYL-AMINE

Di-phenyl-di-amido-phenol *Hydrochloride* $(C_6H_4(NHPh)_2OH)HCl$ [192°] Formed from azophenin, Sn, and HCl in presence of AcOH (Fischer a Hepp, *A* 256, 260). Plates.

PHENYLAMIDO-PHENYL ACETIC ACID

$C_6H_5NO_2$ v $C_6H_5CH(NHPh)CO_2H$ [164°-168°] (S) Formed by saponification of the nitrile (Tiemann a Priest, *B* 15, 2030), and also by boiling bromo phenyl-acetic acid with aniline and alcohol (Stöckemus, *J* 1878, 779). Plates, subliming at 176° as slender white needles (T a F.). V. sl sol water, sol alcohol—HA'HCl·molecules, decomposed by water—HA'HNO₂; thin needles—BaA'; radiating tress.

Ethyl ether EtA' [84°]. Needles.

Amide $C_6H_5CH(NHPh)CONH_2$. Silky

plates, v sol alcohol, nearly insol ligroin.

Nitrile $C_6H_5CH(NHPh)CN$ [85°] Formed by heating an alcoholic solution of the cyanhydrin of benzoic aldehyde with aniline at 100°. Formed also by the action of KCy and HCl on benzylidene-aniline (Cech, *B* 11, 246). Needles or prisms, sol ether and hot alcohol.

Phenyl amido-di-phenyl-acetic acid

$CPh_2(NHPh)CO_2H$ [168°] Formed by passing HCl into an alcoholic solution of benzoic acid and treating the resulting CPh_2ClCO_2Et with aniline (Klinger a Standke, *B* 22, 1212). Needles, v sl sol water. Conc H_2SO_4 forms a yellow solution, becoming crimson on warming.

Methyl ether MeA' [107°]. Monoclinic

Ethyl ether EtA' [115°].

PHENYL-AMIDO-PHENYL-ACRIDINE

$C_6H_5 \begin{matrix} \text{N} \\ \text{C} \end{matrix} \begin{matrix} \text{CH} \\ \text{CH} \end{matrix} \begin{matrix} \text{C} \\ \text{C} \end{matrix} \begin{matrix} \text{CH} \\ \text{CH} \end{matrix} \begin{matrix} \text{N} \\ \text{N} \end{matrix} \begin{matrix} \text{CH} \\ \text{CH} \end{matrix} \begin{matrix} \text{C} \\ \text{C} \end{matrix} \begin{matrix} \text{CH} \\ \text{CH} \end{matrix} \begin{matrix} \text{N} \\ \text{N} \end{matrix}$ [197°] Formed by

heating the di-benzoyl derivative of *m* phenylene di-phenyl di-amine (10 g) with ZnCl₂ (25 g) at 250° (Besthorn a Curtman, *B* 24, 2045). Brownish red needles, m sol alcohol, ether, and benzene. The dilute ethereal solution exhibits a greenish fluorescence. The hydrochloride is v sl sol water, and forms a red solution in alcohol.

DI-PHENYL-DI-AMIDO-TRI-PHENYL-CARBINOL $C_{16}H_{12}N_4O_2$ v

$CPh(OH)(C_6H_5NHPh)_2$. Formed by heating di-phenylamine with benzyl chloride and oxidising the product with arsenic acid. Formed also by heating diphenylamine with ZnCl₂ and benzoyl chloride or benzoic trichloride, the resulting chloride $C_{16}H_{12}N_4Cl_2$ being decomposed by alcoholic NH₃ (Meldola, *C J* 41, 187). Amorphous, v sol ether, sol hot alcohol and acetone. The chloride $C_{16}H_{12}N_4Cl_2$, called 'diphenylamine green,' forms bronzed granules, and, in alcoholic solution, dyes wool bluish green, and gives a dark green pp with H₂PtCl₆. Hot H₂SO₄ converts the chloride into the sulphonic acid $C_{16}H_{12}N_4SO_3$, ppd by water in dark green flakes.

Tri-phenyl tri-amido tri-phenyl-carbinol. The chloride $CCl(C_6H_5NHPh)_3$, or 'diphenylamine blue,' is obtained by heating diphenylamine (1 pt) with oxalic acid (2 pts) (Willm a Girard, *D P J* 221, 192, Hausdörfer, *B* 23, 1963). Formed also by heating pararosaniline with aniline. Brownish red powder, v sol hot aniline and nitro benzene, sl sol cold alcohol.

Hexa-phenyl tri-amido tri-phenyl carbinol $C(OH)(C_6H_5NHPh)_3$. Formed by the action of alcoholic NH₃ on the chloride $CCl(C_6H_5NHPh)_3$, which is got by heating triphenylamine with COCl₂ at 190° (Heydrich, *B* 19, 758). White amorphous pp, forming a bluish-violet solution in H_2SO_4 , turned blue by addition of alcohol.

PHENYLAMIDO-PHENYL-ISOCROTONIC ACID $C_6H_5CH=CHCH(NHPh)CO_2H$ [154°].

Formed by saponification of the nitrile (Peine, *B* 17, 2116). Minute needles, v sol alcohol and ether.

Amide [171°]. Thin plates.

Nitrile $C_6H_5CH=CHCH(NHPh)CN$ [180°] Formed by heating $CHPhCH=CH(OH)CN$ with aniline in alcoholic solution. Crystalline solid, sol hot alcohol.

PHENYLAMIDO-PHENYL-ETHYL-KETONE $C_6H_5(NHPh)CO \cdot C_2H_5$. *Homophenacylamide*.

[88°] Formed from phenyl bromo ethyl ketone and aniline in alcoholic solution (Pampel & Schmidt, *B* 19, 2897) Yellow crystals Yields an acetyl derivative [103°]

PHENYL AMIDOPHENYL HYDRAZINE

*Acetyl derivative $C_6H_5N_2H_2C_6H_5NHAc$
[146°] Gg by reduction of $C_6H_5N_2C_6H_5NHAc$
with alcoholic ammonium sulphide (Schultz, *B*
17, 463) Glistening plates, v sol alcohol and
ether $SnCl_4$ reduces it to aniline and *p*-
phenylene diamine Conc $HClAq$ forms a blue
compound

PHENYLAMIDO DI PHENYL-METHANE

$\text{JH}_2\text{PhC}_6\text{H}_4\text{N}(\text{HPh})$ [c 89°] Formed by heating diphenylamine with benzyl chloride and ZnCl_2 at 100° (Meldola, *C J* 41, 198) White powder, v sol benzene, insol alcohol

Di phenyl-di-amido-tri phenyl-methane

$\text{C}_{21}\text{H}_{18}\text{N}_2$ *see* $\text{CHPh}(\text{C}_6\text{H}_5\text{NHP})_2$, [c 170°]
Formed by heating diphenylamine with benzyl
dene chloride, and also from diphenylamine,
benzoic aldehyde, and ZnCl_2 (Meldola, *C J* 41,
192) White granules, v sol ether, yielding
diphenylamine green on oxidation

Reference — CHLORO - DIPHENYLDIAMIDOTRI
PHENYLMETHANE

PHENYLAMIDO-PHENYL METHYL PYR

IMIDINE CPh $\begin{matrix} \text{N CMe} \\ \diagup \quad \diagdown \\ \text{N} \quad \text{C(NHPh)} \end{matrix} \text{CH} \text{ [150°-153°]}$
 Formed from chloro phenyl methyl pyrimidine
 and aniline (Pinner, *B* 18, 2852) —B'HNO₃
 [87°] —H'HC! [240°] Long slender needles

PHENYLAMIDO PHENYL PROPIONIC
ACID Nitrile $\text{CH}_2\text{CPh}(\text{NHPh})\text{CN}$ [152°]

Formed by digesting an ethereal solution of acetophenone cyanhydrin and aniline at 50° (Jacoby, *B* 19, 1515) White prisms (from alcohol) With alcoholic hydroxylamine it yields the oxim of acetophenone Alcoholic HCl gives *s* tri phenyl benzene

Amide $\text{CH}_3\text{CPh}(\text{NHPh})\text{CO NH}_2$ [119°]

Formed from the nitrile by heating with H_2SO_4 at 100° and pouring into water Insol water

PHENYL AMIDO TRI PHENYL PYRROLE

$\text{CH CPh} \searrow$
 $\text{CPh CPh} \searrow \text{N NPh} \quad [232^\circ] \quad \text{Formed from}$
anhydrazetophenone benzil (1 mol) and phenyl
hydrazine (2 mols) in alcoholic solution at 100°
(Japp. B 21, 551, 22, 2885) Yellow needles

PHENYL AMIDO TRI PHENYL QUINOX-
ALINE DIHYDRIDE

$[14^3]C_6H_3(NHPh) \left\langle \begin{array}{c} NPh.CHPh \\ N=CPh \end{array} \right. [223^\circ] \text{ Formed}$
by heating di-phenyl (1,3,4) tri amido benzene
with benzoïn at 160° (Fischer, B 24, 722)
Yellow plates, sl sol alcohol, forming a yellow
solution with green fluorescence

PHENYLAMIDO PHENYL ROSINDULINE

[$\frac{1}{2}$] $C_6H_4(NPh) \left\langle \frac{N-C}{NPh-C} OH C(NHPh) \right\rangle C_6H_4$
 [192] Formed by heating benzene-azo (α -naphthyl)-amine with aniline and benzene (Fischer & Hopp, *A* 262, 297). Formed also by oxidation of (1,2,3,4) tetra phenyl-tetra amido-naphthalene (Fischer & Hopp, *A* 256, 252). Bronzed plates, forming a dark-green solution in some H_2SO_4 , becoming reddish-violet on dilution with HCl bronzed plates

TRI-PHENYL-TRI-AMIDO-DI-PHENYL-
TOLYL-CARBINOL

$$\text{C}(\text{OH})(\text{C}_6\text{H}_4\text{NHPh})_4(\text{C}_6\text{H}_4\text{NHPh}) \quad \text{Aniline blue.}$$

Lyons blue *Triphenylrosaniline* Formed, to
gether with mono and di phenyl rosaniline by
heating rosaniline acetate or benzoate with
aniline (Girard & De Laire, *D P J* 162, 297,
Hofmann, *Pr* 12, 578, 13, 9) Prepared by
heating rosaniline (1 pt) with aniline (10 pts)
and a little benzoic acid, dissolving in alcoholic
 NH_3 , and pouring into water White pp, insol
water, v sol alcohol, m sol ether Yields di
phenyl amine on distillation. Reduced by zinc
and HCl to tri phenyl tri amido di phenyl tolyl
methane or tri phenyl leucaniline (Hofmann,
C R 57, 25)— $\text{C}_{24}\text{H}_{22}\text{N}_3\text{Cl}$ *Spirit soluble blue*
Opal blue Small brassy crystals, insol water and
ether, sl sol alcohol Becomes brown at 100°
Left with coppery lustre on evaporation of the
alcoholic solution Blue dye— $(\text{C}_{24}\text{H}_{22}\text{N}_3)_2\text{SO}_4$, v
sl sol alcohol The salts of mono phenyl ros-
aniline dye violet, those of di phenyl rosaniline
a bluish violet Tri phenyl rosaniline yields
mono, di, tri, and tetra sulphonic acids, the
Na salts of which are v sol water and dye wool
blue (Nicholson, *Bulk*, *B* 5, 417)

The mono and di sulphonic acids are known as *Nicholson's* or *Alkali blue*, the tri sulphonic acid as *Soluble blue*.

Reference — TRI CHLORO TRI PHENYL-ROBANIL
INE

TRI PHENYL AMIDO PHOSPHIDE OXIDE

PO(NHPh), [208°] Formed from POCl₃ and aniline (Michaelis & Soden, *J* 229, 334) Six sided plates, insol water, sl sol ether and alcohol, v sol HOAc Yields PO(NHC₆H₄Br)₃, crystallising in needles [253°] The compound HO PO(NHPh)₂, [197°] may be got by treating aniline with ether and POCl₃, followed by water

TRI-PHENYL TRI-AMIDO-PHOSPHINE

SULPHIDE $\text{PS}(\text{NHPh})_2$, [153°] Formed from aniline and P_2S_5 below 150° (Knop, B 20, 3352) Monoclinic crystals

PHENYLAMIDO PROPIONIC ACID

$\text{C}_6\text{H}_{11}\text{NO}_3$, ϵ $\text{CH}_2(\text{NHPh})\text{CO}_2\text{H}$ [162*]
Formed by saponification of the nitrile or of the
ether (Tiemann & Stephan, *B* 15, 2036,
Nastvogel, *B* 22, 1792, 23, 2010). Plates (from
hot water), sol alcohol. May be sublimed.
Yields a hydrochloride decomposed by water,
and a white silver salt.

Ethyl ether EtA' (272°) SG $\frac{19.8}{19.8}$ 1.060

Formed by heating α bromo propionic ether with aniline on the water-bath Oil.

Formyl derivative

$\text{CH}_3\text{CH}(\text{NPhCHO})\text{CO}_2\text{H}$ Prepared from
 $\text{NaNPh}(\text{CHO})$ and α -bromo-propionic ether
(Paal & Otten, *B* 23, 2597) Large prisms,
sol alcohol and ether — BaA' , aq white powder

Acetyl derivative $\text{CH}_3\text{CH}(\text{NaCPh})\text{CO}_2\text{H}$, [143°] Plates (from hot benzene), sl sol cold water — $\text{NaC}_6\text{H}_5\text{NO}_2$ 3aq plates — EtA' (294° 298°) Oil, formed from sodium acetalide and a bromo propionic ether

Amide $\text{CH}_2\text{CH}(\text{NHPh})\text{CONH}_2$ [141°]
 Plates, sol. alcohol and hot water

Anilide $\text{CH}_3\text{CH}(\text{NPh})\text{CO NPh}$ [126°].
Needles, v sol hot water

Nitrile $\text{CH}_3\text{CH}(\text{NHPH})\text{CN}$ [92°] Formed by heating the cyanhydrin of acetic aldehyde with aniline (T & S) White plates, sol also hot and ether Weak base — B^+HCl [86°]

Isomerides: v AMIDO-PHENYL PROPIONIC ACID.

Di-phenyl-di-amido-propionic acid. Nitrile
 $\text{CH}(\text{NHPh})_2\text{CH}_2\text{CN}$ [118°]. Formed by heating CHOCH_2CN with aniline for 30 hours at 310° (Chautard, *A. Ch.* [8] 16, 180). Needles, v. sol. water and benzene — B'HOI . — B' , H_2PtCl_6 : small yellow needles, v. sol. water

Reference.—TRI-BROMO PHENYL-AMIDO-PROPIONITRILE

PHENYL- β -AMIDO-PROPYL-PHTHALIMIDE
 $\text{C}_6\text{H}_4\text{C}_6\text{H}_4\text{NCH}_2\text{CHMe.NHPh}$ [93°] Formed from β -bromo-propyl-phthalimide and aniline (Seitz, *B.* 24, 2631). Yellow needles. Decomposed by HCl at 190° into phthalic acid and propylene-diamine

Phenyl- γ -amido-propyl-phthalimide
 $\text{C}_6\text{H}_4\text{C}_6\text{O}_2\text{NCH}_2\text{CH}_2\text{CH}_2\text{NHPh}$ [89°] Formed by heating γ -bromo-propyl phthalimide (54 g) with aniline (37 g) at 150° (Goldenring, *B.* 23, 1168). Yellow crystals (from boiling ligroin). Split up by heating with HCl , yielding phenyl-trimethylene-diamine

PHENYLAMIDO-PROPYL-UREA
 $\text{NHPh.C}_6\text{H}_4\text{NH.CO.NH}_2$ [98°] Formed by warming $\text{C}_6\text{H}_4(\text{NH}_2\text{Cl})\text{NHPh}$ with potassium cyanate at 100° (Goldenring, *B.* 23, 1173). Rose-coloured needles, v. sol. alcohol. Decomposes at 120° in NH_3 and phenyl-trimethylene-urea.

PHENYLAMIDO-PYROTARTARIC ACID
 $\text{CO}_2\text{H.C}_6\text{H}_4\text{CMe}(\text{NHPh})\text{CO}_2\text{H}$ *Anilide methylsuccinic acid* [102°] Obtained by saponification of its ether (Schiller-Wechler, *B.* 18, 1037). White needles (containing aq), v. sol. hot water and alcohol, nearly insol. ether. Its solution in KOH aq is colourless. NH_4aq forms a violet solution. — CuA'' aq — CuA''NH_2 green pp — $\text{H}_2\text{A''HCl aq}$ white crystalline solid

Ethyl ether of the mono-amide
 $\text{CO}_2\text{Et.C}_6\text{H}_4\text{CMe}(\text{NHPh})\text{CONH}_2$ [125°] Formed by digesting $\text{CO}_2\text{Et.C}_6\text{H}_4\text{CMe}(\text{OH})\text{ON}$ with aniline in ether, dissolving the resulting compound $\text{CO}_2\text{Et.C}_6\text{H}_4\text{CMe}(\text{NHPh})\text{ON}$ in H_2SO_4 , and pouring into water. Plates, sol. hot water. Converted into the imide by heating with water, alcohol, acids, or alkalis

Imide $\text{CH}_2\text{CO} > \text{NH}$. $\text{NHPh.CMe.CO} > \text{NH}$. [150°]. Formed as above. White prisms, sol. alcohol and hot water. Yields a nitrosamine $\text{C}_6\text{H}_4(\text{NO})\text{N}_2\text{O}_2$ [173°], an acetyl derivative [285°], and a benzoyl derivative [190°]

Methylimide $\text{CH}_2\text{CO} > \text{NMe}$. $\text{NHPh.CMe.CO} > \text{NMe}$ [108°] Formed from the imide, MeI , MeOH , and KOH . Prisms. Yields a nitrosamine $\text{C}_6\text{H}_4\text{N}_2\text{O}_2$, crystallising in needles [147°]. Forms also $\text{C}_6\text{H}_4\text{MeN}_2\text{O}_2\text{Me.PtCl}_6$, which is sol. aq

Phenylimide $\text{CH}_2\text{CO} > \text{NPh}$. So-called 'n-Phenyl- α -keto- γ -oxy- β - α -dimethyl- β -anilido- α -tetra-hydropyridine carboxylic lactone.' [181°] A product of the action of heat on phenylamido-pyrotartaric acid (Reissert, *B.* 21, 1886; Anschütz, *A.* 261, 140). Prisms (from alcohol). Its acetyl derivative [169°] forms tabular crystals.

Mono-crystals
 $\text{C}_6\text{H}_4(\text{NHPh})(\text{CO}_2\text{H})(\text{CONHPh})$. [150°]. Got by boiling the phenylimide with NaOHAq . Forms with nitrous acid a compound melting at 204° .

Acetyl derivative of the anhydride
 $\text{CH}_2\text{CO} > \text{O}$. $\text{NACPh.CMe.CO} > \text{O}$ [186°] Got by boiling the acid with AcCl (Anschütz) Tables, v. sol. chloroform, al sol. ether

Acetyl derivative of the anilide
 $\text{C}_6\text{H}_4(\text{NACPh})(\text{CO}_2\text{H})(\text{CONHPh})$ [141°] Formed from the preceding body by the action of aniline on its ethereal solution. Prisms

PHENYL-(Py β) AMIDO-QUINOLINE
 $\text{C}_6\text{H}_4\text{CH}(\text{OH})\text{NCHNHPh}$ *Phenyl-quinolinamine* [98°] (above 360°). Formed by heating (Py β)-chloro-quinoline with aniline (Fiedler *id.* a. Weinberg, *B.* 18, 1532). White plates

Reference.—BROMO-PHENYL-AMIDO-QUINOLINE.
PHENYL-AMIDO-QUINOLINEQUINONE-

ANILIDE $\text{CH}(\text{NHPh})\text{C}(\text{CH}_2\text{NHPh})_2\text{CO} > \text{NCH}$ [222°] Formed by adding excess of aniline to an alcoholic solution of $\text{CH}(\text{COCl})\text{CO} > \text{C}_6\text{H}_4\text{N}$ (Hebe-

brand, *B.* 21, 2986). Long needles (from alcohol and HOAc), forms a deep-blue solution in dilute HCl aq — B'HOI . Dark-golden needles — Acetate [199°]. Bronzed needles, decomposed by water — $\text{B'CH}_2\text{N}_2\text{O}_2$. Coppery needles

PHENYLAMIDO-QUINONE *Dianilide*
 $\text{O}_2\text{H}_2(\text{NPh})_2(\text{NHPh})$ [142] [280°] Formed from azophenine and HOI at 160° (Fischer & Hepp, *A.* 256, 261). Reddish-brown needles, forming a blue solution in conc. H_2SO_4 .

Di-phenyl-di-amido-quinone
 $\text{C}_6\text{H}_4(\text{NHPh})_2$ [1435] *Quinone dianilide*
 Formed, together with hydroquinone, by boiling quinone with an alcoholic solution of aniline (Hofmann, *Pr.* 18, 4; Wichelhaus, *B.* 5, 851; Zincke, *B.* 16 1556; Knapp & Schultz, *A.* 210, 178). Formed also by heating chloro-quinone with aniline and HOAc (Niemeyer, *A.* 228, 332) and by strongly heating aniline with di-oxy-quinone (Nietzki & Schmidt, *B.* 22, 1655). Small violet plates, sol. HOAc and aniline, nearly insol. alcohol. May be sublimed. Conc. H_2SO_4 forms a reddish violet solution

Anilide $\text{C}_6\text{H}_4(\text{NHPh})\text{C}(\text{O})\text{NPh}$. [208°].

A product of the action of aniline on quinone in HOAc (Zincke & Hagen, *B.* 18, 785), on o-nitrophenol (Fischer & Hepp, *A.* 262, 247), and on quinone phenylimide (Bandrowski, *M.* 9, 415). Reddish-brown needles, forming a blood-red solution in H_2SO_4 . On warming with alcoholic H_2SO_4 , it yields $\text{C}_6\text{H}_4(\text{NHPh})(\text{OEt})\text{C}(\text{O})\text{NPh}$. Alcoholic potash forms $\text{C}_6\text{H}_4\text{N}_2\text{O}_2$, crystallising in red needles [192°], v. sol. alcohol, and forming a green solution in H_2SO_4 .

Di-anilide $\text{C}_6\text{H}_4(\text{NHPh})_2(\text{NPh})_2$ is **ASOPHENIDE**

DI-PHENYL-DI-p-AMIDO-QUINONE DI-CARBOXYLIC ETHER

$\text{C}_6(\text{NHPh})_2\text{O}_2(\text{C}_6\text{H}_4)_2$ [1,4 2 5 8 6] *Di-anilido-quinone di-carboxylic ether*. [246°]. Formed by the action of aniline in alcoholic solution upon p-di-chloro-quinone-di-carboxylic ether $\text{C}_6\text{Cl}_2\text{O}_2(\text{CO}_2\text{Et})_2$ (Hantzsch & Zeekendorf, *B.* 20, 1812). Glistening garnet-red crystals.

PHENYLAMIDO-SUCCINIC ACID
 $\text{CO}_2\text{H.C}_6\text{H}_4\text{CH}(\text{NHPh})\text{CO}_2\text{H}$ *Phenylsuccinic acid*. [183°] Formed by boiling bromo succinic

acid with aniline, and by the decomposition of an aqueous solution of acid maleate of aniline (Anschütz a. Wirtz, *Am* 9, 248, *A* 289, 140) Small crystals— $\text{H}_2\text{A} \cdot \text{HCl}$ monoclinic prisms.

Phenylimide $\text{C}_6\text{H}_5(\text{NHPh})\text{C}_2\text{O}_2\text{NPh}$. [211°]. Formed by heating maleic anhydride with aniline. Formed also, together with the dianilide $\text{C}_6\text{H}_5(\text{NHPh})(\text{CONHPh})$, [206°], by heating asparagine with aniline (Plutti, *G* 14, 474) Small needles (from alcohol). Yields a nitrosamine [180°]. Alcoholic NH_2 forms the amide-anilide [200°]. The dianilide yields a nitrosamine [190°].

Imide. [158°]. Formed from bromo-succinimide and aniline (Kusseroff, *A* 252, 158)

Ethyl ether $\text{C}_6\text{H}_5(\text{NHPh})(\text{CO}_2\text{Et})_2$. [214°] Oil, forming a crystalline sulphate

Di-phenyl-di-amido-succinic acid $\text{OO}_2\text{H} \cdot \text{CH}(\text{NHPh})\text{CH}(\text{NHPh})\text{CO}_2\text{H}$. [c 190°] Formed by saponifying its ether, which is formed by heating di-bromo succinic ether with aniline in alcohol (Lopatine, *C* R 105, 230; Gorodetzky a. Hell, *B* 21, 1796) Plates (from HOAc), nearly insol. water, m. sol alcohol

Ethyl ether Et_2A . [150°] Needles. Forms with Br a hexa-bromo-derivative [104°]

PHENYL-AMIDO-SULPHO-BENZOIC ACID $\text{C}_6\text{H}_5(\text{NHPh})(\text{SO}_3\text{H})\text{CO}_2\text{H}$ [124] Formed from bromo sulpho benzoic acid and alcoholic aniline (Fischer, *B* 24, 3802) Plates— BaA 5sq— $(\text{NH}_2\text{Ph})\text{HA}$ needles (from water)

PHENYL-AMIDO-THIAZOLE $\text{C}_6\text{H}_5\text{N}_2\text{S}$ t.s. $\text{S.O}(\text{NHPh})\text{CH}=\text{CH} > \text{N}$ [126°] Formed by the action of phenyl-thio-urea on di-chloro-di-ethyl ether (Hantzsch a. Traumann, *B* 21, 940, *A* 249, 47) Small white needles, sl sol water, v sol alcohol

PHENYLAMIDO-TOLUQUINONE $\text{C}_{11}\text{H}_9\text{NO}$, t.s. $\text{C}_6\text{H}_5\text{Me}(\text{NHPh})\text{O}_2$. [145°] Formed in small quantity, together with di-phenyl-amido toluquinone and its phenylimide, by the action of aniline in alcohol and HOAc on toluquinone (Hagen a. Zincke, *B* 16, 1559) Red needles (from dilute alcohol)

Anilide $\text{C}_6\text{H}_5\text{Me}(\text{NHPh})\text{C}_6\text{H}_4\text{N}$ [151°]. Formed by reducing the anilide of di-phenyl-di-amido-toluquinone with SnCl_2 and HOAc (Fischer a. Hepp, *A* 256, 259) Red needles

Di-phenyl-di-amido-toluquinone $\text{C}_6\text{H}_5\text{Me}(\text{NHPh})_2\text{O}_2$. [238°] Formed as above Brown needles (from alcohol), forming a red solution in H_2SO_4

Anilide $\text{C}_6\text{H}_5\text{Me}(\text{NHPh})\text{C}_6\text{H}_4\text{N}$. [178°]

Toluquinone trianilide. Formed as above, and also by heating nitroso-o-cresol (tolu-quinone metho-oxum) (1 pt.) with aniline (4 pts.) and aniline hydrochloride (3 pts.) (O. Fischer a. Hepp, *B* 21, 678) Brown plates with bluish lustre, sol hot alcohol—B'HI. Brownish needles.—B'HI, PtCl₄—B'HBz dark-green needles.

PHENYLAMIDO-TOLYL-ACETIC ACID $\text{C}_6\text{H}_5\text{NO}_2$ t.s. [13] $\text{C}_6\text{H}_5\text{Me}(\text{NHPh})\text{CO}_2\text{H}$. [159°]. Formed by saponifying its nitrile, which is made by heating $\text{C}_6\text{H}_5\text{CH}(\text{OH})\text{CN}$ with aniline in ether at 100° (Bornemann, *B* 17, 1471) Silvery plates (from dilute alcohol), v sol. hot Aq.

Amide $\text{C}_6\text{H}_5\text{N}(\text{CO}_2\text{NH}_2)$ [128°]. Plates.

Nitrile $\text{C}_6\text{H}_5\text{N} \cdot \text{CN}$. [95°]. Plates.

PHENYL AMIDO-TOLYL KETONE

Phthalyl derivative $\text{C}_6\text{H}_5\text{CO} \cdot \text{C}_6\text{H}_4\text{MeN} \cdot \text{C}_6\text{H}_5$. [202°] Formed, together with an isomeride [160°], by heating the phthalyl derivative of *p*-toluidine with benzoyl chloride and ZnCl_2 (Fröhlich, *B* 17 2872). Dimetric crystals, v sol. hot HOAc

DI-PHENYL-AMIDO-*p*-TOLYL-UREA [184] $\text{C}_6\text{H}_5\text{Me}(\text{NH}_2)\text{NH} \cdot \text{CO} \cdot \text{NPh}_2$. [137°]. Formed by reducing the nitro-compound in alcoholic solution with SnCl_2 and HCl (Lellmann a. Bonhöffer, *B* 20, 2123) Needles, solidifying at 165°–170°, and melting a second time at 220°–260°, yielding diphenylene and tolylene-urea at 300°

PHENYLAMIDO ISOVALERIC ACID

$\text{CHMe} \cdot \text{CH}(\text{NHPh})\text{CO}_2\text{H}$ S. 7 at 100°. Formed from bromo isovaleric acid and aniline (Duvillier, *A* Ch. [5] 21, 446, *C* R 88, 425) Scales (from hot water), v sol alcohol and ether—B'HI needles, decomposing at 100°–110°

DI-PHENYL-*o*-DI-AMIDO-*o*-XYLENE $\text{C}_6\text{H}_5(\text{CH}_2\text{NHPh})_2$. [172°] Formed by boiling *o*-xylylene bromide with an alcoholic solution of aniline (Leser, *B* 17, 1825) Small colourless plates Weak base.

PHENYLAMINE v ANILINE

Diphenylamine $\text{C}_6\text{H}_5\text{N} \cdot \text{C}_6\text{H}_5$ NPh_2 . Mol. w 169 [54°] (302°) (Graebe, *A* 236, 362). S.V 208 4 (Lossen, *A* 254, 72), 208 8 (Ramsay)

Formation—1 By the dry distillation of tri-phenyl-rosaniline (Hofmann, *A* 132, 168)—2 By heating aniline (3 mols.) with aniline hydrochloride (2 mols.) for thirty hours at 210° (De Laire, Girard, a. Chapoteaut, *Bl* [2] 7, 360 *C* R 74, 811, 1254, Merz a. Weith, *B* 5, 268; 6, 1511)—3 By distilling *o*-di-phenyl-urea or tri-phenyl-urea (Michler, *B* 9, 715)—4 By heating phenol with aniline-zinc-chloride to 260° (Merz a. Weith, *B* 13, 1298)—5 By heating a mixture of aniline, phenol, and SbCl_5 at 150°–160° (Buch, *B* 17, 2639)

Preparation.—By heating aniline with aniline hydrochloride at 230° and treating the warm product with HClAq followed by water

Properties—Monoclinic plates Forms a colourless solution in H_2SO_4 , changing on warming to blue, when a trace of nitrous or nitric acid is present (test for nitrous acid in H_2SO_4 : Kopp, *B* 5, 284, Plutti, *A* 227, 181) A solution in conc. H_2SO_4 (5 c c) is coloured blue by shaking with a solution (1 c c) containing nitric acid (test for nitric acid Muller, *Bl* [3] 2, 670). With PbO_2 and alcoholic HOAc it gives a bright olive green colour (Lanth, *C* R 111, 976). Picryl chloride forms dark-red needles of $\text{NPh}_2\text{SO}_3\text{H} \cdot (\text{NO}_2)_2\text{Cl}$ [66°] (Herr, *B* 23, 2540)

Reactions—1 When passed through a red-hot tube it yields carbazole, aniline, benzene, and NH_3 (Graebe, *A* 174, 177)—2 COCl_2 forms NPh_2COCl (Michler, *B* 8, 1664)—3 ClCO_2Et forms $\text{NPh}_2\text{CO}_2\text{Et}$ [73°]—4 Chlorine in presence of I yields $\text{NH}(\text{C}_6\text{H}_5\text{Cl}_2)$, and finally $\text{C}_6\text{H}_5\text{Cl}_2$ (Ruoff, *B* 9, 1488)—5 Bromine and I yield tetra-, hexa-, octo-, and deca-bromo-diphenylamines (Gessner, *B* 9, 1506)—6 Alkaline KMnO_4 yields oxalic acid, a resin, and a substance $\text{C}_{12}\text{H}_9\text{N}_2$, crystallising in yellow needles [176°–180°], which yields quinone on oxidation by MnO_2 and conc. HNO_3 , may be reduced to *p*-phenylene-di-phenyl-diamine [185°], and yields a (hexa ?)-bromo-

derivative [248°] (Bandrowski, *M* 7, 375, 8, 475, 9, 418) —7 Yields acridine on warming with AlCl_3 and chloroform HOAc and ZnCl_2 give methylacridine, and other acids act in like manner —8. PCl_5 and ZnCl_2 at 250° yield $\text{C}_{12}\text{H}_9\text{NPO}$, a white powder (Michaelis, *B* 21, 1504, *A* 260, 89) —9 *Acrolein* in alcoholic solution forms $(\text{C}_6\text{H}_5\text{N})_2\text{C}_2\text{H}_4$, an amorphous powder, sol chloroform, capable of combining with Br (Leeds, *B* 16, 1158, *A C J* 4, 32) —10 Heated with oxalic acid it gives diphenylamine blue —11 Heated with sulphur it gives imido-di-phenyl sulphide —12. HNO_3 forms a hexa-nitro-derivative which decomposes carbonates

Salts — B^+HCl^- needles (from alcohol), decomposed by water — $\text{B}^+\text{H SO}_4^-$ [125°] Insol ether and benzene, decomposed by water — Benzene sulphate [117°] —Toluene *p*-sulphonate [64°] (Norton, *Am.* 10, 129, 140)

Formyl derivative $\text{C}_6\text{H}_5\text{NO}$ *see* NPh, CHO [74°] Formed by heating diphenylamine with oxalic or formic acid (Willm & Girard, *B* 8, 1195) Insol water, sol benzene and alcohol Yields acridine on heating with ZnCl_2 .

Acetyl derivative NPh, Ac [108°] Plates (from ligroin), sol. hot water Yields NPh, Bz on heating with BzCl (Pictet, *B* 23, 3013) PCl_5 followed by water forms an acid $\text{C}_6\text{H}_4\text{ONPh}$, aq and an oil converted by alcoholic NH_3 into $\text{C}_{12}\text{H}_9\text{N}_2\text{O}$ [186°] (Claus, *B* 14, 2367)

Thio-acetyl derivative NPh, CS CH₃ [111°] Formed by heating *u*-di-phenyl-acetamide with CS_2 at 100° (*B*) Tables, sl sol water, v e sol ether

Benzoyl derivative NPh, Bz [177°] Formed from diphenylamine and BzCl (Hofmann, *A* 132, 166, Bernthsen, *A* 192, 13, 224, 12, Wallach, *A* 214, 235), and also by the action of Ph, N COCl on benzene in presence of AlCl_3 (Lellmann & Bonhöffer, *B* 19, 3231) Trimetric needles, *a b c* = 950 1 324 Sl sol ether and water, m sol hot alcohol

p-Toluyyl derivative. [155°]

Di-methyl-benzoyl derivative [43 1] $\text{C}_6\text{H}_4\text{Me}_2\text{CONPh}$ [136°] Formed from *o*-xylene, NPh, COCl, and AlCl_3 (Lellmann, *B* 20, 2119)

Nitrosamine NPh, NO [66 5°] Yellow four-sided tables (from benzene-alcohol) (Witt, *B* 8, 855, Fischer, *A* 190, 174) Yields azophenine when heated with aniline and aniline hydrochloride at 120° Its hydrochloride heated with *p* bromo-aniline at 80° yields tetra-bromo-azophenine $\text{C}_{12}\text{H}_8\text{Br}_4\text{N}_2$ [243°] (Ikuta, *A* 243, 285) On heating with aniline it yields PhN, $\text{C}_6\text{H}_4\text{NH}_2$, PhN, NPh, and diphenylamine by heating with alcoholic HCl it is converted into nitroso-diphenylamine.

Tri-phenyl-amine $\text{C}_6\text{H}_5\text{N}$ *see* NPh, Mol w 246. [127°]. Formed by dissolving K or Na in aniline or diphenylamine and digesting the product with bromo-benzene (Merz & Weith, *B* 6, 1514, Heydnich, *B* 18, 2156) Monoclinic crystals (from ether), *a b c* = 991 1: 412, β = 88° 38' Sl. sol hot alcohol, m sol benzene. Does not form salts. AcCl at 100° gives a greenish substance. Cold conc. H_2SO_4 gives a violet colour, changing to blue. Benzotrichloride and ZnCl_2 give a green colouring matter on heating *A* solution in

HOAc is coloured green by a little HNO_3 . Chlorine and I yield $\text{N}(\text{C}_6\text{Cl}_5)_3$, and finally C_6Cl_5 (Ruoff, *B* 9, 1438)

Isomeride of triphenylamine. $\text{C}_{12}\text{H}_9\text{N}$ Got by distilling the compound of cinnamic aldehyde with $(\text{NH}_4)_2\text{HSO}_4$ (Gössmann, *A* 100, 57) Liquid, volatile in a current of H at 140°–140°, sl sol. water, v sol alcohol and ether Forms unstable salts, decomposed by water and alcohol EtI yields B^+EtI^- , whence moist Ag_2O yields only B^+EtOH^- , while platinum chloride forms $\text{B}^+\text{Et}_2\text{PtCl}_4^-$ — $\text{B}^+\text{H}_2\text{PtCl}_4^-$ monometric crystals — $\text{B}^+\text{PtCl}_4^-$

References — AMIDO, DIBROMO, BROMO-NITRO-, CHLORO, CHLORO NITRO-, NITRO, and OXY, DIPHENYLAMINE

DIPHENYLAMINE SULPHONIC ACID $\text{C}_6\text{H}_5\text{NHSO}_3$ *see* NHPH $\text{C}_6\text{H}_5\text{SO}_3\text{H}$ [200°]. Formed, together with the disulphonic acid, by heating diphenylamine with H_2SO_4 at 160° (Merz & Weith, *B* 5, 283, 6, 1512) Formed also by heating NPh, HSO_3H , at 190° for 2 hours (Vignon, *C R* 107, 263) Crystalline mass, becoming blue in air — KA' plates — BaA'_2 plates, sl sol water — PbA'_2 nodules, sl sol water

Diphenylamine disulphonic acid $\text{NH}(\text{C}_6\text{H}_4\text{SO}_3\text{H})_2$ — BaA'_2 2aq nodules, v e sol water Hydrolysed by HCl aq at 200° (Girard, *Bl* [2] 23, 2)

Diphenylamine disulphonic acid $\text{C}_6\text{H}_5(\text{NHPH})(\text{SO}_3\text{H})_2$ [12 4] Formed from bromobenzene disulphonic acid and aniline in glycerol (Fischer, *B* 24 3807) V e sol aq — BaA'_2 3aq

Anilide $\text{C}_6\text{H}_5(\text{NHPH})(\text{SO}_2\text{NHPH})_2$ [223°] Yellowish crystals, insol water

Triphenylamine trisulphonic acid $\text{N}(\text{C}_6\text{H}_4\text{SO}_3\text{H})_3$ Formed by adding powdered triphenylamine to fuming H_2SO_4 at 60° (Herz, *B* 23, 2541) — $\text{Na}_3\text{A}'''$ crystalline powder (from 95 p c alcohol), v e sol water, insol alcohol

PHENYL-AMMELINE *v* Cyanuric acid in the article CYANIC ACID

PHENYL-AMYL-AMINE *see* AMYL ANILINE Di-phenyl-isomyl-amine $\text{NPh}_2\text{C}_6\text{H}_{11}$ [330°–340°] Formed from diphenylamine, amyl alcohol, and HCl (Girard, *Bl* [2] 23, 2) Gives a blue colouring matter when heated with oxalic acid and H_2SO_4

DIPHENYL-DI-ISOMYL-TETRAZINE $\text{NPh}(\text{C}_6\text{H}_{11})\text{N}$, $\text{NPh}(\text{C}_6\text{H}_{11})$ [86 5°] Formed from phenyl isomyl hydrazine in ether and HgO (Michaelis & Philips, *A* 252, 286) Yellowish crystals

PHENYL-AMYLENE CHPh CHPr (210°–215°) Formed by passing bromine vapour into amyl benzene at 150° and distilling the product (Schramm, *A* 218, 392) Yields a dibromide [54°]

Phenyl-isomylene CHPh CHPr (201°) at 787 mm S G \approx 878 Made in like manner from isomyl-benzene (*S*) Yields a dibromide [129°]

Phenyl-amylene CHPhEt CH CH_3 (173°) S G \approx 846 Formed by boiling $\text{CHPhEt.C}_6\text{H}_5\text{Br}$ with water or alcoholic potash (Dafert, *M* 4, 621). Liquid, readily converted into the polymeride C_6H_5 , (208°–213°), S G \approx 96°, V D. 10.2 (calc 10.1).

DI-PHENYL-AMYLENE DIKETONE $(\text{C}_6\text{H}_5\text{CO CH}_2\text{CH}_2)_2\text{CH}_2$ *see* Di-benzoyl-pentane. [68°]. Formed by the action of boiling KOH is

MeOH on $\text{CH}_2\text{Bz CH}_2\text{CH}_2\text{CH}_2\text{CHBz CO}_2\text{Et}$, which is got from sodium benzoyl-acetic ether and phenyl-bromo-butylketone (Kipping, W H Perkin, jun, *O J* 55, 849) Long colourless needles, insol. water, m. sol. cold alcohol

Ds. osm [176°]. Small needles

PHENYL-ISOAMYL-HYDRAZINE

$\text{NPh}(\text{C}_6\text{H}_5)_2\text{NH}_2$ (260°) Formed from iso-amyl bromide and sodium phenyl-hydrazine in benzene (Michaelis & Philips, *A* 252, 284) Reduces warm Fehling's solution. Ac_2O yields $\text{NPh}(\text{C}_6\text{H}_5)_2\text{NHAc}$ [125°] crystallising in scales

DI-PHENYL-AMYLIDENE DISULPHONE $\text{C}_6\text{H}_5(\text{SO}_2\text{Ph})_2$ [181°] Formed from NaOH , $\text{CH}_3(\text{SO}_2\text{Ph})_2$ and EtI (Fromm, *A* 253, 163) Crystalline, v. sol. hot alcohol.

PHENYL AMYL KETONE $\text{C}_6\text{H}_5\text{COCH}_2\text{Pr}$ (240°) at 720 mm Formed by heating isobutyl-benzoyl-acetic ether with dilute alcoholic potash (W H Perkin, jun, a Calman, *C J* 49, 166) Oil, with aromatic odour

PHENYL AMYL KETONE CARBOXYLIC ACID $\text{C}_6\text{H}_5\text{COCH}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{CO}_2\text{H}$ [82°] Formed by boiling the compound $\text{CH}_2\text{Bz CH}_2\text{CH}_2\text{CH}_2\text{CHBz CO}_2\text{Et}$ with KOH in MeOH (Perkin a Kipping, *C J* 55, 350) Plates (from light petroleum), or needles (from water) Yields an oxim [75°] Ag^+ amorphous pp

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PHENYL ISOAMYL OXIDE $\text{C}_6\text{H}_5\text{O C}_4\text{H}_9$ (225°) Got from phenol (Cahours, *A* 78, 227)

DI-PHENYL-ISOAMYL-PHOSPHINE OXIDE $\text{P}(\text{C}_6\text{H}_5)_2(\text{C}_4\text{H}_9)_2\text{O}$ [97°] Formed from $\text{PPH}_3\text{C}_4\text{H}_9$ and Ag_2O (Michaelis & Soden, *A* 229, 317) Needles, sol water and ether

DI-PHENYL-ISOAMYL-THIO-SEMICARBAZIDE $\text{C}_6\text{H}_5\text{NPh NHCS NPh}$ [160°] Formed from phenyl isoamyl-hydrazine and phenyl thiocarbimide (Michaelis & Philips, *A* 252, 285) Yellow needles

PHENYL-AMYL-THIO UREA $\text{C}_6\text{H}_5\text{NH}_2\text{N}_2\text{S}$ $\text{NHPh CO NH CH}_2\text{CMe}_2$ [136°] Formed from the corresponding amylamine and phenyl-thiocarbimide (Freund & Lenze, *B* 28, 2868)

PHENYL AMYL UREA $\text{C}_6\text{H}_5\text{NH}_2\text{N}_2\text{O}$ $\text{NHPh CO NH CH}_2\text{CMe}_2$ [155°] Formed from phenyl cyanate and the corresponding amylamine in alcoholic solution (Freund & Lenze, *B* 23, 2867, 24, 2158) White needles

PHENYL-ANGELIC ACID $\text{C}_{11}\text{H}_{12}\text{O}_4$ $\text{CHPh CHEt CO}_2\text{H}$ Mol w 176 [104°] Formed by heating benzoic aldehyde with sodium butyrate and Ac_2O at 100° (Perkin, *C J* 31, 391, 32, 661, 35, 136, *Slocum*, *A* 227, 53) Formed also by heating benzoic aldehyde with butyryl chloride at 125° (Fittig, *A* 153, 364) Needles, v sol ligroin (unlike cinnamic acid) Melts at 81° after having been fused — Ba^+ needles, m sol hot water — Ca^+ — Ag^+ white pp

Chloride $\text{C}_{11}\text{H}_{11}\text{COCl}$ Oil

Amide $\text{C}_{11}\text{H}_{11}\text{CONH}_2$ [128°]

Phenyl-angelic acid

$\text{CHPh CH CH}_2\text{CH}_2\text{CO}_2\text{H}$ Formed from $\text{CHPh CH OH CH CO}_2\text{H}$ by reduction with sodium-amalgam (Perkin, Baeyer & Jackson, *B* 18, 123) Liquid. — Ag^+ white pp

Phenyl-angelic acid $\text{CHPh CH CHMe CO}_2\text{H}$ [110 5°] Formed by boiling the dibasic acid $\text{CHPhBr CH}(\text{CO}_2\text{H})\text{CHMe CO}_2\text{H}$ with water (Penfield, *A* 216, 123), and by distilling $\text{CHPh} \begin{smallmatrix} \text{CH}(\text{CO}_2\text{H}) \\ \text{O} \text{---} \text{CO} \end{smallmatrix} \text{CHMe}$ (Fittig a Liebmman, *A* 255, 262) Plates (from water) — Ba^+ aq. needles, v sol water

Phenyl-angelic acid $\text{CHPh OMe CH}_2\text{CO}_2\text{H}$ [113°] Formed by distilling the lactonic acid $\text{CHPh} \begin{smallmatrix} \text{OMe}(\text{CO}_2\text{H}) \\ \text{O} \text{---} \text{CO} \end{smallmatrix} \text{CH}_2$ (F a L). Thin tables, sl sol water — Ba^+ , groups of needles, sol. water

Phenyl angelic acid Nitrate $\text{C}_6\text{H}_5\text{CHPh CN}$ (c 265°) Formed by heating phenyl-acetonitrile with NaOH and allyl iodide (Buddeberg, *B* 23, 2068) Oil, converted by NaOEt and benzyl chloride into the benzyl derivative $\text{C}_6\text{H}_5\text{CPh}(\text{CH}_2\text{Ph})\text{CN}$ (c 325°)

PHENYL-ANTHRACENE $\text{C}_{20}\text{H}_{12}$, $\text{c c C}_6\text{H}_5\text{Ph}$ [153°] (417°) Formed by heating phenyl-anthranol with zinc-dust (Baeyer, *A* 202, 61), and by the action of chloroform and AlCl_3 on benzene (Friedel, Crafts, a Vincent, *BI* [2] 40, 97; *A Ch.* [6] 1, 495) Leaflets (from alcohol), forming solutions with blue fluorescence Reduced by P and HI to a crystalline dihydride [120°], which is oxidised by CrO_3 in HOAc to phenyl-oxanthranol

PHENYL ANTHRANOL $\text{C}_{16}\text{H}_{10}\text{O}$ c c H $\begin{smallmatrix} \text{CPh} \\ \text{O}(\text{CH}) \end{smallmatrix} \text{C}_6\text{H}_5$ [141°-144°] Formed by dissolving tri phenyl methane *o*-carboxylic acid in H_2SO_4 and ppg with water (Baeyer, *A* 202, 57). Golden needles, sol hot alcohol Its ethereal solution shows greenish yellow fluorescence

Acetyl derivative [166°] Golden needles, turned red by conc KOHaq

Reference — *DI-CHLORO-PHENYL ANTHRANOL*

PHENYL-ARSINE *v* vol 1 p 319 The sulphides PhAsS [162°] and $\text{Ph}_2\text{As}_2\text{S}_3$ [130°] have been prepared by Schulte (*B* 15, 1956)

PHENYL AZIMIDO-COMPOUNDS *v* AZIMIDO-COMPOUNDS

DI-PHENYL-TETRAZINE $\text{C}_{11}\text{H}_{12}\text{N}_4$ c c $\text{NPh} \begin{smallmatrix} \text{N CH} \\ \text{CH N} \end{smallmatrix} \text{NPh}$ [180°] Mol w 240 by Raoult's method (calc 236) Formed by the action of chloroform and alcoholic potash on phenyl hydrazine (Ruhemann, *C J* 53, 850, 55, 243) White needles (from alcohol) HNO_3 yields $\text{C}_6\text{H}_5(\text{NO}_2)_2\text{N}$ (above 300°) H_2SO_4 forms $\text{C}_6\text{H}_5(\text{SO}_3\text{H})_2\text{N}$ Bromine gives $\text{C}_6\text{H}_5\text{Br}_2\text{N}$ [220°], $\text{C}_6\text{H}_5\text{Br}_2\text{N}$ [131°], and $\text{C}_6\text{H}_5\text{Br}_2\text{N}$, decomposing at 224°

Salts — B^+HCl needles — $\text{B}^+\text{H}_2\text{PtCl}_6$ — B^+MeCl [244°] Needles — $\text{B}^+\text{Me}_4\text{PtCl}_6$ — B^+MeI [214°] Yellow needles, v sol alcot l

PHENYL AZO-COMPOUNDS *v* Azo-compounds

PHENYL-TETRAZOLE CPhHN , c c $\text{N} \begin{smallmatrix} \text{N NPh} \\ \text{N CH} \end{smallmatrix} \text{N}$ Formed by heating its carboxylic acid [138°] at 155° (Bladin, *B* 18, 2907) Heavy oil, v sol. alcohol and ether Explodes when strongly heated Sol acids, but reppd on dilution

PHENYL-TRIAZOLE CARBOXYLIC ACID $\text{CH} \begin{smallmatrix} \text{N NPh} \\ \text{N C CO}_2\text{H} \end{smallmatrix}$ [184°]. Formed by boiling

phenyl-hydrazine dicyanide $\text{NH}_2\text{NPh CO}_2\text{NH}$ with formic acid and heating the resulting formyl derivative with alcoholic potash (Bladin, B 23, 3788). Silvery plates (from water).

Methyl ether MeA' [118°] Prisms.

Phenyl-triazole dicarboxylic acid

$\text{CO}_2\text{H C} \begin{smallmatrix} \text{N NPh} \\ \text{N C CO}_2\text{H} \end{smallmatrix}$ Formed by oxidising phenyl-methyl-triazole carboxylic acid with alkaline KMnO_4 (Bladin, B 23, 3785). The free acid is unstable, readily changing to the preceding acid. The acid K and Na salts also readily give off CO_2 .— CuA'' 4aq blue needles.— $\text{Ag}_2\text{A}''$ 3aq? Bulky white pp

Methyl ether MeA'' [167°] Needles

Ethyl ether EtA'' [82°] Needles

Di phenyl-triazole carboxylic acid

$\text{CPh} \begin{smallmatrix} \text{N NPh} \\ \text{N C CO}_2\text{H} \end{smallmatrix}$ Formed by saponification of the nitrile (Bladin, B 22, 797). Crystals (containing EtOH), decomposing at 172° – 182° . V sol. alcohol, m sol ether.— CuA' ,— AgA' white pp

Methyl ether MeA' [159°] Needles

Ethyl ether EtA' [165°] Needles

Nitrile $\text{C}_6\text{H}_5\text{CN}$ [156.5°] Formed from phenyl-hydrazine dicyanide, alcohol, and benzoin aldehyde. Needles, v sol benzene

Amide $\text{C}_6\text{H}_5\text{CONH}_2$ [196°]. Formed from the nitrile by the action of dilute (3 p.c.)

H_2O_2 and KOH aq . Plates (from alcohol) or needles (from water)

Amidozine $\text{C}_6\text{H}_5\text{C}(\text{NOH})\text{NH}_2$ [214°].

Formed from the nitrile and hydroxylamine

(Bladin, B 22, 1752). Prisms (from alcohol)

Yields an acetyl derivative [177°] which yields

$\text{C}_6\text{H}_5\text{Ph} \begin{smallmatrix} \text{N O} \\ \text{N} \end{smallmatrix} \text{CMe}$ [163°], and a benzoyl de

rivative [180°] which yields in like manner

$\text{C}_6\text{H}_5\text{Ph} \begin{smallmatrix} \text{N O} \\ \text{N} \end{smallmatrix} \text{CPh}$ [206°].— B'HCl : crystal

line

PHENYL-TETRAZOLE CARBOXYLIC ACID

$\text{N} \begin{smallmatrix} \text{N NPh} \\ \text{N C CO}_2\text{H} \end{smallmatrix}$ [188°]. Got by saponification

of its nitrile, which is formed by the action of nitrous acid on phenyl-hydrazine dicyanide

(Bladin, B 18, 2907). Colourless needles, v sol

alcohol.— KA' plates, v sol water.— CuA' , 2aq

— AgA' colourless crystalline pp

Methyl ether MeA' [116°]. Plates

Ethyl ether EtA' [74°]. Needles

Amide CN, Ph CO NH_2 [168°] Formed

from the nitrile by treatment with hydrogen

peroxide. Crystals, al sol cold water

Amidozine $\text{CN, Ph C}(\text{NOH})\text{NH}_2$ [177°].

Formed from the nitrile and hydroxylamine

(Bladin, B 22, 1755). Scales, v al sol water.

Yields an acetyl derivative [208°] and a benzoyl

derivative [206°], both crystallising in needles

END OF THE THIRD VOLUME.

